

Universität Heidelberg  
Interdisciplinary Center for Scientific Computing  
Engineering Mathematics and Computing Lab

Bachelor Thesis

# Continuous Modeling of Extracellular Matrix Invasion by Tumor Growth

Name: Maximilian Bing  
Matriculation number: 3606060  
Advisor: Professor Vincent Heuveline  
Date of submission: April 30, 2024

Hiermit versichere ich, dass ich die Arbeit selbst verfasst und keine anderen als die angegebenen Quellen und Hilfsmittel benutzt und wörtlich oder inhaltlich aus fremden Werken Übernommenes als fremd kenntlich gemacht habe. Ferner versichere ich, dass die übermittelte elektronische Version in Inhalt und Wortlaut mit der gedruckten Version meiner Arbeit vollständig übereinstimmt. Ich bin einverstanden, dass diese elektronische Fassung universitätsintern anhand einer Plagiatssoftware auf Plagiate überprüft wird.

---

Abgabedatum:

## Zusammenfassung

Krebszellen können sich vom Primärtumor lösen und das umgebende Gewebe abbauen. Kontinuierliche mathematische Modelle werden verwendet, um diesen Prozess besser zu verstehen.

In diesem Zusammenhang basieren die Modelle in der Regel auf mindestens drei Schlüsselkomponenten: den Tumorzellen, dem umgebenden Gewebe oder der extrazellulären Matrix (ECM) und den matrixabbauenden Enzymen (MDE). Diese Variablen werden in dem hier behandelten Modell in einem System partieller Differentialgleichungen gekoppelt, um den komplexen Prozess der Tumorinvasion zu beschreiben.

Aufgrund der hohen Anzahl der freien Parameter solcher Modelle, hat eine Konfiguration dieser Parameter hohen Einfluss auf die produzierten Ergebnisse. Diese Arbeit untersucht den Einfluss der Wahl dieser Konfigurationen, sowie den, der Dimension auf die Ergebnisse einer Simulation.

Um realitätsnahe biologische Szenarien zu simulieren, verlangen Simulationen mindestens zwei am besten drei räumlichen Dimensionen. Da in der Literatur fast ausschließlich eindimensionale Experimente beschrieben werden, untersuchen wir unser Modell nur in höheren Dimensionen: zwei- oder dreidimensional. Darüber hinaus wurde hauptsächlich die homogene Struktur der extrazellulären Matrix (ECM) behandelt, jedoch für eine heterogene extrazelluläre Matrix Struktur nur solche Fälle analysiert, die wenig biologische Relevanz haben. Die Struktur der epithelialen Schicht und der benachbarten extrazellulären Matrix ist jedoch in biologischem Gewebe deutlich organisierter als in den meisten später gezeigten Simulationen und anderen Beispielen aus der Literatur. Diese Organisiertheit kann zu erheblichen Veränderungen der Ergebnisse führen, selbst wenn die Parameter des Systems konstant gehalten werden.

Diese Arbeit soll einen Einstiegspunkt geben, um das zu grundliegende Modell für realistische Experimente zu nutzen. Die Parameteranalyse soll hierbei auch helfen eine angebrachte Wahl der Parameter zu erleichtern.

## Abstract

Cancer cells can detach from the primary tumor and degrade the surrounding tissue. Continuous mathematical models are been utilized for a better understanding of the process.

In this context the models typically rely on at least three key components: tumor cells, the surrounding tissue or extracellular matrix (ECM) and matrix-degrading enzymes (MDE). These variables are coupled in the investigated model in a system of partial differential equations to describe the complex process of tumor invasion.

Due to such a model's high degree of freedom, the configuration of the model parameters significantly influences the resulting outcomes. This study examines the influence of choosing these configurations, as well as the dimensionality, on the results of a simulation.

To simulate realistic biological scenarios, simulations require at least two, preferably three, spatial dimensions. Since the literature predominantly describes one-dimensional experiments, we only investigate our model in higher dimensions: two or three dimensions. Furthermore, primarily, the homogeneous structure of the extracellular matrix (ECM) has been addressed, but for a heterogeneous extracellular matrix structure, only cases with little biological relevance have been analyzed. However, the structure of the epithelial layer and the adjacent extracellular matrix is significantly more organized in biological tissue than in most simulations shown later and other examples from the literature. This organization can lead to significant changes in the results, even when the parameters of the system are kept constant.

This work aims to provide a starting point for using the fundamental model for realistic experiments. The parameter analysis should also help facilitate an appropriate choice of parameters.

## Contents

<b>1</b>	<b>Introduction</b>	<b>5</b>
<b>2</b>	<b>Theoretical Basics</b>	<b>6</b>
2.1	Basics of Tumor Biology . . . . .	6
2.2	Mathematical Methods in Oncology . . . . .	8
<b>3</b>	<b>Modeling</b>	<b>10</b>
3.1	Mathematical Formulation . . . . .	10
3.2	Numerical Formulation and Parameters . . . . .	12
<b>4</b>	<b>Experiments and Results</b>	<b>14</b>
4.1	Dimension Variation . . . . .	17
4.2	Parameter Analysis without Proliferation and Renewal - Homogenous ECM	24
4.2.1	Parameter Analysis . . . . .	26
4.3	2D Results with Proliferation and Renewal - Homogenous ECM . . . . .	44
4.3.1	Basecase Analysis . . . . .	46
4.3.2	Parameter Analysis . . . . .	47
4.4	2D Results with Proliferation and Renewal - Heterogenous ECM . . . . .	59
<b>5</b>	<b>Conclusion and Discussion</b>	<b>62</b>

## 1 Introduction

Modeling tumor growth plays a crucial role in understanding the complex mechanisms governing the development and progression of cancer diseases. Since cancer is one of the leading death causes worldwide and many of its forms are incurable, challenges in the area of Oncology require researchers to have a deep understanding in the biological foundation. This Bachelor thesis is dedicated to analyzing Anderson et al.'s [1, 2] model for modeling tumor invasion.

The dynamics of tumorous growth are an intricate system influenced by numerous biological and chemical factors, genetic pre-dispositions, the surrounding tissue of cancer cells, angiogenic processes and interactions with the immune system. Integrating these factors in mathematical models allows us to decode these complex interactions with quantification and helps us understand the fundamental mechanisms surrounding cancerous diseases.

Mathematical models are an essential part of Oncology; they are used to quantify biological phenomena and help to predict and understand tumor development and treatment response. In Mathematical Oncology, we differentiate between continuous, discrete and hybrid models [3].

For the continuous type, cells and tissue are described over time with partial differential equations modeling continuous quantities like, in our case, tumor cell density, extracellular matrix concentration or matrix-degrading enzyme concentration.

In the discrete case, an entity-based model is used, pursued with the goal of better understanding the phenomena on the cell level. This approach allows the researcher to better implement a cell's biological effects with its outer circumstances, like interaction with other cells, nutrients or other microorganisms. As the name implies, use these models discrete values to describe the temporal course of events.

Hybrid models try to combine both approaches to offer efficient systems capturing cell level events and continuous changes in outer circumstances.

This work investigates how a continuous model proposed by Anderson et al. [1, 2] to analyze tumor development performs in the case of different dimensions and free parameter values. The model examines the third and fourth stages of a cancer disease: tumor progression, where the tumor cells grow more extensive and take on more aggressive behavior due to further genetic instabilities and tumor invasion, with the tumor cells gaining the ability to penetrate and invade the surrounding tissue [4].

From examples of the original paper, we can already see that the model's results vary with the dimensionality of the space, we are modeling the partial differential equations in.

Our primary focus is investigating how the model's free parameters influence tumor dynamics growth.

Another point of interest is comparing simulations of two dimensions with those of three dimensions of extracellular matrix invasion by tumor growth.

Additionally to the variation of dimensions, we will briefly examine how the geometry of the extracellular matrix will influence tumor development.

## 2 Theoretical Basics

### 2.1 Basics of Tumor Biology

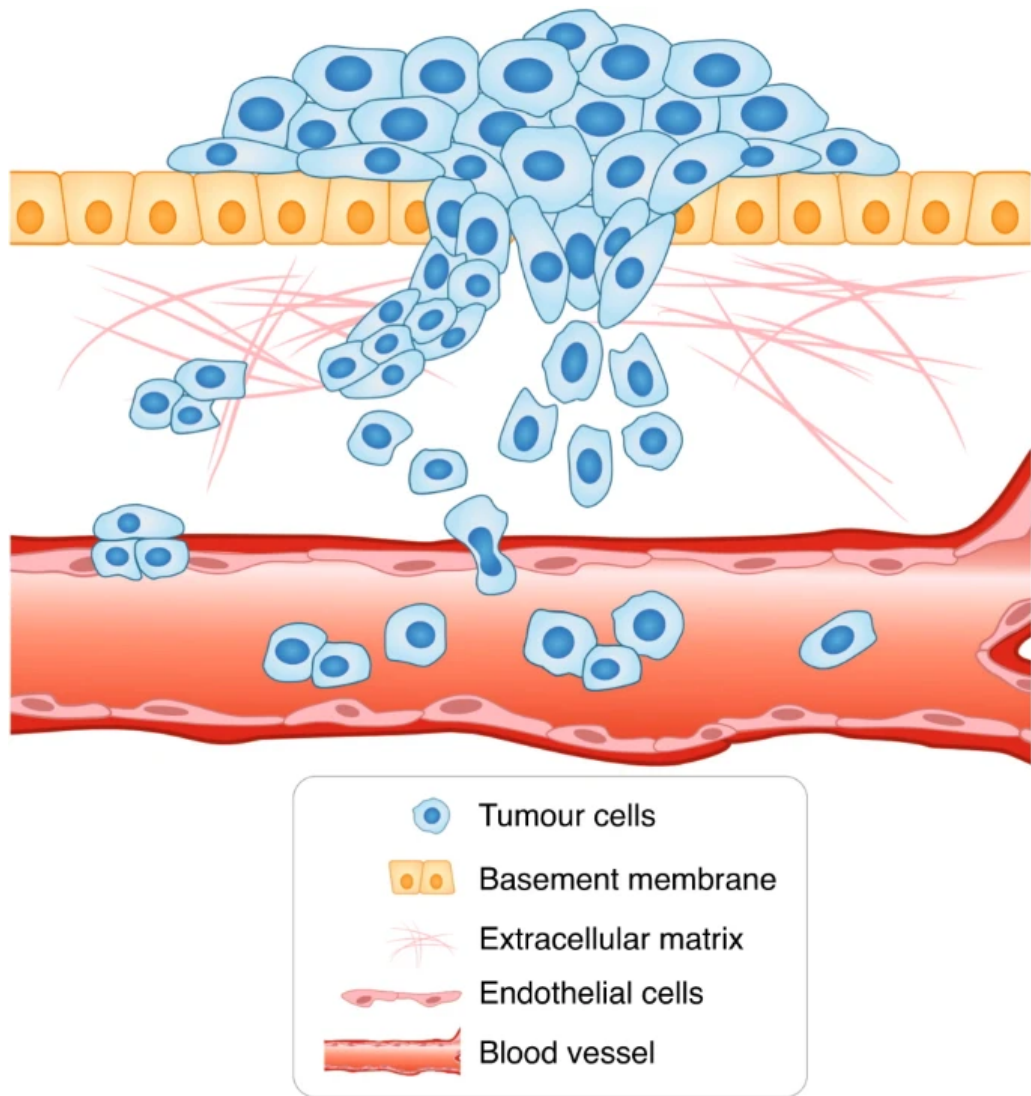


FIGURE 1. Tumor Invasion Stage

The body of a living human being is made up of more than 200 different types of cells; the coordination between the cells and their surroundings keeps the body running. Each of these cells is built from the genetic information encoded in the DNA in the cells' nuclei. Though the nucleotide sequence of DNA is well-checked and maintained throughout the cell's life, mutations still occur that cause changes in a cell's DNA. These mutations may be of a positive, negative or neutral nature. In the case of a harmful mutation, this alteration of the DNA may cause diseases, with cancer being one of them. The failure of the complex system managing cell birth, proliferation and cell death (apoptosis) causes cancer, resulting in uncontrolled cell proliferation in a local area. Abnormal growth of

body tissue is called a tumor, yet these can be malignant or benign.

Cancer diseases can be categorized medically into five stages. First is the tumor initiation phase, where it comes to the above explained genetic mutations of normal cells. The next stage is the tumor promotion stage, in which the mutated cells of phase one may experience further genetic alterations resulting from uncontrolled growth and proliferation of the cancerous cells. The third stage is the tumor progression stage, where the cancerous cells progress in growing and proliferating, reaching a critical mass and forming a tumor at a local site of the body. Fourth comes the invasion stage, shown in figure 1. Here, the tumor can invade surrounding tissue by breaking through the basement membrane, invading the extracellular matrix inside and entering the blood circulation or lymphatic systems. Next, the tumor cells that have invaded the blood circulation of the lymphatic system spread throughout the body and form new tumors. This stage is called Metastization. To grow tumors further, they need access to more nutrients and oxygen. During angiogenesis, a tumor develops its own blood vessels, securing its nutritional provision.

In our model, the focus lies on the third and fourth stages: tumor progression and tumor invasion. The tumor progression stage is characterized by the tumor growing more extensive and the cancerous cells taking on more aggressive behavior by starting to invade the surrounding area. While they keep growing uncontrolled, they are also affected by further genetic instabilities, which lead to more mutations, possibly developing resistance mechanisms against, for example, degrading factors. Already in this stage, the affected area is exposed to heavy tissue damage and functional disabilities. In the next stage, tumor invasion stage, the malignant cells gain the ability to penetrate and invade the surrounding tissue furhter. The tumor cells break through the normal tissue barrier and infiltrate neighboring structures. In order to do so, the cancer cells produce so-called matrix-degrading enzymes which break down the extracellular matrix. The degradation of the extracellular matrix helps local spreading and destroys otherwise healthy tissue and cells in the affected area

The most important factors influencing those two phases are the genetic dispositions of the tumor cells towards proliferation and the evasion of apoptosis, programmed cell death, which increase the invasive potential. Another critical factor is the geometry of the extracellular matrix, as well as the exact macromolecules that make it up. A solid immune biological defense reaction also helps the body defend against spreading cancer cells. Hence, evasion of detection and destruction of the tumor cells plays a vital role in the first stages. To invade the affected area, the malignant cells need to be able to move freely and quickly. In order to do so, cancer cells can gain the ability to lose adhesion properties, which healthy cells usually have, to allow migrating into the surrounding tissue.

Another organization principle for cancerous diseases is found in the Hallmarks of Cancer of Douglas Hanahan and Robert Weinberg [4], shown in figure 2. They describe a set of functional capabilities, eight hallmark capabilities and two enabling characteristics commonly acquired by cancer cells. They contribute to their ability to grow uncontrollably, evade the immune system and metastasize. These hallmarks are sustaining proliferative signaling, evading growth suppressors, avoiding immune destruction, enabling replicative immortality, tumor-promoting inflammation, activating invasion and Metastization, in-

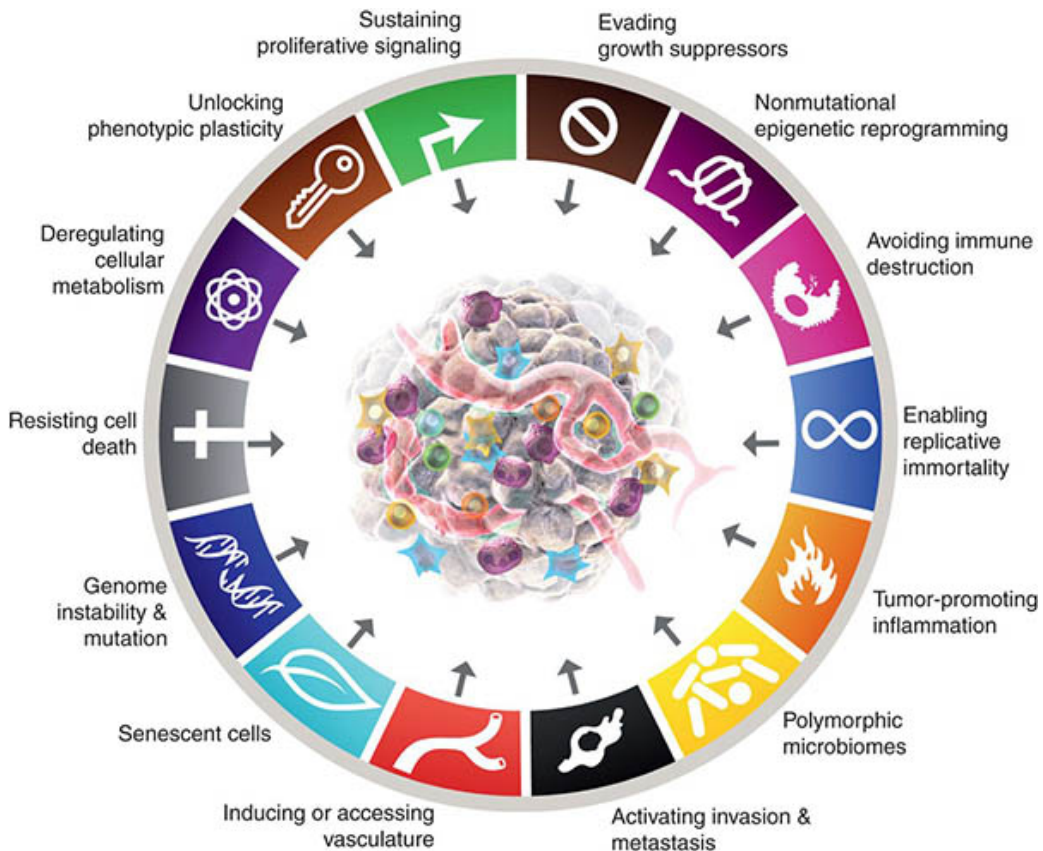


FIGURE 2. Hallmarks of Cancer by Hanahan and Weinberg [4]

ducing or accessing vasculature, genome instability and mutation, resisting cell death, deregulating cellular metabolism. These Hallmarks of Cancer together contribute to the development and progression of cancer and provide targets for therapeutic intervention and research efforts aimed at understanding and treating the disease.

In our model, we see many of the capabilities incorporated. Proliferative signaling only relies on the tumor cells themselves and not on typically other hormones or molecules, allowing them to proliferate rapidly without external growth signals. Immune destruction and induced cell death are avoided, optimizing them to survive and accumulate genetic mutations that promote tumor growth. The tumor cells invade the surrounding tissue, modeled by the extracellular matrix.

## 2.2 Mathematical Methods in Oncology

Mathematical methods and models in Oncology play a crucial role in analyzing, understanding and predicting cancer development. Since the objective of this research underlies complex and intricate biochemical systems and mechanisms, many models exist that find their respective applications in distinct areas of this research field. These methods can be coarsely divided into three sections: continuous, discrete and hybrid models [3].

For describing tumor growth, exponential and logistic growth models are often used,

the latter allowing limiting factors, such as space or nutritional supply to play a role during modeling. These methods are a subclass of the differential equations approach, which bases their functionality on an ordinary or partial differential equation, studying the continuous approach. Like our model, they are not limited to only one equation but can include many, incorporating systematic dependencies on other factors. These models generally deal with continuous quantities like densities or concentrations, for example, spatial and temporal nutritional supply or drug concentration, as well as their effects on the affected area over time.

Discrete models use discrete entities to describe the behavior of tumor cells and their interactions with surrounding tissue. They allow us to model a wide range of biological and chemical processes which are hard to describe continuously. Commonly used types in Mathematical Oncology are, for example, Cellular Automata or Agent-Based Models. Cellular Automata represents cells as entities with states on a grid, with each cell being allowed to change states according to a set of rules based on its own current state and the states of the neighbors. In contrast, Agent-Based Models enable the differentiation of cell types and allow movement that is not restricted to a grid, implementing complex mechanics at cell-cell or cell-environment levels. Using these discrete models allows researchers to focus on biological effects during modeling, which are hard to describe in continuous models. With these approaches, we can simulate genetic and evolutionary events, for example, they are studying the genetic alternations of tumor cells on cell level or the interaction between healthy and cancerous cells.

Hybrid models combine both methods above, using continuous and discrete approaches. Like in the model proposed by Franssen et al. [5] or Anderson et al. [1], these approaches allow to incorporate the exactness of continuous models with a wide range of biological effects described by discrete models. They for example allow to model the motility and proliferation of tumor cells to be modeled by continuous variables, while enabling effects such as the formation of blood vessels, angiogenesis.

However, not all models try to model tumor growth; others are concerned with optimality regarding drug dosages or radiation exposition, offering personalized treatment. Machine Learning and Data Mining methods are used analyzing large datasets to identify patterns and predict outcomes to optimize treatment for individual patients. Though these methods are used in all kinds of applications, also for, spacial and temporal cancer development. Putting all these methods together gives us a powerful toolbox to simulate and understand cancer biology. As the last years have shown, they are applied in a wide range of areas, offering insight into all areas of cancer research [6]. Therefore, it is essential to come up with methods and evaluate their usefulness and meaningfulness in different research areas.

## 3 Modeling

### 3.1 Mathematical Formulation

The model proposed by Anderson et al. [1, 2] and Chaplain et al. [1, 5, 7], extended with terms for modeling cell proliferation and extracellular matrix renewal consists of a system of coupled partial differential equations:

$$\frac{\partial c}{\partial t} = D_c \Delta c - \chi \nabla \cdot (c \nabla e) + \mu_1 c \left(1 - \frac{c}{c_0} - \frac{e}{e_0}\right) \quad (1)$$

$$\frac{\partial e}{\partial t} = -\delta m e + \mu_2 c \left(1 - \frac{c}{c_0} - \frac{e}{e_0}\right) \quad (2)$$

$$\frac{\partial m}{\partial t} = D_m \Delta m + \mu_3 c - \lambda m \quad (3)$$

with zero-flux boundary conditions:

$$\zeta \cdot (-D_c \nabla c + c \chi \nabla e) = 0 \quad (4)$$

$$\zeta \cdot (-D_m \nabla m) = 0 \quad (5)$$

where the free parameters are  $D_c$ ,  $D_m$ ,  $\chi$ ,  $\delta$ ,  $\mu_1$ ,  $\mu_2$ ,  $\mu_3$ ,  $\lambda$  and  $\zeta$  is an appropriate outward unit vector. These free parameters as well as their non-dimensionalized versions are all assumed to be positive real numbers.

The variable  $c$  describes the tumor cell density,  $e$  the concentration of the extracellular matrix and  $m$  the matrix-degrading enzyme concentration. All of those functions are mathematically defined to be mapping a one-, two- or three-dimensional spacial value  $x$  and a temporal value  $t$  to a scalar value describing the respective quantity at a specific point in space and time  $(x, t)$ :

$$\{c, e, m\} : \mathbb{R}^i \times \mathbb{R} \rightarrow \mathbb{R}, i \in \{1, 2, 3\}$$

To derive the expression of the tumor cell density  $c$ , we are going to assume that the tumor cell's motility is subject to two influences: haptotaxis and random movement. Haptotaxis is a directed migratory response of cells to gradients of fixed or bound chemicals [1] and random movement is influenced by, for example, mechanical stress, electric charge or other physical effects [8]. We must define flux to get an expression of how much or how fast the tumor cells move. Flux is defined to be the amount of a substance that crosses a unit area in a unit time. Incorporating the two assumed influencing factors into our mathematical model; we define the haptotactic flux  $J_{hapt}$  and random flux  $J_{random}$ :

$$\begin{aligned} J_{hapt} &= \chi c \nabla e \\ J_{random} &= -D_c \nabla c \end{aligned}$$

$\chi$  is the haptotactic flux coefficient and  $D_c$  is a random mobility coefficient. These parameters could also be a function of, for example, extracellular matrix and or matrix-degrading enzyme concentration. Knowing that cells grow over time and proliferate, we

want to respect this in our model with a term for tumor cell proliferation:

$$\mu_1 c \left(1 - \frac{c}{c_0} - \frac{e}{e_0}\right)$$

The idea is that this term describes the cell proliferation with a logistic growth model respecting spacial limiting factors of already present extracellular matrix molecules and tumor cells,  $\mu_1$  describes the rate at which tumor cell proliferation and growth happens. In the initial model proposed by Anderson et al. [1, 2] and Chaplain et al. [1, 5, 7], they did not respect proliferation of tumor cells and extracellular matrix renewal. They applied a conservation equation for the tumor cells, which yields:

$$\begin{aligned}\frac{\partial c}{\partial t} &= -\nabla \cdot (J_{hapto} + J_{random}) \\ \frac{\partial c}{\partial t} &= -\nabla \cdot (\chi c \nabla e - D_c \nabla c) \\ \frac{\partial c}{\partial t} &= D_c \Delta c - \chi \nabla \cdot (c \nabla e)\end{aligned}$$

The extended model incorporates proliferation and renewal into this conservation formula, resulting in equation 1:

$$\begin{aligned}\frac{\partial c}{\partial t} &= -\nabla \cdot (J_{hapto} + J_{random}) + \mu_1 c \left(1 - \frac{c}{c_0} - \frac{e}{e_0}\right) \\ \frac{\partial c}{\partial t} &= D_c \Delta c - \chi \nabla \cdot (c \nabla e) + \mu_1 c \left(1 - \frac{c}{c_0} - \frac{e}{e_0}\right)\end{aligned}$$

To model the extracellular-matrix concentration  $e$ , we assume that the enzymes degrade the extracellular matrix upon contact in an exponential way. The equation models this assumption:

$$\frac{\partial e}{\partial t} = -\delta m e$$

where  $\delta$  is the constant describing this degradation process. For the extended model we add a term describing the renewal process of the extracellular matrix, which is also being modeled by logistic growth incorporating limiting factors of present tumor cells and extracellular matrix molecules:

$$\frac{\partial e}{\partial t} = -\delta m e + \mu_2 c \left(1 - \frac{c}{c_0} - \frac{e}{e_0}\right)$$

with  $\mu_2$  being the coefficient describing the rate of the renewal process.

Modeling the matrix-degrading enzyme concentration  $m$ , we combine a diffusion term with production and decay terms. The diffusion term is described like for the tumor cells:

$$J_{random} = -D_m \nabla c$$

The production term depends on the tumor cell density  $c$  and the decay term on the extracellular matrix concentration  $m$ . In both cases production and decay we are assuming exponential growth. Incorporating this gives us equation 3:

$$\begin{aligned}\frac{\partial m}{\partial t} &= \nabla \cdot J_{random} + \mu c - \lambda e \\ \frac{\partial m}{\partial t} &= D_m \Delta c + \mu_3 c - \lambda m\end{aligned}$$

$\mu_3$  and  $\delta$  describing production and decay coefficients.

### 3.2 Numerical Formulation and Parameters

To make solving the model easier, we will first non-dimensionalize all the equations 1 to 5 in a standard way to rescale the space domain to a unit size domain  $\Omega$ . For one space dimension this results in the unit interval  $[0, 1]$ , for two the unit square  $[0, 1] \times [0, 1]$  and for three spacial dimensions the unit cube  $[0, 1] \times [0, 1] \times [0, 1]$ .

We start with non-dimensionalizing the three continuous variables  $c, e, m$ :

$$\begin{aligned}\tilde{c} &= \frac{c}{c_0} \\ \tilde{e} &= \frac{e}{e_0} \\ \tilde{m} &= \frac{m}{m_0}\end{aligned}$$

Next we rescale distance with an appropriate length scale  $L$  and time with  $\tau = \frac{L^2}{D}$  [2], which will be described more detailed in section 4.

Modifying the system's free parameters  $D_c, \chi, \delta, D_m, \mu_3, \lambda$  gives us:

$$d_c = \frac{D_c}{D}, \quad \gamma = \chi \frac{e_0}{D}, \quad \eta = \tau m_0 \delta, \quad d_m = \frac{D_m}{D}, \quad \alpha = \tau \mu_3 \frac{c_0}{m_0}, \quad \beta = \tau \lambda.$$

with  $D$  being a reference chemical diffusion coefficient.

These modifications make the new system of coupled partial differential equations, where henceforth the tildes are dropped and we assume  $t$  as  $\tau$  for simplicities' sake:

$$\frac{\partial c}{\partial t} = d_c \Delta c - \gamma \nabla \cdot (c \nabla e) + \mu_1 c (1 - c - e) \quad (6)$$

$$\frac{\partial e}{\partial t} = -\eta m e + \mu_2 e (1 - c - e) \quad (7)$$

$$\frac{\partial m}{\partial t} = d_m \Delta c + \alpha c - \beta m \quad (8)$$

with also updated zero-flux boundary conditions:

$$\zeta \cdot (-d_c \nabla c + c \gamma \nabla e) = 0 \quad (9)$$

$$\zeta \cdot (-d_m \nabla m) = 0 \quad (10)$$

where the free parameters are as described above and  $\zeta$  stays an appropriate outward unit normal vector.

In order to use the Finite Element Method, we will change to the variational formulation. If we assume each species to be in the Hilbert space  $H^1(\Omega)$ , the variational formulation can be derived by multiplying with a test function  $\varphi_j, j \in \{c, e, m\}$ , integrating over the domain  $\Omega$  and using integration by parts and the Gauss theorem. Using the variational formulation will give us a broader solution space and reduce the solution's requirements regarding differentiability. With  $(\cdot, \cdot)$  denoting the  $L^2$ -scalar product on  $\Omega$  the following equation system results:

$$\left( \frac{\partial c}{\partial t}, \varphi_c \right) = -d_c (\nabla c, \nabla \varphi_c) + \gamma (c \nabla e, \nabla \varphi_c) + \mu_1 \left( c \left( 1 - \frac{c}{c_0} - \frac{e}{e_0} \right), \varphi_c \right) \quad (11)$$

$$\left( \frac{\partial e}{\partial t}, \varphi_e \right) = -\eta (m e, \varphi_e) + \mu_2 \left( e \left( 1 - \frac{c}{c_0} - \frac{e}{e_0} \right), \varphi_e \right) \quad (12)$$

$$\left( \frac{\partial m}{\partial t}, \varphi_m \right) = -d_m (\nabla m, \nabla \varphi_m) + \alpha (c, \varphi_m) - \beta (m, \varphi_m) \quad (13)$$

## 4 Experiments and Results

All experiments considering the ECM homogenous start with the same initial values, as seen in figure 3. Experiments observing the effects of a heterogenous ECM use different initial values, as seen in figure 4.

Solving the numerical model HiFlow<sup>3</sup>[9] will be used with the weak form given with equations 11 - 13. ParaView [10] is used to evaluate the numerical simulation results, producing informative plots to compare the evolution of the simulation in time. For this, we rely on the tool Plot Over Line to give results for the three variables of tumor cell density, extracellular matrix density and matrix-degrading enzyme concentration. This tool also allows us to compare all three variables in one plot, whereas using 2D plots, we would need one for each variable at a time. Using it makes the results better readable and allows a clearer quantitative insight into the experiments, as shown in figure 6.

This work starts with replicating numerical simulations done by other papers in higher dimensions. Since there were only 1D simulations done previously, the model will be adjusted so that the Plot Over Line graphs mimic the plots given by the previous experiments. This will serve two purposes: first, it will verify the correct implementation of the model and second, it will give us a starting point by which we can vary the parameters, investigating the phenomena this model exhibits.

We will start with examining 2D experiments with homogenous ECMS, using our model with the parameters  $\mu_1$  and  $\mu_2$  set to zero, considering a case with no proliferation of tumor cells and no renewal of the extracellular matrix molecules. After this, we will introduce proliferation and renewal, incorporating  $\mu_1$  and  $\mu_2$  into the variation. Due to the immense computational effort, we will do a slimmer replication 3D experiments and point out the effects of changing the dimension. After examining the model with homogenous ECM initial condition, we will move on to a case considering a heterogenous ECM in two spacial dimensions.

Looking at the parameter estimates from [2] to non-dimensionalise the time, we see that with  $L \in [0.1\text{cm}, 1\text{cm}]$  and  $D \approx 10^{-6}\frac{\text{cm}^2}{\text{s}}$ ,  $t = \frac{L^2}{D}$  would give us a big temporal range,  $t_{min} = 1000\text{s}$  and  $t_{max} = 1000000\text{s}$  in which the simulations take place. However, using estimates taken from [11] and [5] for the length scale, with  $L = 0.2\text{cm}$  gives us a concrete value for our non-dimensional time,  $t = 40000\text{s}$ .

Since Anderson et al. do not specify the exact value for  $L$ , we needed to determine appropriate configurations for  $dt$  used in our simulation to get comparable results. We used a timestep of  $dt = 0.01$ , corresponding to  $400\text{s}$  and let the simulations run for a dimensionless time of  $t = 8$  corresponding to  $320000\text{s} = 88\text{ days}$ . Looking at the results below, we found that for every unit of time in Anderson et al.'s experiments, in our experiments, 0.4 units of time have passed.

Another challenge are the diffusion and haptotaxis coefficients. Since they depend on the dimension we are in, we first have to find our own estimates as a baseline value. We will use these baseline values as a foundation for the Parameter Analysis. In the Parameter Analysis we will vary one or more, at most 3 parameters at time to get an overview of their effects.

For all the plots of the experiments, the red curve indicates the tumor cell density, the

blue curve the ECM concentration and the green curve the MDE concentration.

Before starting with the experiments we need to discuss the system's initial conditions. We state them for three dimensions, but for two or one dimension the  $y$  or  $z$  terms need to be left out.

We assume for that at dimensionless time  $t = 0$ , there is already a nodule of cells located at the center of the unit domain  $\Omega$  that has produced a concentration of matrix-degrading enzymes and has already degraded the extracellular matrix at the center. We therefore define the initial conditions for the tumor cell density as:

$$c(x, y, z, 0) = \exp\left(-\frac{(x - 0.5)^2 + (y - 0.5)^2 + (z - 0.5)^2}{0.01}\right)$$

and the initial conditions of the matrix-degrading enzymes are described as:

$$m(x, y, z, 0) = 0.5 \exp\left(-\frac{(x - 0.5)^2 + (y - 0.5)^2 + (z - 0.5)^2}{0.01}\right)$$

Regarding the structure of the extracellular matrix we are using two cases; first a homogenous ECM strucutre and second a heterogenous ECM structure. The strucutre of the homogenous extracellular matrix looks as follow:

$$e(x, y, z, 0) = 1 - 0.5 \exp\left(-\frac{(x - 0.5)^2 + (y - 0.5)^2 + (z - 0.5)^2}{0.01}\right)$$

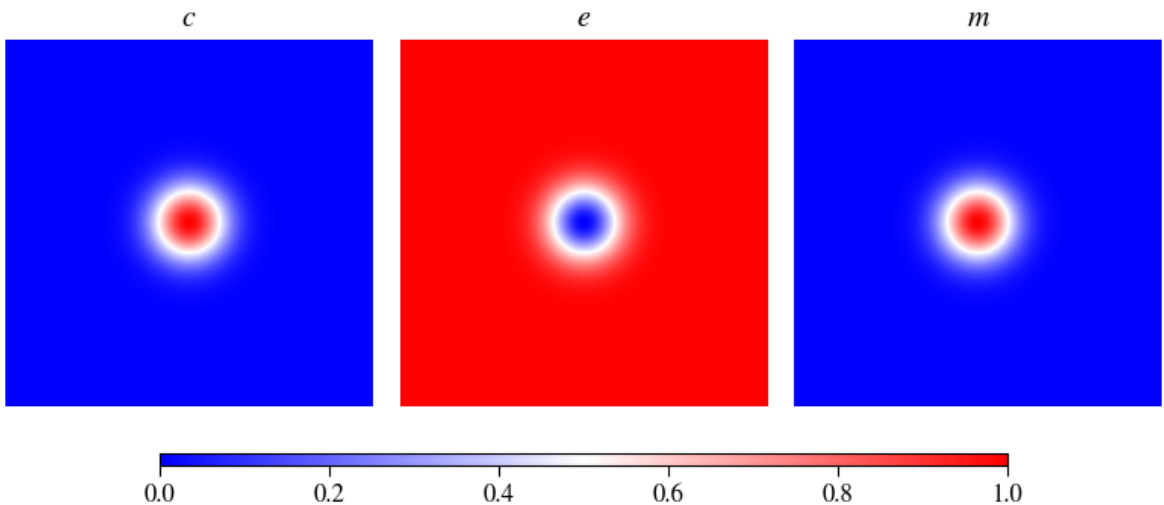


FIGURE 3. Visualization of the initial value distribution for an experiment in two space dimensions with a homogenous extracellular matrix

Figure 3 shows a 2D plot of the initial conditions using the homogenous extracellular matrix structure. The images describing the tumor cell density and the matrix-degrading

enzyme concentration have high values in the center. The image describing the extracellular matrix has high values everywhere except the center since the matrix-degrading enzymes have already degraded the ECM.

The other case we are investigating is using a heterogenous structured extracellular matrix. In section ??, we investigate how a more realistic structure of the ECM will affect the results of a 2D simulation. Since we are looking at a heterogenous extracellular matrix only for in experiments with two spacial dimensions we are stating the initial conditions one the variable  $e$  also for two spatial dimensions. For this scenario, we also assume a nodule of tumor cells in the center of the simulation that has produced a concentration of matrix-degrading enzymes. Contrary to the homogenous ECM strucutre, we assume that the tumor cells are located at a basement membrane that they already have invaded and are now pushing into the extracellular matrix behind it, located on the left side looking from the center. These assumptions are described in figure ?? . While the initial conditions on tumor cell density and matrix-degrading enzyme concentration are the same for both cases of ECM strucutre, we have to change the initial values of the variable  $e$  to the following:

$$e(x, y, 0) = (1 - 0.5 \exp\left(-\frac{(x - 0.5)^2 + (y - 0.5)^2 + (z - 0.5)^2}{0.01}\right)) \frac{1}{1 + \exp\left(-\frac{2}{0.1(x-0.5)}\right)}$$

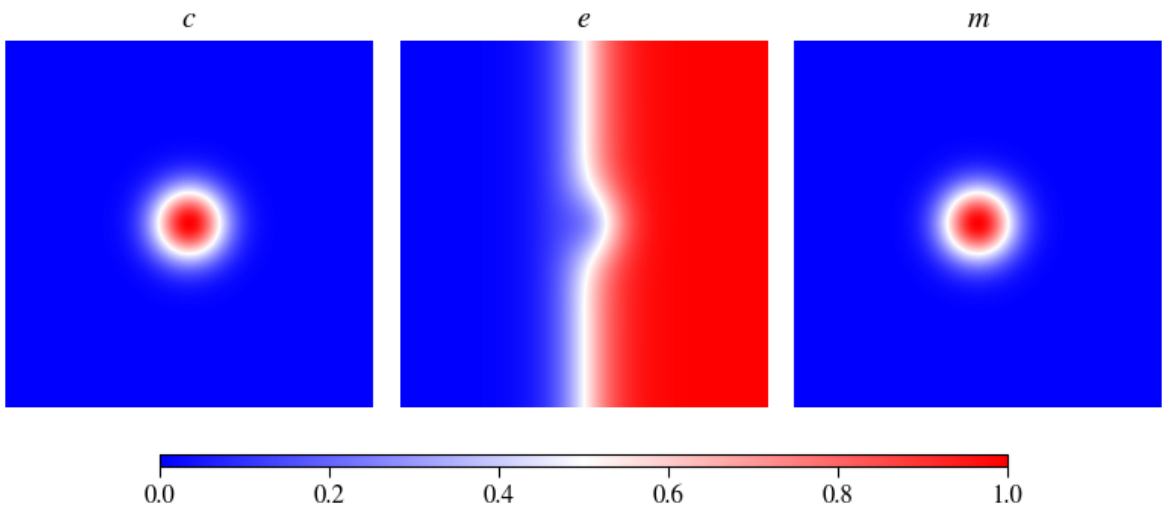


FIGURE 4. Visualization of the initial value distribution for an experiment in two space dimensions with a heterogenous extracellular matrix

Figure 4 describes these modifications on  $e$  and shows the inital conditions for our system using a heterogenous ECM structure.

#### 4.1 Dimension Variation

Before starting with the Parameter Analysis, we need to discuss how changing the dimension varies the results and what adjustments we need to take, to make our model mimick the one given by Anderson et al. Therefore we start with using the parameters given by Anderson et al.'s first experiment [2]. Figure 5 shows a screenshot of this experiment, unfortunately in low resolution, since the original paper had not included any digital data containing the diagrams of their results. In the figure, you can see that after  $t = 1$  in

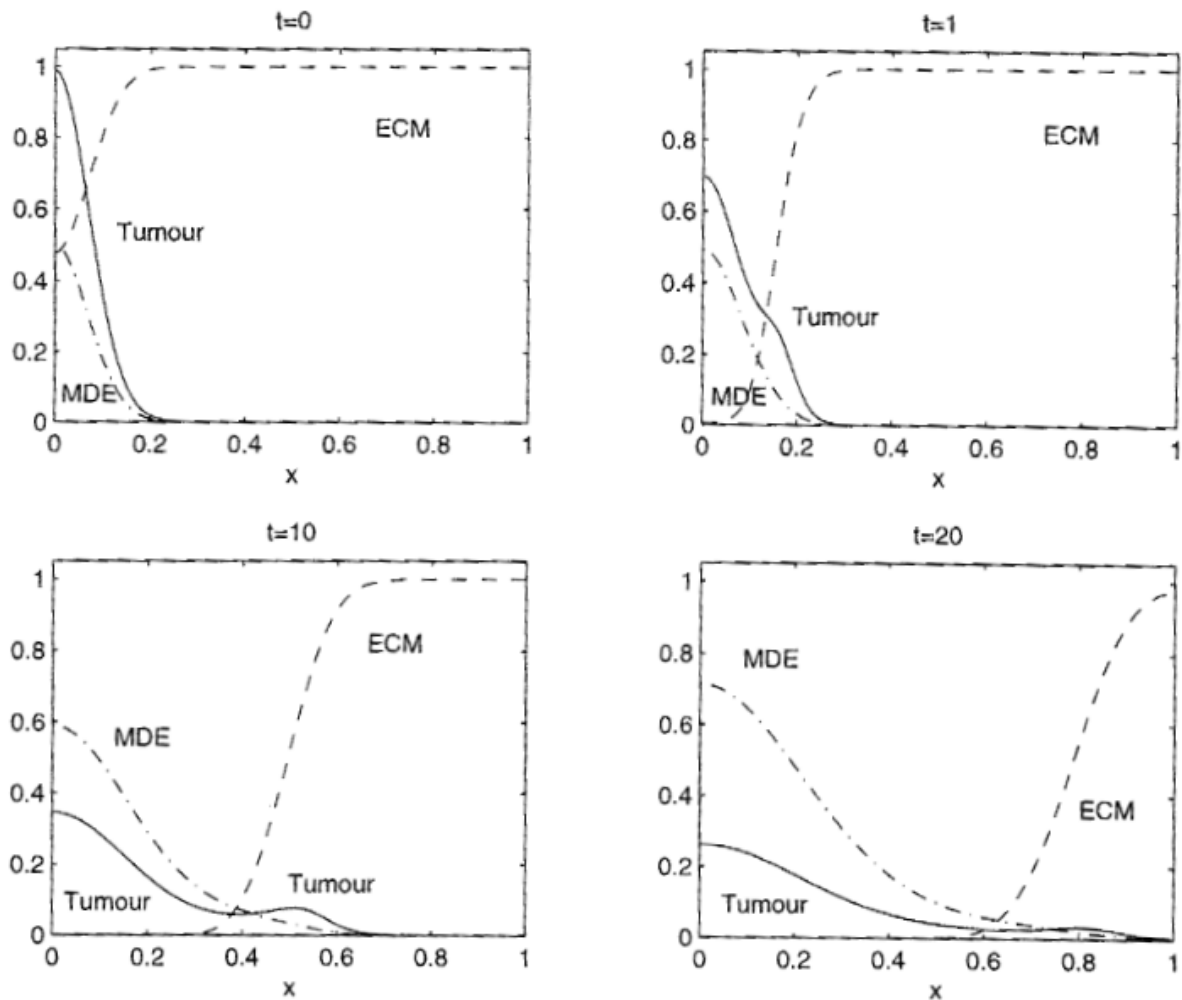


FIGURE 5. Anderson's first one dimensional experiment using the parameter values  $d_c = 1 \cdot 10^{-3}$ ,  $d_m = 1 \cdot 10^{-3}$ ,  $\gamma = 0.005$ ,  $\eta = 10$ ,  $\alpha = 0.1$ ,  $\beta = 0$ ,  $\mu_1 = 0$ ,  $\mu_2 = 0$

their timescale, the tumor cells develop a division at the part invading the tissue. This division propagates to a pointy peak at a later point in time. The concentration of the matrix-degrading enzymes increases continuously and the concentration of the extracellular matrix decreases continuously. Their  $x$ -axis is stretched or rescaled to show an interval from 0 to 1. In our case, the interval on the plots has only  $x$ -values from 0 to 0.5.

Replicating this experiment in two and three dimensions, we start with the same

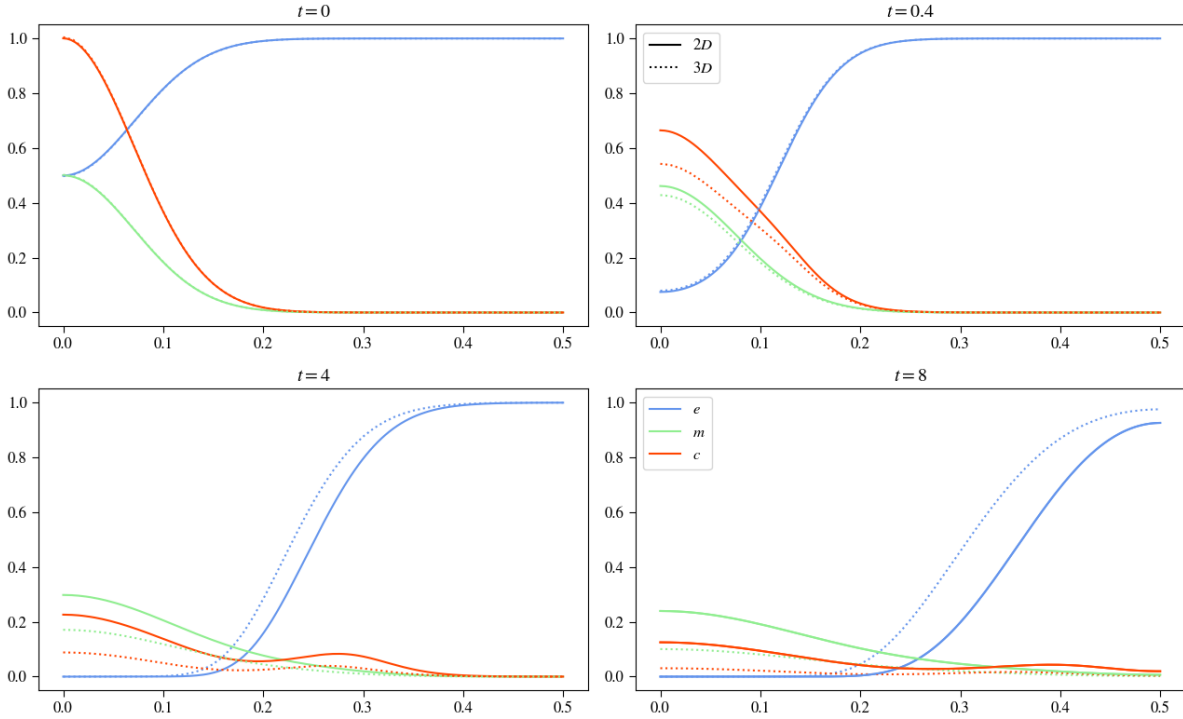


FIGURE 6. Results using Anderson et al.’s parameters produced by applying the Plot Over Line tool in two and three dimensions

parameters as Anderson et al. had used,  $d_c = 1 \cdot 10^{-3}$ ,  $d_m = 1 \cdot 10^{-3}$ ,  $\gamma = 0.005$ ,  $\eta = 10$ ,  $\alpha = 0.1$ ,  $\beta = 0$ . Figure 6 and adjust them to make our simulation match Anderson et al.’s as good as possible.

Starting from the initial values, we see that after  $t = 0.4$ , a very small division of the tumor cells is starting to form for both dimensions. This division increases strongly for the experiment done in two dimensions and minimal for the experiment done in three dimensions at  $t = 4$ . At  $t = 8$ , both curves visibly flatten. We see that the division of tumor cells detaching from the primary tumor are not as pregnant as in the one-dimensional experiment of Anderson et al. The interplay of the diffusive and haptotactic factors determines how considerable this division will be that detaches from the primary lump of the tumor cells and invades the tissue faster. It will also decide how pointy this secondary lump of tumor cells will be. Biologically, this relation between diffusion and haptotaxis translates to the invasion pace of the tumor cells, the rate at which how much of them will remain at the primary tumor and how much will invade further into the tissue degrading the extracellular matrix.

Looking at the concentration of matrix-degrading enzymes, we see it is visibly lower here than in Anderson et al.’s experiment. We see little increase over time. We can change this by changing the production factor of  $\alpha$  or by adjusting the motility of the tumor cells producing them exponentially. The factor  $\alpha$  determines how fast the tumor cells produce the matrix-degrading enzymes, degrading the extracellular matrix and allowing the tumor cells to invade the tissue further.

Only the extracellular matrix concentration seems to mimic the behavior of Anderson et al.'s experiment. It decreases continuously and as the last image shows, there is still a considerable amount left.

To replicate Anderson et al.'s results, we will start adjusting the parameters mentioned exemplary in two spacial dimensions: the tumor cells' diffusion and haptotaxis coefficients and the production rate of the matrix-degrading enzymes. We are focusing on the values the variables will take at the origin  $x = 0$ . Since Anderson et al. used in the original paper the model without tumor cell proliferation and extracellular matrix renewal, we are using the same model during the dimensional variation.

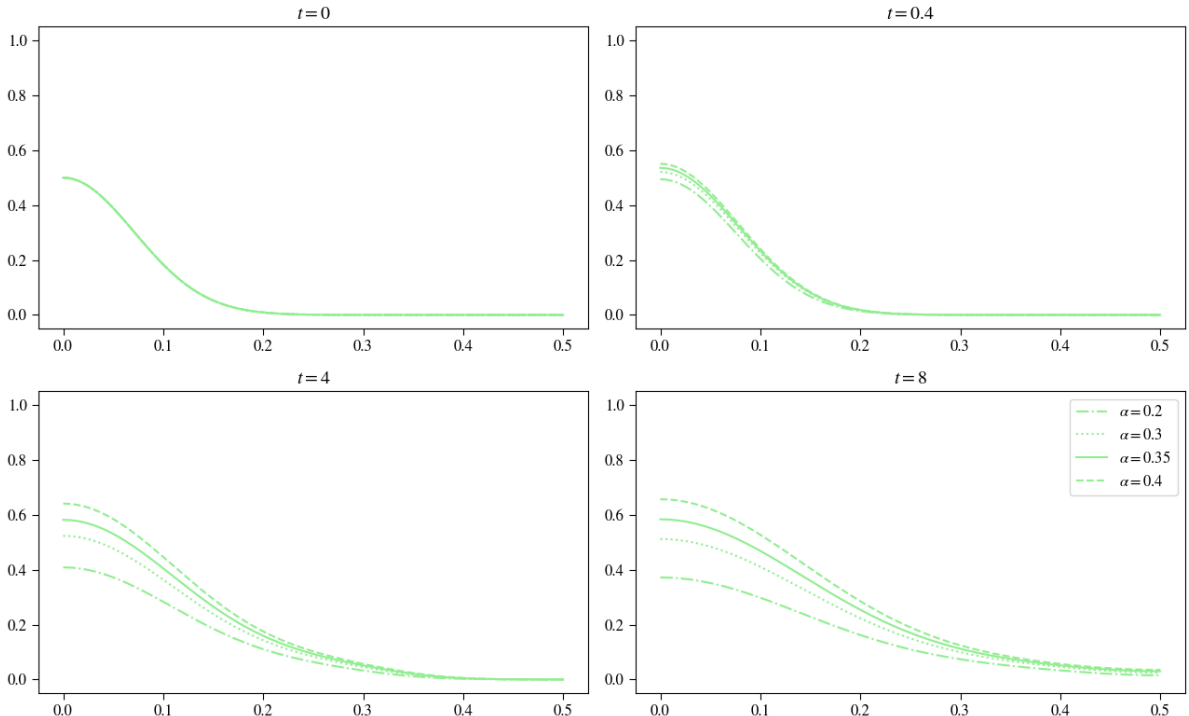


FIGURE 7. Comparison of  $\alpha$  values to replicate Anderson et al.'s experiment

Starting with comparing different values for  $\alpha$ , we see in figure 7 a comparison of how this affects the curve for the matrix-degrading enzyme concentration. A maximum difference between the compared values of 0.2 already causes drastic changes. In Anderson et al.'s experiment, we observed a value of approximately  $m(0, 1) = 0.5$  after, in their timescale,  $t = 1$  at the origin. This value is mimicked best for a value of  $\alpha = 0.2$  in our experiments. However, in the later points in time, we see that this value for  $\alpha$  is insufficiently low in terms of increase, though choosing  $\alpha$  higher than  $\alpha = 0.4$  results in an accelerated ECM degradation that is too fast. Taking a value between those two allowed our simulations to exhibit the observed behavior. We choose a value of  $\alpha = 0.35645$  to use in the later experiments as a basecase.

Looking at the diffusion coefficient for the tumor cells, in figure 8, you can see different  $d_c$  values regarding the tumor cell concentration. It is clear that with decreasing  $d_c$ , the sharpness of the secondary lump of cells that begins to invade the tissue drastically

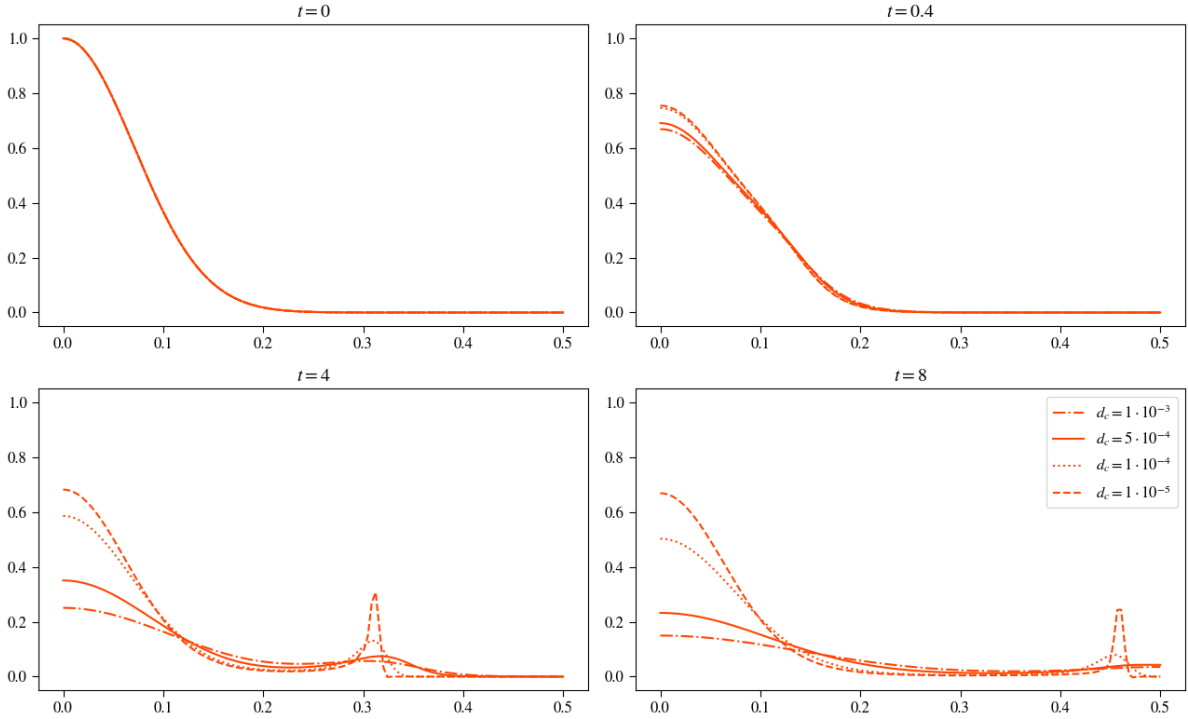


FIGURE 8. Comparison of  $d_c$  values to replicate Anderson et al.’s experiment

increases due to increased effects of haptotaxis, which is now the main factor in controlling tumor motility. However, we also observe that with decreasing  $d_c$ , the remaining lump of tumor cells at the origin increases due to little diffusion here. Over time, we found that  $d_c = 5 \cdot 10^{-4}$  describes Anderson et al.’s experimental results best for our simulations, with a roughly matching density of tumor cells that remains at the origin at all times and, for our case, balanced effects of haptotaxis and diffusion.

At last, we want to adjust the haptotactic pull slightly; for this, we are looking at a comparison of different  $\gamma$  values in figure 9 depicting the tumor cell density. Comparing our results with Anderson et al.’s, we want a bigger secondary lump that invades the tumor cells faster than the one remaining at the origin. We need to increase  $\gamma$  to increase the haptotactic pull that controls tumor cell motility. The figure shows that increasing  $\gamma$  yields these effects, but increasing it too much results in a too-fast invasion pace of the surrounding tissue. Increasing  $\gamma$  slightly to  $\gamma = 0.0055$  is sufficient to produce the desired effects.

These adjustments leave us with the final configuration for replicating the system in two dimensions. We see the important effects met when comparing the two dimensional experiment with the original experiment. We see that the production of the matrix-degrading enzymes fits the original experiment and the motility of the tumor cells also matches, with balanced effects of haptotaxis and diffusion. However comparing the three-dimensional experiment to original, there are still major differences. The second timestep  $t = 0.4$  shows a tumor cell density that is considerably lower in the 3D experiment than in the 2D experiment, which is can also be observed at the later points in time. Due to

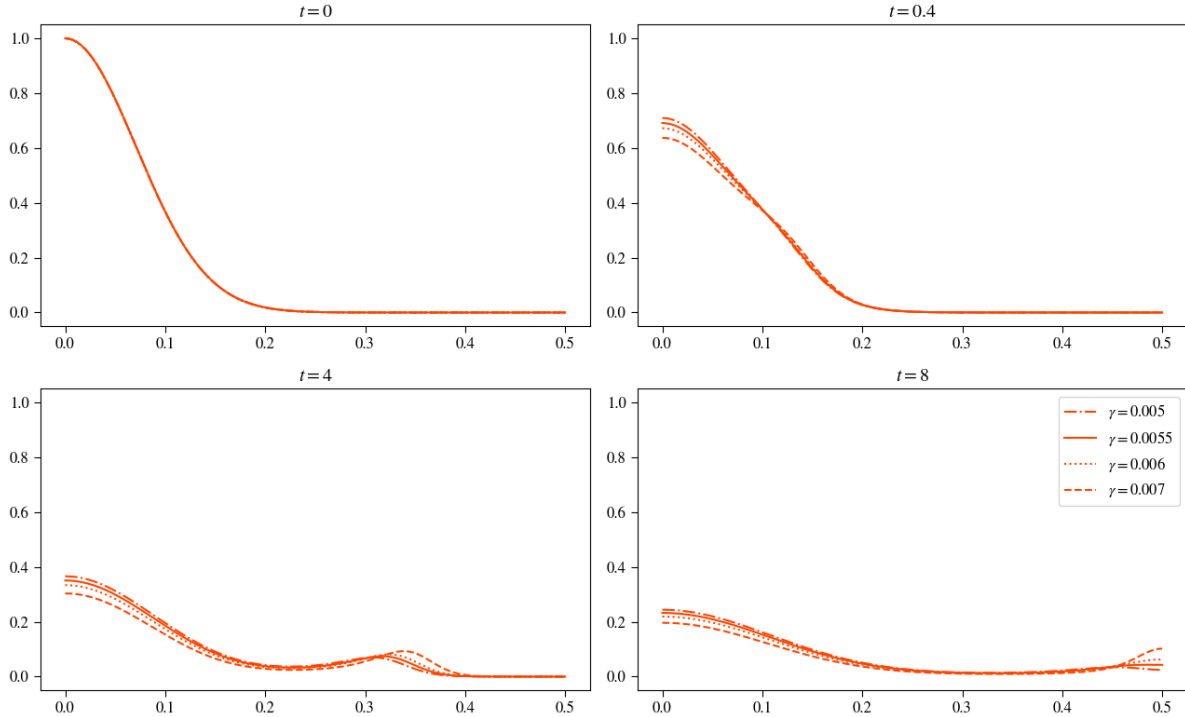
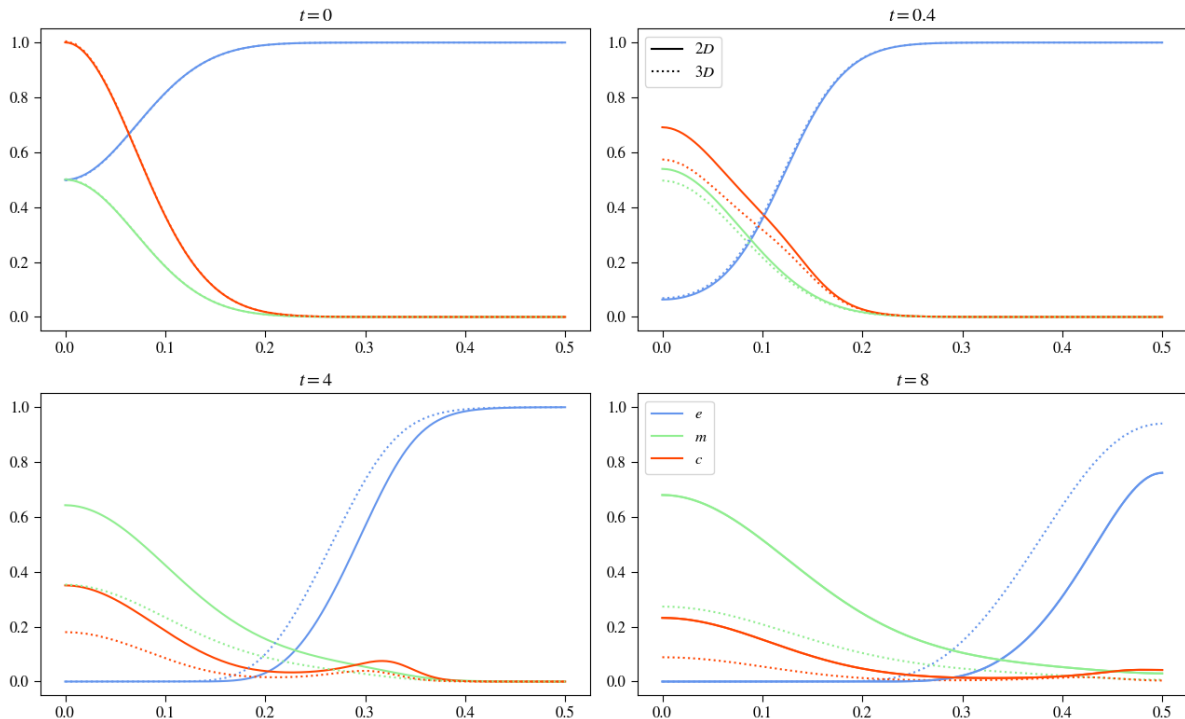
FIGURE 9. Comparison of  $\gamma$  values to replicate Anderson et al.'s experiment

FIGURE 10. 3D Replication of using the 2D basecases values

the exponential production of matrix-degrading enzymes by the tumor cells, the change in tumor cell density also effects the MDE concentration massively, resulting also in a lower concentration. The also reduced MDE concentration causes a slower degradation process of the extracellular matrix.

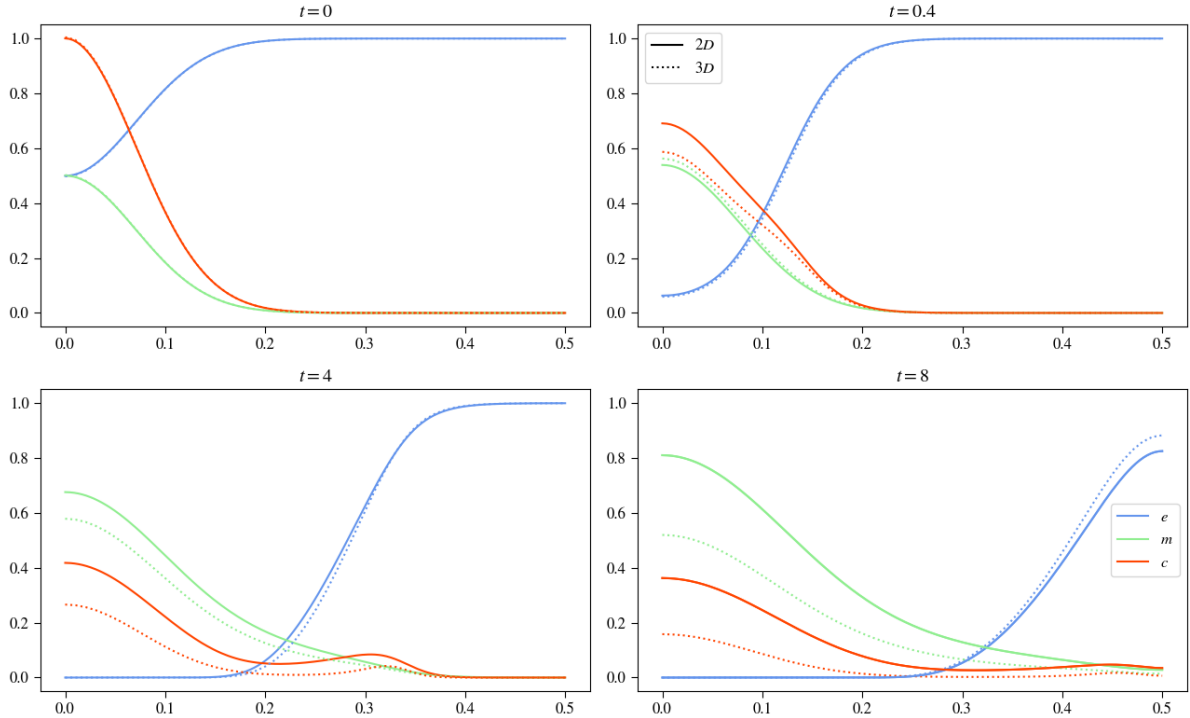


FIGURE 11. Comparing 2D basecase without proliferation and 3D basecase

In figure 11, we used the same schematic we did above, changing  $d_c, \gamma$  and  $\alpha$  to make the 3D results simulations match the results produced in 2D or 1D. Challenging in this process was, as mentioned previously, that the reduced tumor cell density causes drastic changes in both MDE production and ECM degradation. Settling on the values:  $d_c = 3 \cdot 10^{-4}$ ,  $\gamma = 0.0058$ ,  $\alpha = 0.6$ , yields sufficiently matching results.

Now the question arises why we have to change those values to replicate results. We can argue here with the parameter analysis, but also with an algebraic hypothesis.

Since  $\beta$  is assumed for most experiments to be zero, also in replication one-dimensional results in two dimensions, we also set it to zero in this case.

Looking at the parameter analysis for  $d_m$  we saw that this parameter has little influence compared to other on the simulation, excepting the overall MDE concentration in certain regions, which is neglectable when in those regions the ECM is already fully degraded. Also regarding the spacial occupation and distribution of the matrix-degrading enzymes we saw that a more important factor is the motility of the tumor cells producing them in their wake.

Inspecting the term concerning  $\eta$  we see that it depends on local values for the ECM and MDE concentrations;  $-\eta m(x, y, z, t)e(x, y, z, t)$ . In the experiments regarding replicating we see that we can influence the MDE concentration well with adjusting the motility

of the tumor cells and their production of the MDEs. In figure 11 you see that the concentration of the matrix-degrading enzymes roughly matches over time for both dimensions. In the plots  $m(x, y, z, t)e(x, y, z, t)$  therefore also roughly match, resulting in similar ECM degradation rate. Because of this, we also do not change  $\eta$ .

As mentioned above the tumor cell motility plays a crucial role during temporal development of the whole system. It is therefore a key component to match these curves. The motility is governed by diffusion and haptotaxis:

$$d_c \Delta c - \gamma \nabla \cdot (c \nabla e) = d_c \Delta c - \gamma (\nabla c \cdot \nabla e + \Delta e)$$

	1D	2D	3D
Diffusion	$\frac{\partial^2 c}{\partial x^2}$	$\frac{\partial^2 c}{\partial x^2} + \frac{\partial^2 c}{\partial y^2}$	$\frac{\partial^2 c}{\partial x^2} + \frac{\partial^2 c}{\partial y^2} + \frac{\partial^2 c}{\partial z^2}$
Haptotaxis	$\frac{\partial c}{\partial x} \frac{\partial e}{\partial x} + \frac{\partial^2 e}{\partial x^2}$	$\frac{\partial c}{\partial x} \frac{\partial e}{\partial x} + \frac{\partial c}{\partial y} \frac{\partial e}{\partial y} + \frac{\partial^2 e}{\partial x^2} + \frac{\partial^2 e}{\partial y^2}$	$\frac{\partial c}{\partial x} \frac{\partial e}{\partial x} + \frac{\partial c}{\partial y} \frac{\partial e}{\partial y} + \frac{\partial c}{\partial z} \frac{\partial e}{\partial z} + \frac{\partial^2 e}{\partial x^2} + \frac{\partial^2 e}{\partial y^2} + \frac{\partial^2 e}{\partial z^2}$

TABLE 1. Overview of Diffusion and Haptotaxis terms across varying dimensions

Table 1 gives an overview of the diffusion and haptotaxis terms for the treated dimensions in this work. For the diffusion we see that with increasing the dimension we get one additional term, assuming for simplicity  $\frac{\partial^2 c}{\partial x^2} = \frac{\partial^2 c}{\partial y^2} = \frac{\partial^2 c}{\partial z^2} = \kappa(t)$ . Also assuming a constant factor  $M$  for diffusion, this yields in one spacial dimension  $M = d_{c,1}\kappa(t)$ , equivalent to  $d_{c,1} = \frac{M}{\kappa(t)} = 1 \cdot 10^{-3}$ ; in two spacial dimensions  $d_{c,2} = \frac{M}{2\kappa(t)} = \frac{1}{2}d_{c,1} = \frac{1}{2}10^{-3} = 5 \cdot 10^{-4}$  and in three dimensions  $d_{c,3} = \frac{M}{3\kappa(t)} = \frac{1}{3}d_{c,1} = \frac{1}{3}10^{-3} = 3.33 \cdot 10^{-4}$ .

If we applied the same hypothesis for  $\gamma$ ; assuming a constant factor  $H$  for it and assuming  $\frac{\partial c}{\partial x} \frac{\partial e}{\partial x} = \frac{\partial c}{\partial y} \frac{\partial e}{\partial y} = \frac{\partial c}{\partial z} \frac{\partial e}{\partial z} = \iota$ ,  $\frac{\partial^2 e}{\partial x^2} = \frac{\partial^2 e}{\partial y^2} = \frac{\partial^2 e}{\partial z^2} = \theta$ , we would get in one dimension:  $\gamma_1 = \frac{H}{\iota + \kappa(t)} = 0.005$ , in two:  $\gamma_2 = \frac{H}{2\iota + 2\kappa(t)} = \frac{1}{2}\gamma_1 = 0.0025$  and in three dimensions:  $\gamma_3 = \frac{H}{3\iota + 3\kappa(t)} = \frac{1}{3}\gamma_1 = 0.00166$ . We see from experimental results that using  $\gamma = 0.002$  and  $d_c = 5 \cdot 10^{-4}$  yields a curve for the tumor cells that has no secondary lump of cells detaching from the primary tumor. Adjusting  $\gamma$  we cannot employ the same hypothesis, since the equation describing haptotaxis also incorporates a term regarding the gradients of both curves;  $\nabla c \cdot \nabla e$  not only the laplacian of the extracellular matrix concentration  $\Delta e$ . Depending on the point in space and time we look at  $\nabla c(x, y, z, t) \cdot \nabla e(x, y, z, t)$ , this term can damping increasing or no effect at all. We therefore resort to determining this value experimentally with a value of  $\gamma = 0.0057$ .

Further improvements to make the experiments match better could be made on varying  $\mu_1$  and  $\mu_2$  to either match the tumor cell density more accurately or the degradation rate of the extracellular matrix.

The conclusions we can draw from adjusting the parameters with varying the dimensions are that the most important factor is the tumor cell density. The motility is crucial for the as well the shape of the MDE curve as well as the concentration of it, due to the tumor cells producing them exponentially, but this we can also adjust with the production coefficient  $\alpha$ . Diffusion coefficient can be calculated algebraically, as seen above, but for Haptotaxis the coefficient must be determined experimentally. It is to say that

due though it is possible to make the simulations across dimensions match at one point in time, it is due to the many different parameters and terms impossible to make them perfectly fit over time.

Figure	Linestyle	$d_c$	$\gamma$	$\eta$	$d_m$	$\alpha$	$\beta$
6	—	$1 \cdot 10^{-3}$	0.005	10	$1 \cdot 10^{-3}$	0.1	0
7	—	$1 \cdot 10^{-3}$	0.005	10	$1 \cdot 10^{-3}$	0.2	0
7	...	$1 \cdot 10^{-3}$	0.005	10	$1 \cdot 10^{-3}$	0.3	0
7	—	$1 \cdot 10^{-3}$	0.005	10	$1 \cdot 10^{-3}$	0.35	0
7	- - -	$1 \cdot 10^{-3}$	0.005	10	$1 \cdot 10^{-3}$	0.4	0
8	—	$1 \cdot 10^{-3}$	0.005	10	$1 \cdot 10^{-3}$	0.3546	0
8	...	$1 \cdot 10^{-4}$	0.005	10	$1 \cdot 10^{-3}$	0.3546	0
8	—	$5 \cdot 10^{-4}$	0.005	10	$1 \cdot 10^{-3}$	0.3546	0
8	- - -	$1 \cdot 10^{-5}$	0.005	10	$1 \cdot 10^{-3}$	0.3546	0
9	—	$5 \cdot 10^{-5}$	0.005	10	$1 \cdot 10^{-3}$	0.3546	0
9	...	$5 \cdot 10^{-4}$	0.0055	10	$1 \cdot 10^{-3}$	0.3546	0
9	—	$5 \cdot 10^{-4}$	0.006	10	$1 \cdot 10^{-3}$	0.3546	0
9	- - -	$5 \cdot 10^{-4}$	0.007	10	$1 \cdot 10^{-3}$	0.3546	0
10	—	$5 \cdot 10^{-4}$	0.0055	10	$1 \cdot 10^{-3}$	0.3546	0
10	...	$5 \cdot 10^{-4}$	0.0055	10	$1 \cdot 10^{-3}$	0.3546	0
11	—	$5 \cdot 10^{-4}$	0.0055	10	$1 \cdot 10^{-3}$	0.3546	0
11	—	$3.3 \cdot 10^{-4}$	0.0058	10	$1 \cdot 10^{-3}$	0.6	0

TABLE 2. Overview of all experiments conducted for the model without proliferation and renewal producing 2D output

In table 2 all experiments with their corresponding parameter configurations and linestyles in the respective figures are listed.

## 4.2 Parameter Analysis without Proliferation and Renewal - Homogenous ECM

This section describes the Parameter Analysis done in two spacial dimensions on a homogenous extracellular matrix structure. In this section we are using the basecase for established in the previous section with the parameters as used in 10.

Figure	Linestyle	$d_c$	$\gamma$	$\eta$	$d_m$	$\alpha$	$\beta$
12	—	$1 \cdot 10^{-4}$	0.0055	10	$1 \cdot 10^{-3}$	0.3546	0
12	...	$1 \cdot 10^{-3}$	0.0055	10	$1 \cdot 10^{-3}$	0.3546	0
13	...	$5 \cdot 10^{-4}$	0.002	10	$1 \cdot 10^{-3}$	0.3546	0
13	—	$5 \cdot 10^{-4}$	0.008	10	$1 \cdot 10^{-3}$	0.3546	0
13	- - -	$5 \cdot 10^{-4}$	0.01	10	$1 \cdot 10^{-3}$	0.3546	0

continued on next page

Figure	Linestyle	$d_c$	$\gamma$	$\eta$	$d_m$	$\alpha$	$\beta$
14	—	$5 \cdot 10^{-4}$	0.1	10	$1 \cdot 10^{-3}$	0.3546	0
15	....	$5 \cdot 10^{-4}$	0.0055	2	$1 \cdot 10^{-3}$	0.3546	0
15	—	$5 \cdot 10^{-4}$	0.0055	12	$1 \cdot 10^{-3}$	0.3546	0
15	....	$5 \cdot 10^{-4}$	0.0055	20	$1 \cdot 10^{-3}$	0.3546	0
16	....	$1 \cdot 10^{-3}$	0.0055	10	0.00001	0.3546	0
16	—	$1 \cdot 10^{-4}$	0.0055	10	0.001	0.3546	0
16	---	$1 \cdot 10^{-5}$	0.0055	10	0.1	0.3546	0
17	....	$5 \cdot 10^{-4}$	0.0055	10	$1 \cdot 10^{-3}$	0	0
17	—	$5 \cdot 10^{-4}$	0.0055	10	$1 \cdot 10^{-3}$	0.6	0
17	....	$5 \cdot 10^{-4}$	0.0055	10	$1 \cdot 10^{-3}$	1.0	0
18	....	$5 \cdot 10^{-4}$	0.0055	10	$1 \cdot 10^{-3}$	0.3546	0.1
18	—	$5 \cdot 10^{-4}$	0.0055	10	$1 \cdot 10^{-3}$	0.3546	0.01
18	....	$5 \cdot 10^{-4}$	0.0055	10	$1 \cdot 10^{-3}$	0.3546	0.005
19 - left	....	$5 \cdot 10^{-5}$	0.001	10	$1 \cdot 10^{-3}$	0.3546	0
19 - left	—	$5 \cdot 10^{-5}$	0.01	10	$1 \cdot 10^{-3}$	0.3546	0
19 - right	....	$1 \cdot 10^{-3}$	0.001	10	$1 \cdot 10^{-3}$	0.3546	0
19 - right	—	$1 \cdot 10^{-3}$	0.01	10	$1 \cdot 10^{-3}$	0.3546	0
20 - left	....	$5 \cdot 10^{-4}$	0.0055	2	$1 \cdot 10^{-5}$	0.3546	0
20 - left	—	$5 \cdot 10^{-4}$	0.0055	20	$1 \cdot 10^{-5}$	0.3546	0
20 - right	....	$5 \cdot 10^{-4}$	0.0055	2	$1 \cdot 10^{-3}$	0.3546	0
20 - right	—	$5 \cdot 10^{-4}$	0.0055	20	$1 \cdot 10^{-3}$	0.3546	0
21 - left	....	$5 \cdot 10^{-4}$	0.0055	10	$1 \cdot 10^{-3}$	0.1	0.005
21 - left	—	$5 \cdot 10^{-4}$	0.0055	10	$1 \cdot 10^{-3}$	0.1	0.1
21 - right	....	$5 \cdot 10^{-4}$	0.0055	10	$1 \cdot 10^{-3}$	1.0	0.005
21 - right	—	$5 \cdot 10^{-4}$	0.0055	10	$1 \cdot 10^{-3}$	1.0	0.1
22 - left	....	$5 \cdot 10^{-4}$	0.0055	10	$1 \cdot 10^{-5}$	0.1	0.005
22 - left	—	$5 \cdot 10^{-4}$	0.0055	10	$1 \cdot 10^{-5}$	0.1	0.1
22 - right	....	$5 \cdot 10^{-4}$	0.0055	10	$1 \cdot 10^{-5}$	1.0	0.005
22 - right	—	$5 \cdot 10^{-4}$	0.0055	10	$1 \cdot 10^{-5}$	1.0	0.1
23 - left	....	$5 \cdot 10^{-4}$	0.0055	10	$1 \cdot 10^{-3}$	0.1	0.005
23 - left	—	$5 \cdot 10^{-4}$	0.0055	10	$1 \cdot 10^{-3}$	0.1	0.1
23 - right	....	$5 \cdot 10^{-4}$	0.0055	10	$1 \cdot 10^{-3}$	1.0	0.005
23 - right	—	$5 \cdot 10^{-4}$	0.0055	10	$1 \cdot 10^{-3}$	1.0	0.1

TABLE 3. Overview of all experiments conducted for the model without proliferation and renewal producing 2D output

Table 3 gives a detailed overview of all the experiments done in this section and the parameters used to produce the results. We are considering the model without the proliferation of the tumor cells or renewal of the extracellular matrix. Therefore, parameters  $\mu_1$  and  $\mu_2$  are set to zero. In most figures, more than one experiment will be described; the linestyle in table 3 determines which experiment exactly is described by the set of

parameters.

#### 4.2.1 Parameter Analysis

Before we start with the parameter analysis, we will discuss the mathematical intuition concerning the system of partial differential equations, equations 6 to 10.

The equation governing the tumor cell density incorporates only two coefficients in this version regarding its motility, diffusion and haptotaxis. As mentioned during replicating Anderson et al.'s experiment, the relation between those two factors determines if a secondary lump of tumor cells detaches from the main lump and invades the tissue faster than the remaining cells but also how large this lump will be. We saw this behaviour in figure 8, varying  $d_c$  whilst keeping  $\gamma$  constant. Diffusive motility depends on the laplacian of the tumor cells themselves,  $\Delta c = \frac{\partial c}{\partial x} + \frac{\partial c}{\partial y} + \frac{\partial c}{\partial z}$ , which is a fundamental tool in sciences of all sorts to describe the effects of spacial rate of change of a scalar field quantity, in our case tumor cell density, at a specific point in space. Typically, this operator has high values where the respective quantity changes rapidly. We have a similar haptotaxis term:  $\nabla \cdot (c \nabla e) = \nabla c \cdot \nabla e + \Delta e$ . This relation means that haptotactic effects are strong not only where the concentration of the extracellular matrix changes rapidly but also where both gradients for tumor cell density and extracellular matrix assimilate in direction. However, these effects can also annihilate each other.

The equation describing the concentration of the extracellular matrix models its exponential decay, taking the concentration of the matrix-degrading enzymes into account. In spatial and temporal positions where both ECM and MDE concentrations are high, we can expect a fast degradation of the extracellular matrix.

The equation modeling the matrix-degrading enzymes combines motility with production/decay terms. The motility of the MDEs is modeled by the same diffusion as the tumor cells, mimicking their behavior in this regard. The tumor cells are responsible for producing the MDEs; the production is modeled by natural decay and both production and decay are modeled using exponential approaches.

From the replicated results shown in figure ??, we saw that the results can vary strongly if we vary the parameters one at a time. We will first look at how changing one parameter at a time affects the output of the simulation. After this, we will consider changing multiple parameters simultaneously in the cross-variation section. For varying the parameters, we assume the other parameters of the system are constant and use the baseline experiment as their values.

#### $d_c$ Variation

This parameter describes the diffusive properties of the tumor cells. As Chaplain et al. assumed in [11], we are also assuming an even distribution of this parameter with  $d_c \sim U[1 \cdot 10^{-5}, 1 \cdot 10^{-3}]$ . However, our experiments encountered numerical instabilities reducing the parameter further than  $5 \cdot 10^{-5}$ .

As described, the intuition is that decreasing  $d_c$  will increase the effects of haptotaxis and make  $\gamma$  more influential. This means the tumor cells will drift faster outward with a bigger secondary lump, forming a pointier leading edge. On the other hand, if we increase

$d_c$ , the effects of haptotaxis will diminish and the tumor cells will be subject to higher diffusion, distributing them more evenly in the tissue. Additionally, there will be less division from the primary lump of cells. Looking at the experiments in figure 12, we can

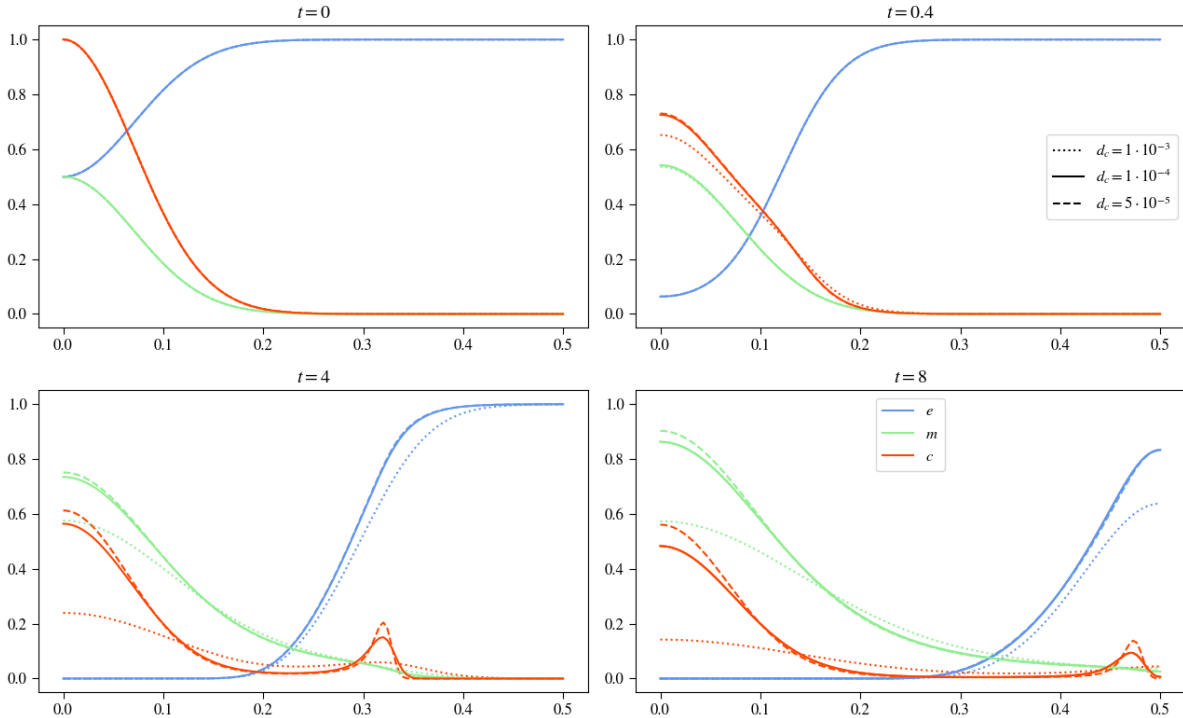


FIGURE 12. Results of varying  $d_c$  in the basecase parameter set, keeping the other parameters constant, using the Plot Over Line tool

see these assumptions met. The smaller  $d_c$  gets, the higher the influence of haptotaxis; therefore,  $\gamma$  will be and vice versa.

Considering the tumor cell density curves shown in red, we can see minor differences after already  $t = 0.4$ . For the two lower values of  $d_c$ , the solid and dashed curves, we see the tumor cells having a higher concentration at the origin than for the biggest value of  $d_c$ , dotted curve, nearly overlaying each other. The red dashed curve is considerably lower at the origin, though it is stretched out more than the other two, indicating a faster invasion rate. The other curves describing EDM and MDE concentration do not show any deviations from each other at this point.

Looking at the plot results, at time  $t = 4$ , we see the previously observed effects increase. The curves of the tumor cells confirm that with increasing  $d_c$ , the remaining lump of cells at the origin decreases, distributing the tumor cells more evenly in space and also reducing the effect of haptotaxis, making the secondary lump, which is still visible for the highest diffusion term, less sharp. At this point, the other two curves also show differentiating behavior. The ECM is degraded faster with increasing  $d_c$  and the slope the ECM describes is less steep. Looking at the extracellular matrix, we see minor differences between the lower two values for  $d_c$ . Due to the tumor cells' exponential production of the MDEs and the more even spread of the tumor cells, we see them taking on a lower

concentration at the origin. However, we can also observe that they have spread farther out than the MDE concentrations describing the experiments with lower  $d_c$  values.

Studying the last image of  $t = 8$ , we see no new effects, only the already mentioned propagated; with increasing  $d_c$ , we see a more even spread of the tumor cells and a reduction of the secondary lump, leading to the invasion of the tissue. For the MDE concentration, we observe less concentration in total due to the exponential growth rate, especially at the origin. However, we also see a more even distribution of them and a slightly faster invasion pace. This causes the extracellular matrix degradation to work faster.

Regarding the sensitivity of this parameter, the higher the value is, the more sensitive the system reacts. Though the lower two values for  $d_c$  are only separated by  $5 \cdot 10^{-5}$  and we can unfortunately not experiment with  $d_c = 1 \cdot 10^{-5}$  due to numerical instabilities, the differences between those two were minimal compared to the difference between the higher two values of  $d_c$ .

From a biological point of view, this increase in diffusion might be caused by a change in temperature, electric potential or mechanical pressure differences. The higher diffusion results in a more even spread of the two actively moving quantities of tumor cells and matrix-degrading enzymes, which degrade their surrounding tissue, the ECM, at a faster rate.

### $\gamma$ Variation

$\gamma$  describes the effects of haptotaxis; it is assumed that it is evenly distributed in  $\gamma \sim U[0.001, 0.01]$ , like Anderson et al. [2] assumed. Like Anderson et al., we will also look at values exceeding this region, though most likely losing their biological meaning. Inspecting the effects of  $\gamma$ , we can assume the countering effects on the tumor cells as for varying  $d_c$ ; selecting higher values for  $\gamma$  will increase the effects of haptotaxis, creating a larger secondary lump of cells that is being pulled faster into the tissue. The experiments described in figure 13 verify the expected behavior.

After  $t = 0.4$ , we already see apparent changes for varying  $\gamma$ ; the higher  $\gamma$  is set, the greater the division for the tumor cells is observable, which will later form the secondary lump. The lowest value for  $\gamma$ , undercutting the one for the basecase, shows no signs of a leading edge forming that invades the tissue separately. The two higher values for  $\gamma$  show little deviation. Considering the other two variables, extracellular matrix concentration and matrix-degrading enzyme concentration, no change is visible, still overlaying each other at this temporal point.

The following image shows the simulation after  $t = 4$  timesteps; we can see changes in all three variables. While the tumor cell density for the values of 0.008 and 0.01 differ slightly by the number of cells that are left at the origin and the distance they have already invaded the surrounding tissue, the curve for the lowest  $\gamma$  value, as at  $t = 0.4$  does not show a division of the tumor cells that invades the tissue separately, which causes the tumor cells to stay centered around  $x = 0$ , resulting in a higher density of cells there compared to the results of the other experiments. With increasing  $\gamma$ , the invasion speed also increases, as the dashed line for the tumor cell density shows. For the MDE curve, we also observe that the lower  $\gamma$  is, the more concentration is at the origin due to the higher

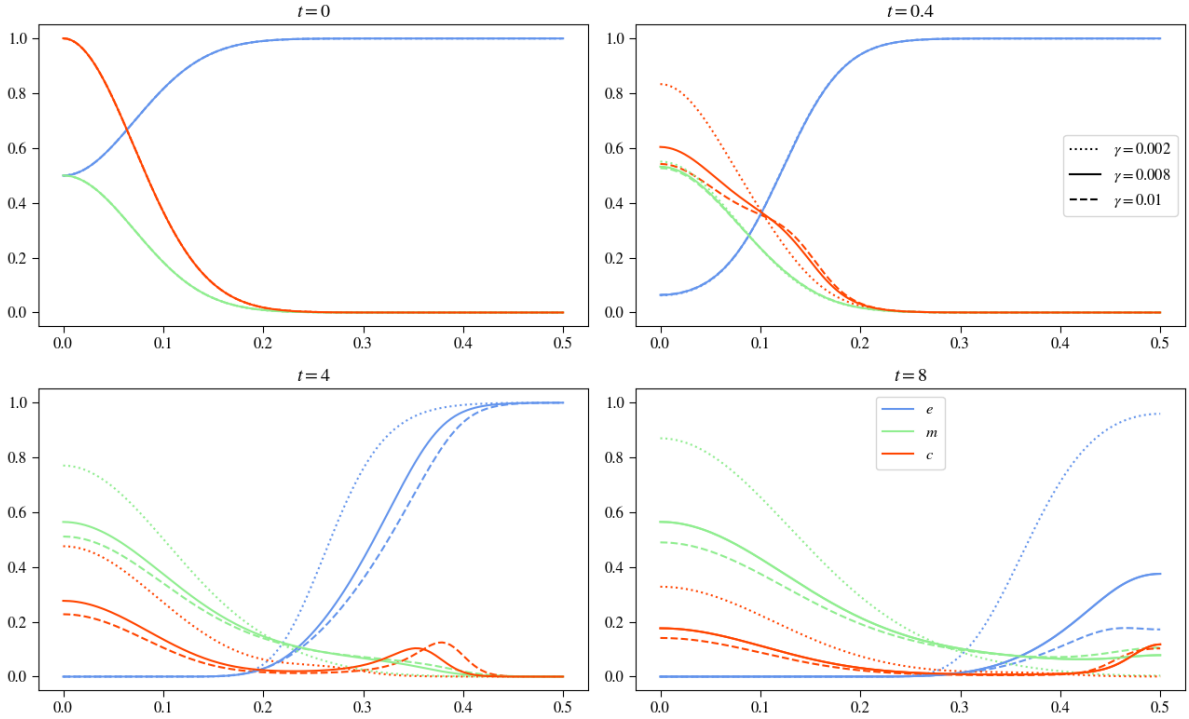


FIGURE 13. Results of varying  $\gamma$  in the basecase parameter set, keeping the other parameters constant, using the Plot Over Line tool

remaining density of tumor cells at the origin. The ECM concentration shows behavior similar to the MDE concentration; with increasing  $\gamma$ , the ECM is faster and more evenly degraded; due to the faster invasion of the tissue, the production of matrix-degrading enzymes also happens in regions farther away from the center. As we saw varying  $d_c$ , only a little MDE concentration is needed to degrade the extracellular matrix efficiently. Therefore, the ECM degradation process also happens faster here.

In the last image at  $t = 8$ , we see the observations from previous points in time confirmed; the higher  $\gamma$ , the faster the invasion pace of the tumor cells and the more of a division forms with fewer tumor cells staying at the origin. The behavior of the tumor cells causes a higher concentration of MDEs at the origin due to exponential growth and a steeper decline moving outwards. With rising  $\gamma$ , the ECM is getting degraded faster.

Out of curiosity, we will take a step further and increase  $\gamma$  again by one power to  $\gamma = 0.1$ . As previously observed, the haptotactic effects of pulling the cells into surrounding tissue increase, causing an even faster invasion pace. However, in this case, the invasion pace of the tumor cells is so high that no cells are left at the origin; everything is being pulled into the surrounding tissue. Before finishing the simulation after  $t = 8$ , the tumor cells have reached the domain  $\Omega$ 's border. The cells are being reflected at the border due to the boundary conditions of our model 9 and 10. In figure 14, you can see that after  $t = 2$ , the border is reached and at  $t = 3$ , the cells have been reflected to move into the corners, where the ECM concentration is highest. At  $t = 3$ , the pace of the ECM degradation of the matrix-degrading enzymes has not been able to keep up with

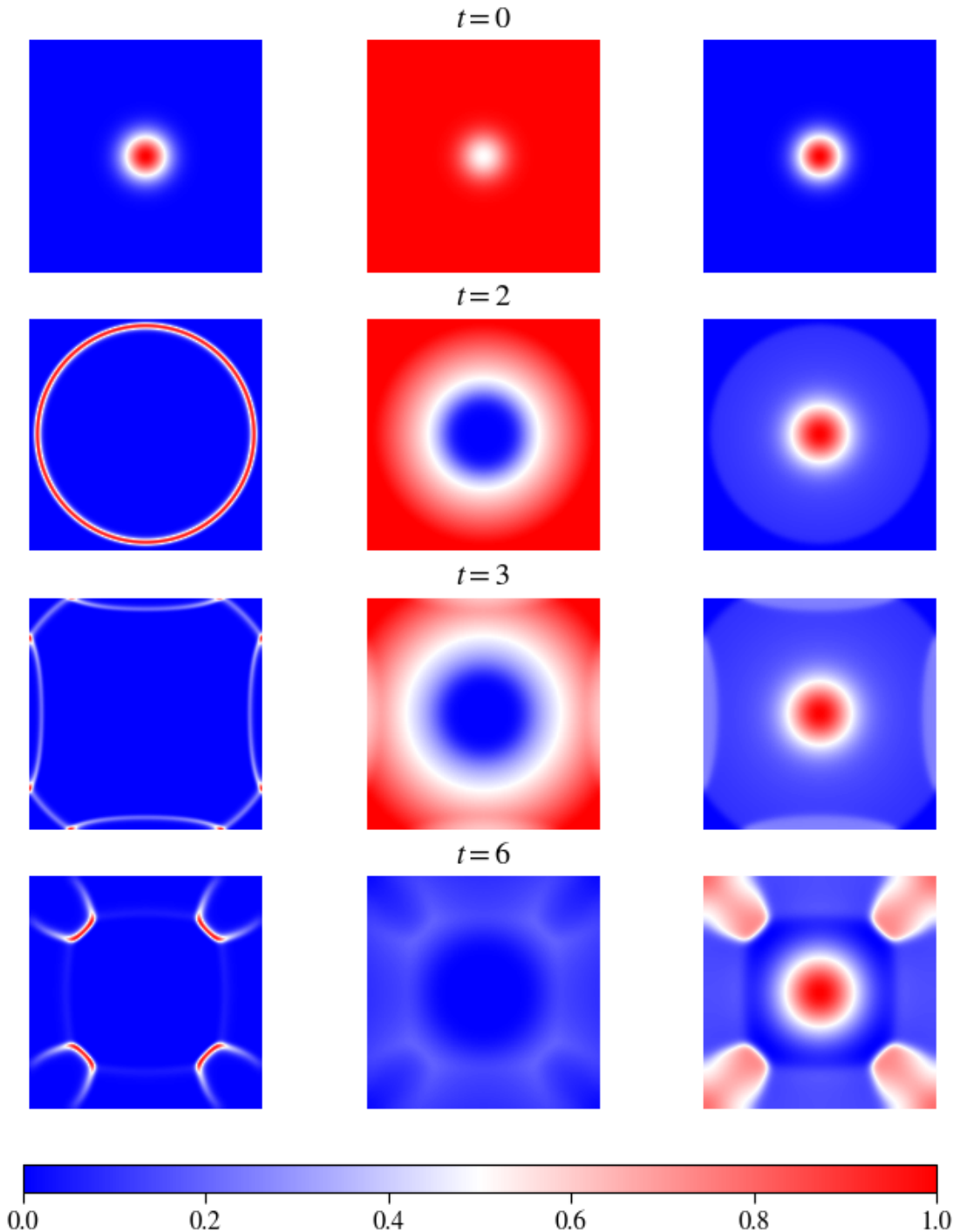


FIGURE 14. 2D plot showing experiment for  $\gamma = 0.1$ , left tumor cell density, right ECM concentration

the invasion pace of the tumor cells. After being pulled into the corners at  $t = 3$  and

degrading the ECM there, at  $t = 6$ , the tumor cells are being pulled back toward the center of the simulation.

Though this behavior matters little from a biological perspective due to the system's boundary conditions reflecting the movement and high invasion pace of the tumor cells, it is still interesting to investigate this case from a numerical perspective.

Though the intuition is met that with increasing  $\gamma$ , the invasion pace of the tumor cells and matrix-degrading enzymes also rises, we get unexpected behavior in the last experiment. There is more of a total ECM concentration left than in the experiment using  $\gamma = 0.01$ . Studying those experiments biologically, we know that the term extracellular matrix describes a whole class of different molecules, minerals or proteins, like collagens or enzymes. The different properties of these elements cause varying effects of haptotaxis. For example, were the haptotactic effects measured on laminin considerably lower than then ones measured on fibronectin, according to Aznavoorian et al. [12]. Different constellations of extracellular matrix composition will be encountered in different human body sites, which will cause the tumor cells, as seen in the numerical experiments, to behave differently.

### $\eta$ Variation

The parameter  $\eta$  controls the degradation process of the extracellular matrix molecules. Since Anderson et al. [2] used a value of  $\eta = 10$  on all their experiments, we assumed an even distribution in  $\eta \sim U[0, 20]$ . The degradation process of the extracellular matrix is modeled using an exponential decay hypothesis, so we can expect that increasing  $\eta$  also increases the system's sensitivity concerning the parameter  $\eta$ . With its role in controlling the degradation, it will also heavily influence the motility of the tumor cells and, therefore, the motility and production rate of the matrix-degrading enzymes. Slower degradation will result in a higher density of tumor cells at the origin, exponentially producing matrix-degrading enzymes. This, in turn, will increase the ECM degradation. With increasing  $\eta$ , the ECM is faster degraded and, therefore, might provoke a faster invasion rate of the tumor cells of the surrounding tissue. The higher  $\eta$  is, the fewer MDEs are needed to degrade the ECM efficiently.

Inspecting the results in figure 15, it is most striking that the slower degradation rate causes a slower tumor invasion.

We see significant differences for both curves, tumor cell density and ECM concentration after  $t = 4$ . Inspecting the experiment with  $\eta = 2$ , we see that the ECM has only degraded a little, exposing the tumor cells to stronger haptotactic pull by the ECM in regions closer to zero, compared to the other experiments. Though it may look like no secondary lump of cells is being formed, the contrary is the case; even more cells are pulled into the tissue, with the ECM slowly receding into the tissue, which smoothes out the bump the other two curves show for the tumor cell density. This results in only one lump of tumor cells that invades the tissue without one remaining at the origin. The same effect is observable when comparing the tumor cell density curves for the higher  $\eta$  values. With fewer tumor cells pulled by the ECM, the higher  $\eta$  gets. Only the curve for the matrix-degrading enzymes has not been affected by the variation of  $\eta$  until now.

The next image at  $t = 4$  propagates the effects on the tumor cells and the ECM. The

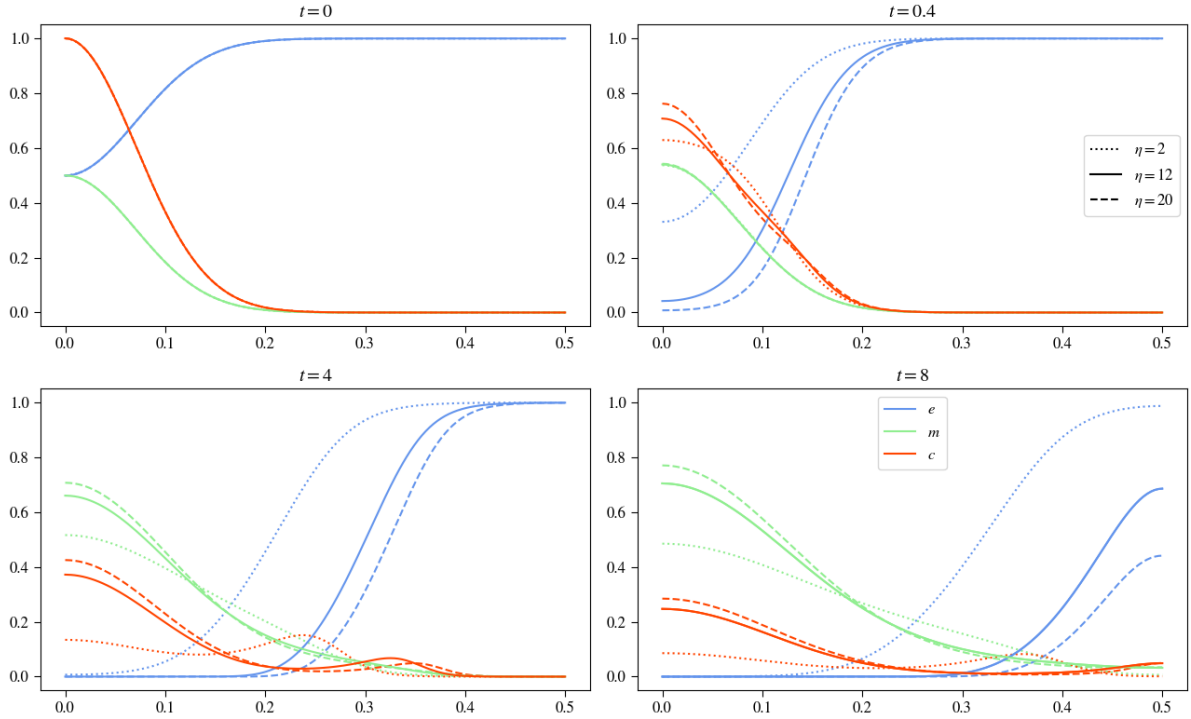


FIGURE 15. Plots show results for varying  $\eta$  while keeping the other parameters constant; in the images you can see the effects of  $\eta = 20$  in the dashed curve,  $\eta = 0$  in the dotted curve and  $\eta = 12$  in the solid line.

slower the degradation process, the more tumor cells invade the tissue and the less of a division is observable. The movement now also affects the concentration of the matrix-degrading enzymes. With the fewer remaining tumor cells at the origin producing the MDEs, we also see a more minor concentration at the origin. However, the distribution of the tumor cells for the lowest case of varying  $\eta$  has a more even distribution of MDEs.

The same goes for the last image, showing the experiments at  $t = 8$ . The more even the distribution of tumor cells and matrix-degrading enzymes across space, the slower the ECM is degraded.

As mentioned, varying  $\gamma$  does the term extracellular matrix include various organic or inorganic compounds. Therefore, the build-up and properties of these compounds vary strongly and motivate this comparison of degradation rate. Some compounds may be degraded faster, while others are complex and need more time to degrade.

### $d_m$ Variation

$d_m$  describes the diffusion coefficient for the matrix-degrading enzymes. As estimates, we use Anderson et al.'s and Franssen et al.'s values, which assume an even distribution of  $d_m$  in  $\sim U[1 \cdot 10^{-5}, 1 \cdot 10^{-3}]$ . Having set  $\beta = 0$ , equation 8 modeling the temporal development of the matrix-degrading enzymes concentration only depends on  $c$  concerning motility as well as in production. As we saw in experiments before, we can expect a high concentration

of MDEs in regions with a high density of tumor cells and we can expect increased motility where the density of the tumor cells rapidly changes. Increasing  $d_m$  will cause a faster, more even spread of the MDEs in the surrounding area and therefore, the ECM will also be degraded faster.

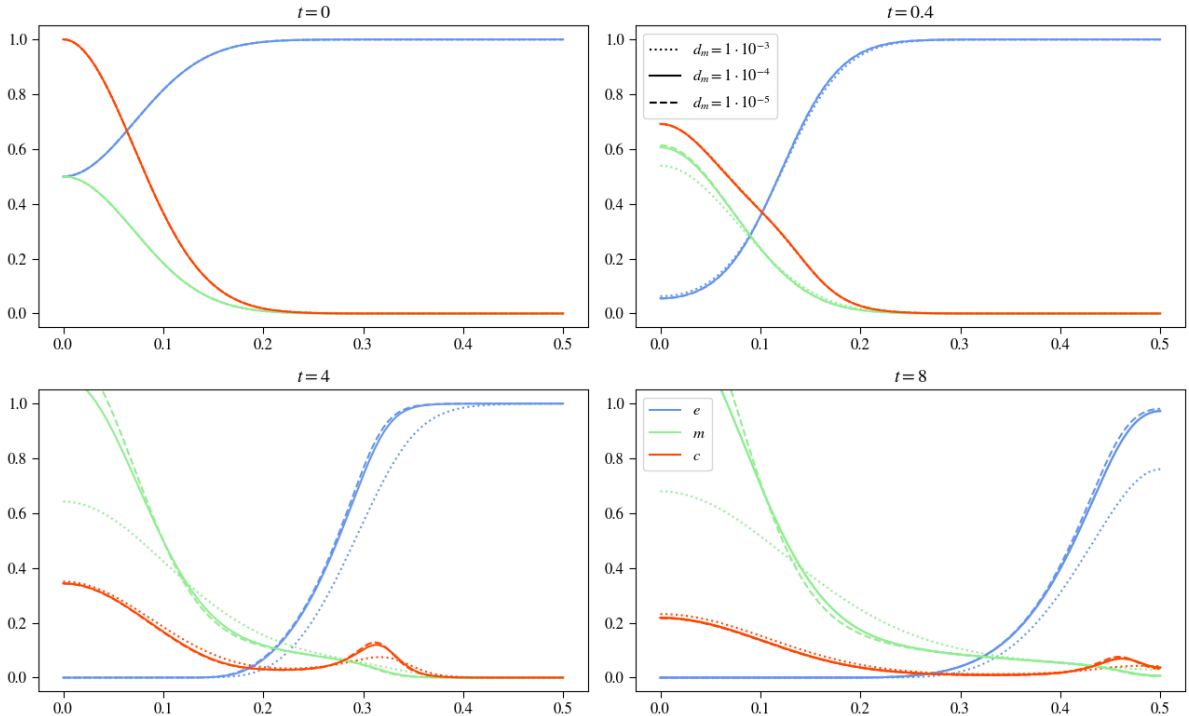


FIGURE 16. Plots show results for varying  $d_m$  while keeping the other parameters constant; in the images you can see the effects of  $d_m = 0.1$  in the dashed curve,  $d_m = 0$  in the dotted curve and  $d_m = 1 \cdot 10^{-3}$  in the solid line.

Inspecting the results in figure 16, we observe that in the second image after  $t = 0.4$ , the curves of the tumor cell density and the ECM are mostly untouched, though looking closely, we can see that for the highest  $d_m$  value, dotted line, the ECM has at the origin higher and farther out lower concentrations of the ECM. This is caused by the more even distribution of the matrix-degrading enzymes, which visibly reduce the concentration at the origin and invade the surrounding tissue slightly faster at this point.

The following image shows the experiments after  $t = 4$  and we see the differences between the lowest  $d_m$  value experiment with the other two more pregnant. The MDE concentration has, relative to the other two, strongly decreased at the origin, though it has invaded the tissue further. The ECM has degraded faster and as we saw in the  $\eta$  variation, the faster ECM degradation reduces the effects of haptotaxis. The other two experiments differ only visibly regarding the MDE concentration, though it exceeds the limits of the plot. The concentration at the origin for the lowest diffusive value is considerably higher than for the middle value.

The last image confirms the above mentioned effects, with the most substantial deviations between the highest  $d_m$  experiment and the other two. Again, a more even spread

of the MDEs with a lower concentration at the origin, a faster ECM degradation and tumor cell density curves that, compared to the other experiments, have less of a division at its leading edge. The two experiments with lower  $d_m$  values still have nearly overlaying tumor cell density and ECM curves. Only the MDE concentration differs between the two, showing a more even spread for the higher diffusion value, though with a lower value at the origin.

Seeing those effects, we saw our intuition met, though it is to say that looking at the lower two values for  $d_m$ , the differences here were modest. Therefore, the higher the value for  $d_m$  is, the more sensitive the system reacts to this change.

These changes in diffusion, like in the section varying the diffusion coefficient of the tumor cells, might be caused by a multitude of physical influences, such as temperature, voltage, pressure, etc. The results show that this change causes a faster degradation of the extracellular matrix molecules but a less aggressive invasion of the tumor cells.

### $\alpha$ Variation

The parameter  $\alpha$  influences how fast the tumor cells produce matrix-degrading enzymes. We assume it to be evenly distributed with  $\alpha \sim U[0, 1.0]$  since Anderson et al. assumed the same range in the original paper. Trying to replicate Anderson et al.'s experiment, we already saw how varying  $\alpha$  affects the simulation results. With growing  $\alpha$ , we will see a higher concentration of MDEs, especially in regions with high tumor cell density. More MDEs will cause faster degradation of the ECM first due to having more of them, but also because since more of them are subject to diffusion, they will spread faster in the tissue. Faster ECM degrading could mean an increased invasion pace of the tumor cells. As we saw in the previous experiments varying  $d_c$ , the MDE concentration can take on values higher than one. We can also expect this here when  $\alpha$  is sufficiently high.

The second image of figure 17 describes the experiments after  $t = 0.4$ . We can clearly see the differences in the MDE curves for the different values of  $\alpha$ . The other curves also show slight deviations. The higher  $\alpha$  is, the faster the ECM degradation. Looking closely, we can observe for the tumor cell density that the increased rate of ECM degradation results, as previously already seen, in a curve with less of a bump that will later form the division invading the surrounding tissue due to more minor haptotactic influences at the origin.

The next point in time, at  $t = 4$ , shows the previously mentioned effects in a reinforced way. For  $\alpha = 0$ , the MDEs have no producing factor and the curve flattens due to diffusion, though the extracellular matrix has degraded visibly, yet at a considerably slower rate. This experiment has the lowest density of tumor cells at the origin, though the most significant secondary lump of cells invades the tissue. This behavior is due to the extended exposition of strong haptotactic influences near the origin region, which pulls more cells outward to invade. The solid curves describing  $\alpha = 0.6$  show that at this point, they have almost reached a concentration of one at the center, which will be exceeded later. Compared to the lowest  $\alpha$  experiment, the faster ECM concentration has also pulled less of the tumor cells outward, though at a faster invasion pace. Looking at the experiment with the highest  $\alpha$  value, the MDE concentration already exceeds one, the ECM degradation is happening faster and the invasion of the tumor cells is happening

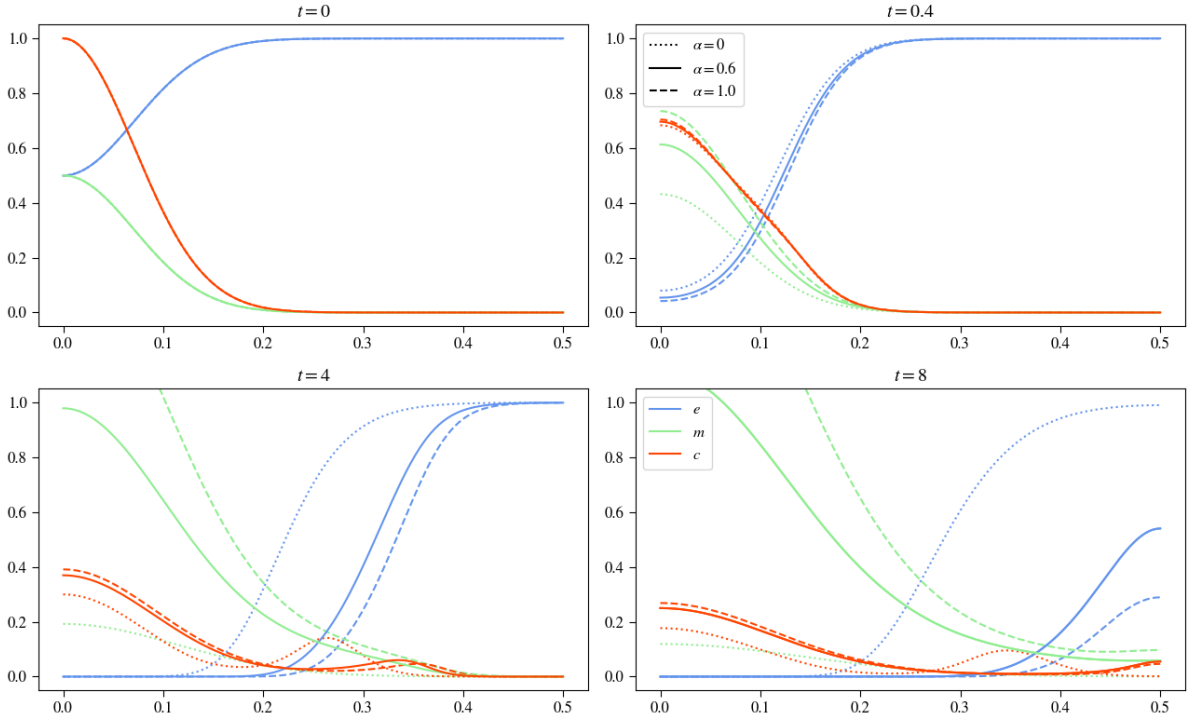


FIGURE 17. Plots show results for varying  $\alpha$  while keeping the other parameters constant; in the images, you can see the effects of  $\alpha = 1.0$  in the dashed curve,  $\alpha = 0$  in the dotted curve and  $\alpha = 0.6$  in the solid line.

faster, though with a lower density.

In the last timestep, we see that the MDE curve of  $\alpha = 0.6$  and  $\alpha = 1.0$  exceeded one. For the dotted curve of the MDE, we have a good example of diffusion distributing the concentration throughout space without changing its overall volume. At the border regions, we see that the dotted curve is also the only one that has yet to degrade any ECM in this area, while the other two experiments show that there is only a little ECM concentration left to degrade. The tumor cells confirm the, in the previous timestep mentioned, effects of faster invasion pace with rising  $\alpha$  though with at a thinner density.

Our initial assumptions were correct with a faster degradation pace due to higher MDE concentration and, therefore, a faster invasion pace of the tumor cells. However, it is interesting to see that the more minor  $\alpha$ , the more tumor cells are being pulled into the tissue.

While it makes sense from a numerical perspective that the concentration of MDEs can exceed one, it might make sense to introduce a finer grid or adapt the model in other ways since, judging from a continuous perspective, it does not make sense that at a certain point in space, there is more than one entity occupying this space.

The production of matrix-degrading enzymes can have many biological causes. During many natural processes like tissue repair or remodeling, the extracellular matrix must be degraded, controlled by cells producing enzymes that, in turn, are responsible for producing matrix-degrading enzymes. This control flow can be interrupted by malignant

cells stimulating the production of MDEs without any repair or remodeling tasks to perform. Considering the absence of or reduced MDE production, we can regard this case as the consequence of a drug treatment. On the other hand, since there are plenty of different extracellular matrix molecules, they may also have different production rates.

### $\beta$ Variation

The factor  $\beta$  controls the decay of the matrix-degrading enzymes and the results of its variation are shown in figure 18. Using Kolev et al.’s estimate for  $\beta$  in [13], we experimented with  $\beta = 0.07$ . Considering the results, we can assume an even distribution of  $\beta$  in  $\beta \sim U[0.005, 0.1]$ . Interestingly,  $\alpha$  and  $\beta$  are their respective distribution and the experiments will confirm that they are not of the same magnitude. While the MDEs are produced by the tumor cells, their decay is controlled by the MDEs themselves. This model assumes that the tumor cells do not proliferate, which makes their total amount constant. In contrast, the MDEs are produced by the tumor cells and are expected to change in amount over time due to production or decay. This makes the decay rate of the MDEs more variable because of the changing amount of total MDEs, causing the change of effective magnitude.

With introducing  $\beta$  foremost, we can assume that the MDE curve will be lower, influencing the ECM degrading process and, therefore, the invasion pace and the diffusion-haptotaxis effects on the tumor cells.

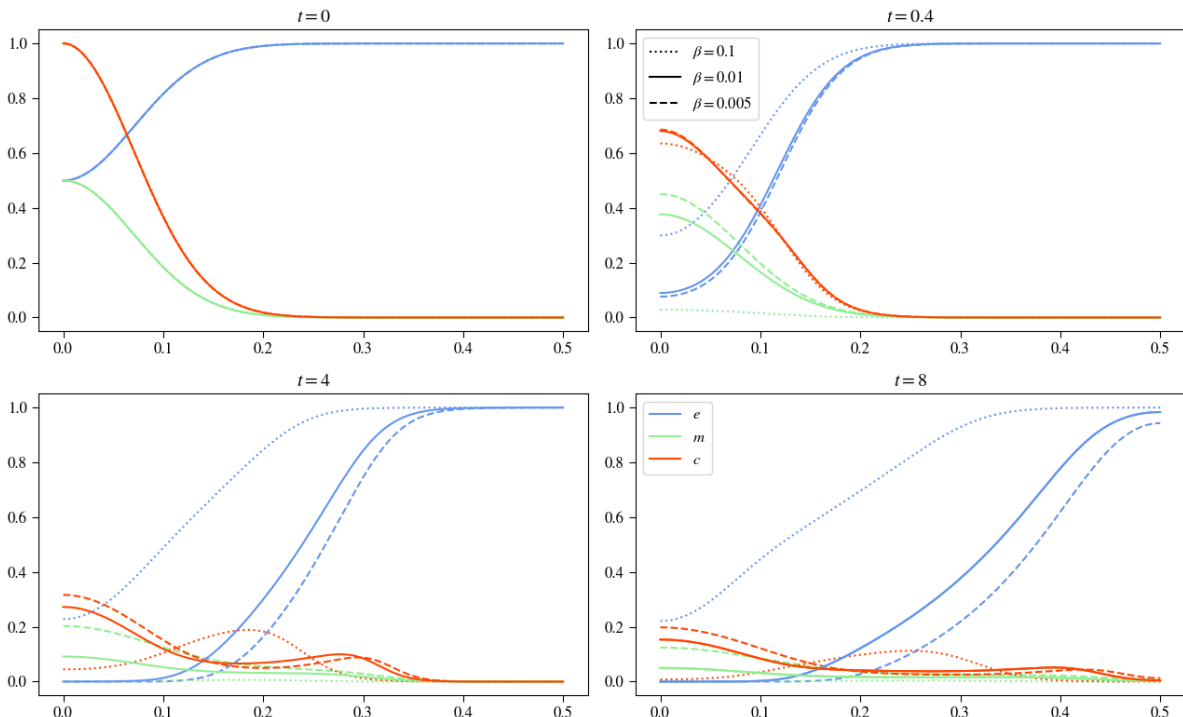


FIGURE 18. Plots show results for varying  $\beta$  while keeping the other parameters constant; in the images, you can see the effects of  $\beta = 0.005$  in the dashed curve,  $\beta = 0.1$  in the dotted curve and  $\beta = 0.01$  in the solid line.

As we see from the results in figure 18, a value of  $\beta = 0.1$  is sufficient to after already  $t = 4$  reduce the MDE concentration to nearly zero. In our case, this decay rate proves too high, outpacing production entirely, with matrix-degrading enzyme concentration vanishing spacially and temporarily completely. The immediate decay of the MDEs causes a drastically slower ECM degradation, yet it has not stopped entirely since the tumor cells still produce matrix-degrading enzymes. This slow degradation of the extracellular matrix, as we saw previously, creates a haptotactic pull that lasts a lot longer in regions around the origin, which pulls, in this case, all of the cells outward to invade the tissue, leaving no primary lump of tumor cells at the origin. Since more MDEs are produced in regions with high tumor cell density, this also causes a more even degradation of the extracellular matrix, causing a more substantial stretch of the tumor cells.

In the other experiments, we can observe that with decreasing  $\beta$  and slowing down the decay of the matrix-degrading enzymes, first, the ECM degradation accelerates and this causes the effects of haptotaxis and diffusion to develop the two lumps of tumor cells, one staying at the center the other invading the tissue as we saw in all previous experiments.

Since the extracellular matrix can comprise many different organic and inorganic compounds, the degrading enzymes are also very diverse. This diversity results in different decay rates, so choosing the right one for a specific experiment can be crucial. While this variation describes the effects of different enzymes, it can also describe, like for the production of the MDEs, the influence of a drug to accelerate the decay of the matrix-degrading enzymes.

### Cross-Variation

Having varied every model parameter, we saw accelerating effects on the invasion and countering effects. Now, it will be interesting to see how either supporting factors interplay and how countering factors affect the simulations, for example, how to increase both  $\alpha$  and  $\beta$  if some balance can be found. Another exciting experiment will be on how diffusion and haptotaxis behave when increasing or decreasing both factors controlling them.

### $d_c - \gamma$ Variation

We, therefore, started with varying  $d_c$  and  $\gamma$  and studied for simplicity and overview reasons only the values on their distribution's borders interplaying, shown in figure 19.

Having set  $d_c = 5 \cdot 10^{-5}$  and  $\gamma = 0.001$ , we see no division of the tumor cells; the effects of haptotaxis are too small, leaving the tumor cells only subject to diffusion, which results in an even distribution process over time, which also causes a slower invasion pace. Because the tumor cells stay in a lump with their maxima at the origin  $x = 0$ , the MDEs also take on their maximum there, moving farther out and distributing very evenly. This staying with values around the origin of the MDEs causes a slower ECM degradation.

Increasing  $\gamma = 0.01$ , we see that the effects of haptotaxis are now pregnantly visible with a very sharp maximum seen at  $t = 4$ , which equals the maxima of the remaining tumor cell lump at  $x = 0$ . A more substantial influence of haptotaxis leads to a faster invasion pace of the tumor cells into the tissue. It allows the creation of matrix-degrading

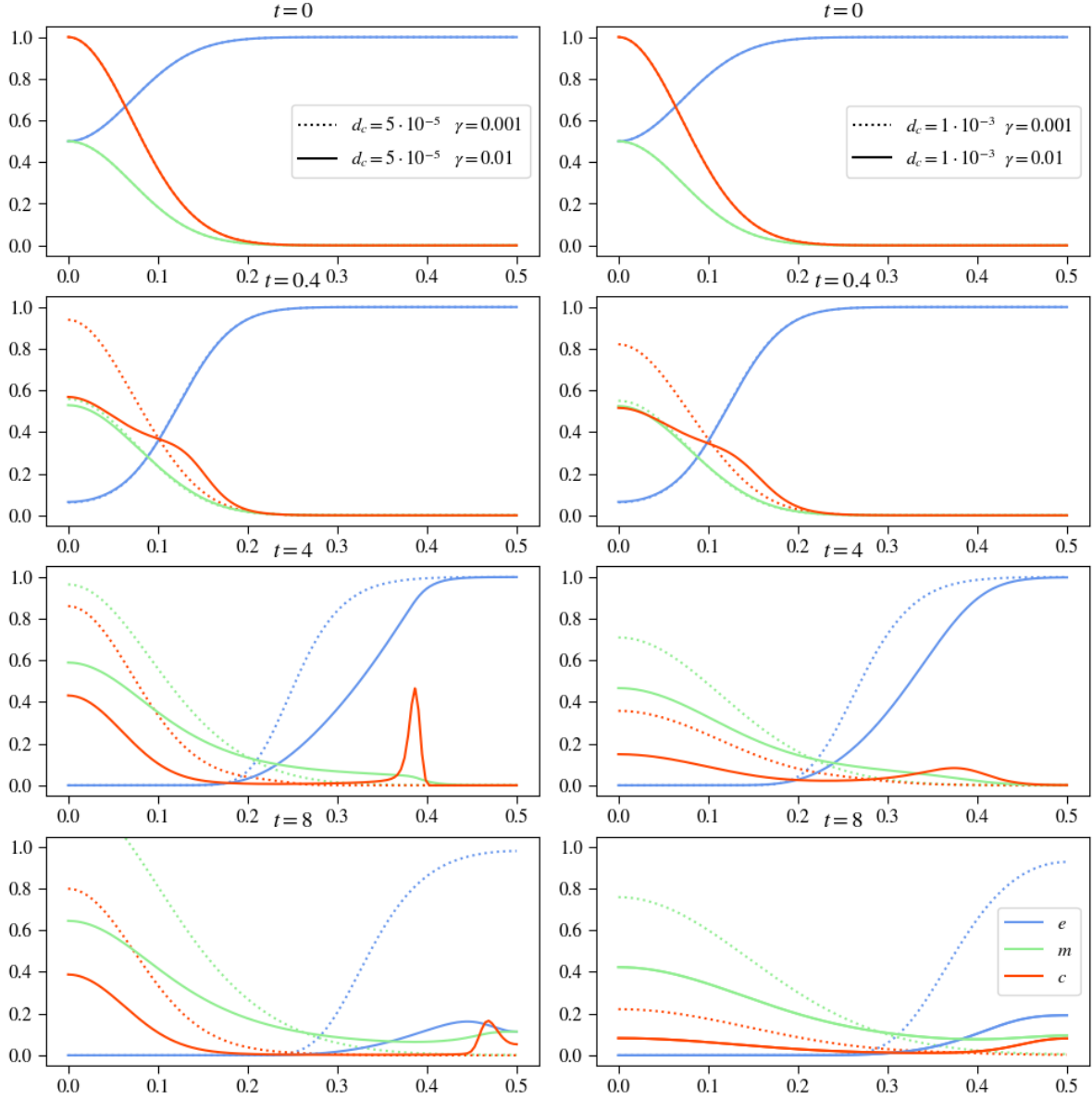


FIGURE 19. Plots show results for varying both  $d_c$  and  $\gamma$  while keeping the other parameters constant; in the images on the left,  $d_c$  is set to  $d_c = 0.00005$  with the solid line showing  $\gamma = 0.01$  and the dotted line  $\gamma = 0.001$  on the right  $d_c$  is set to  $d_c = 0.1$  with the solid line showing  $\gamma = 0.01$  and the dotted line  $\gamma = 0.001$ .

enzymes in their wake, causing a more even distribution compared to  $\gamma = 0.001$  and a faster ECM degrading process.

Looking at the right side of the plot 19, we see the results for  $d_c = 1 \cdot 10^{-3}$ . Comparing  $\gamma$  values with this  $d_c$  value, we can, as on the left side, see that the experiment showing  $\gamma = 0.001$  does not form a division for the tumor cells, with the main lump of cell staying at the origin. No secondary site of tumor cells is formed that invades the tissue faster. This is also shown on the ECM concentration for the dotted experiment, with a considerably slower degradation process and when comparing both dotted experiments on left and right

side images. Staying near the center of the tumor cells also causes a much higher MDE concentration there compared to the  $\gamma$ -increased experiment.

Increasing both  $d_c$  and  $\gamma$ , we see the effects of haptotaxis more potent with the secondary site of tumor forming. However, comparing this experiment with the experiment having high  $\gamma$  and low  $d_c$  values shows that the division has no peaky leading edge.

These experiments give us an excellent example of the relationship between haptotaxis and diffusion. We see that with decreasing  $d_c$  but increasing  $\gamma$ , we get a leading edge that is very pointy. The higher value of  $\gamma$  was sufficient to form a secondary tumor site invading the tissue separately. Decreasing  $\gamma$  has proven for high and low  $d_c$  values to prevent a secondary lump of cells from forming.

### $d_m - \eta$ Variation

Comparing both columns for varying  $d_m$  in figure 20, though looking at the same  $\eta$  valued experiments, we can only see minor differences regarding the shape of the tumor cell density curve and ECM concentration curve. On the left side, we see that the ECM degradation is happening slower, which causes a different relation regarding the diffusion-haptotaxis effects. The curves on the left side are not as evenly stretched as on the right, with a more significant hill at the secondary lump of cells. This behavior is due to the extended exposition of more substantial haptotaxis effects on the tumor cells in regions closer to the origin.

The ECM degradation is happening slower on the left side due to minor diffusive motility, which is slowing the MDE invasion process.

Interestingly, the MDE concentration for lower  $d_m$  values is considerably higher for both experiments than for the ones with higher  $d_m$  values. Since the tumor cells produce the MDEs, it is apparent that only the slightest visible change in the tumor cell density is sufficient to cause such drastic changes in the production of the MDEs due to exponential growth.

These experiments conclude that the diffusive properties of the matrix-degrading enzymes play only a minor role in degrading the extracellular matrix. Tumor cells producing them in their wake play a more crucial role in the degradation process. An intriguing observation in this experiment is that minimal differences in tumor cell density are sufficient to create major differences in the MDE concentration due to their exponential production.

### $\alpha - \beta$ Variation

Looking at figure 21 we see experimental results varying both  $\alpha$  and  $\beta$ . For low MDE production and low MDE decay, we can see that the curve for the MDEs is still visible at up to  $t = 4$ ; at  $t = 8$ , it is zero. We see that the ECM degrading happens faster than for high  $\beta$  values; therefore, the tumor cells develop two lumps, with one invading the tissue and the other staying at  $x = 0$ . The maxima for both lumps are lower than in previous experiments, though the cells seem to be more evenly distributed between the two lumps. Increasing  $\beta = 0.1$ , the MDE curve appears to be zero after already  $t = 0.4$  and stays there until the end of this experiment. This low concentration of MDEs causes a slower ECM degrading process, leading the tumor cells to develop only one lump, invading the

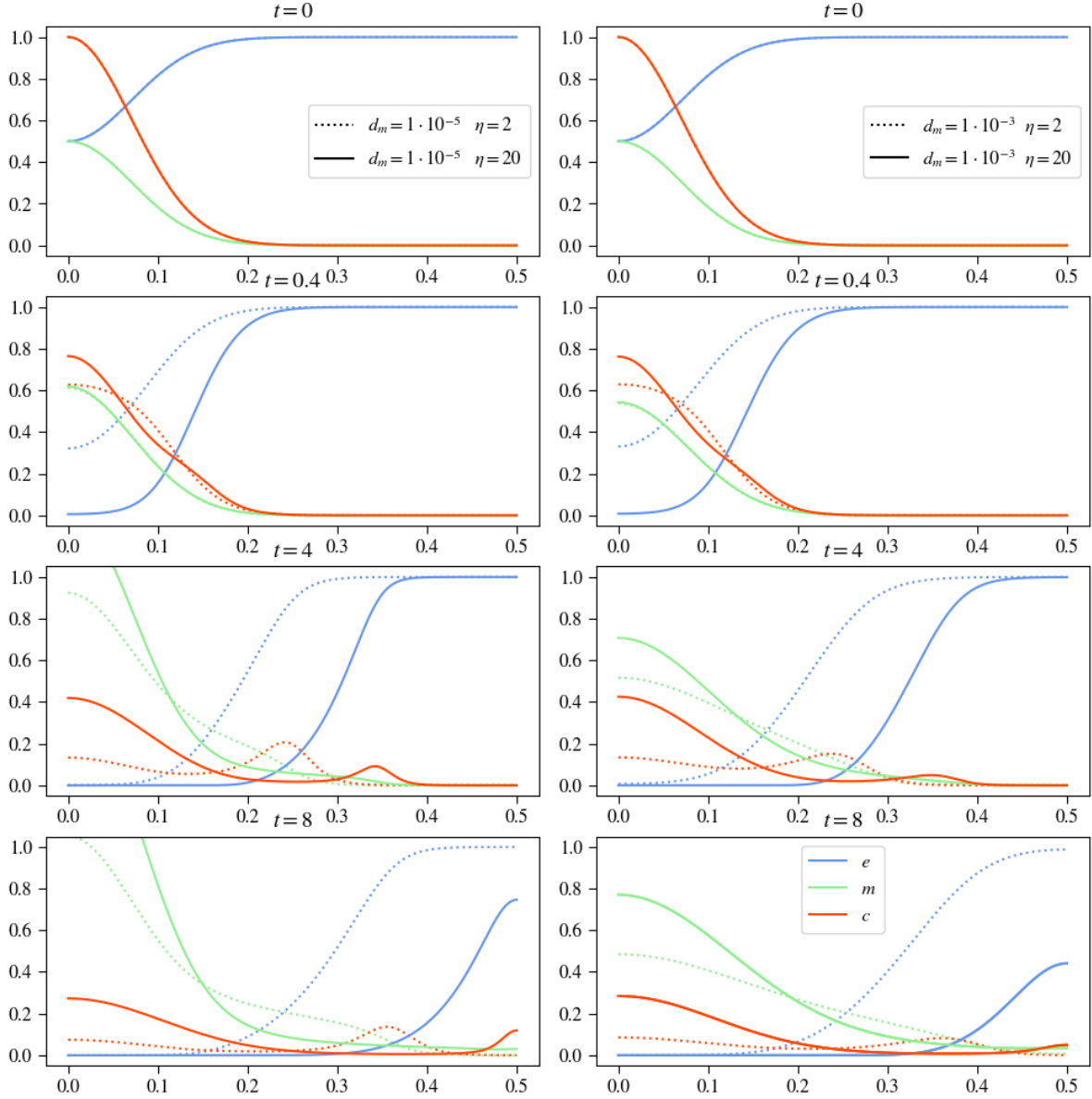


FIGURE 20. Plots show results for varying both  $d_m$  and  $\eta$  while keeping the other parameters constant, in the images on the left  $d_m$  is set to  $d_m = 1 \cdot 10^{-5}$  with the solid line showing  $\eta = 2$  and the dotted line  $\eta = 20$  on the right  $d_m$  is set to  $d_m = 1 \cdot 10^{-3}$  with the solid line showing  $\eta = 2$  and the dotted line  $\eta = 20$ .

space, with its maxima moving at the center of this lump. For  $\alpha = 0.1$ , both values for  $\beta$  have proven too high, decaying the matrix-degrading enzymes too fast to keep up with production.

On the other hand, increasing  $\alpha$  to 1.0 and keeping  $\beta = 0.05$ , we see that production outweighs decay, with at the end of the experiment, the MDEs still have a concentration of about 0.4 at  $x = 0$ . For this experiment, we see that the tumor cells develop two lumps, indicating that diffusion and haptotaxis effects are also in some balance and ECM degradation seems to resemble, due to similarities with the basecase for the MDE curve,

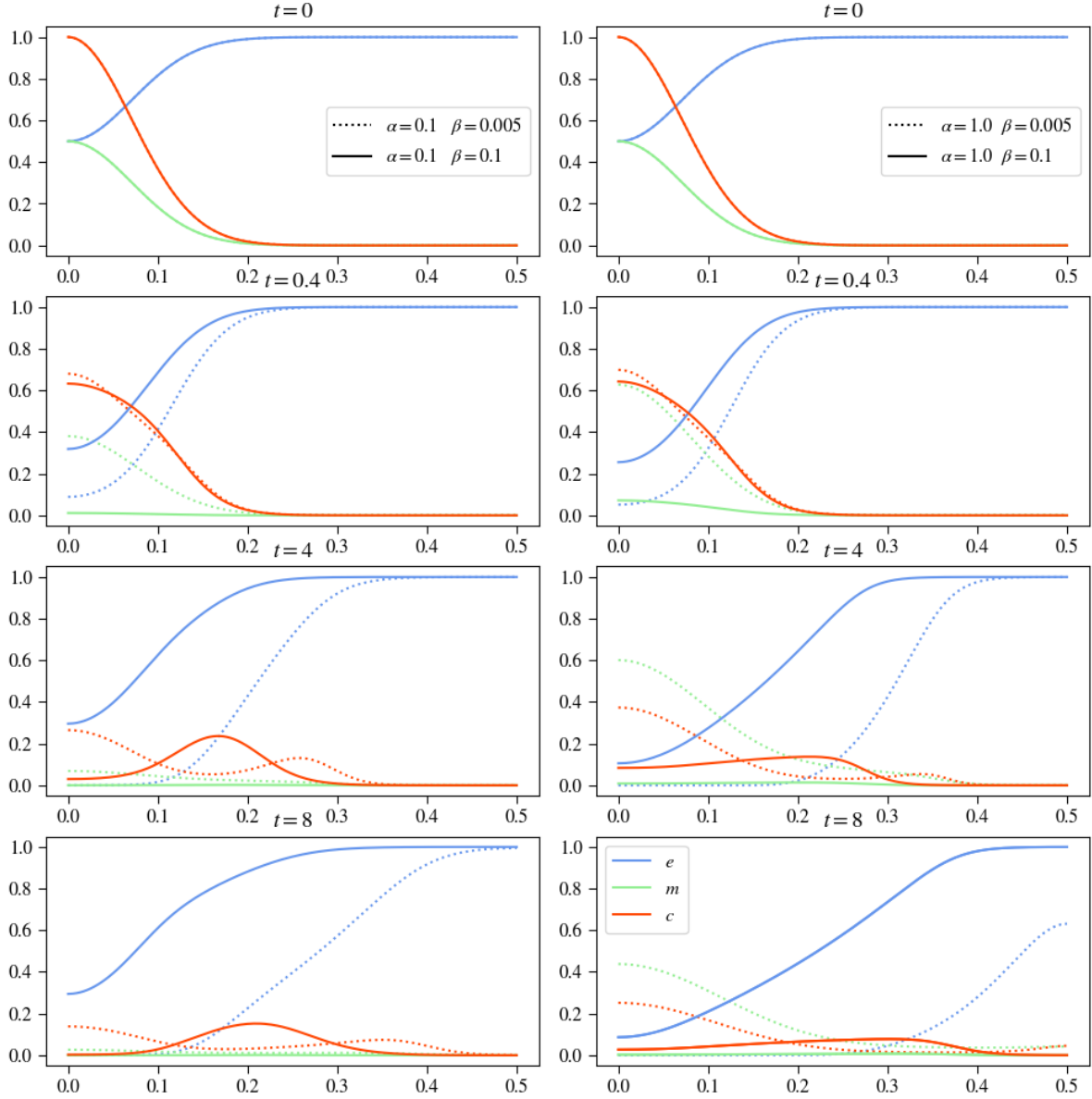


FIGURE 21. Plots show results for varying both  $\alpha$  and  $\beta$  whilst keeping the other parameters constant, in the images on the left  $\alpha = 0.1$  with the solid line showing  $\beta = 0.005$  and the dotted line  $\beta = 0.1$  on the right  $\alpha = 1.0$  with the solid line showing  $\beta = 0.005$  and the dotted line  $\beta = 0.1$ .

also the ECM degradation of the basecase experiment. Increasing both  $\alpha$  and  $\beta$ , we see in the solid line of the right column of figure 21 that decay outweighs production again, after  $t = 0.4$  we can only see a small remaining portion of matrix-degrading enzymes at the origin. This causes slower ECM degradation and forms only one lump of tumor cells due to the strong effects of haptotaxis, though this singular lump is stretched flat along the x-axis.

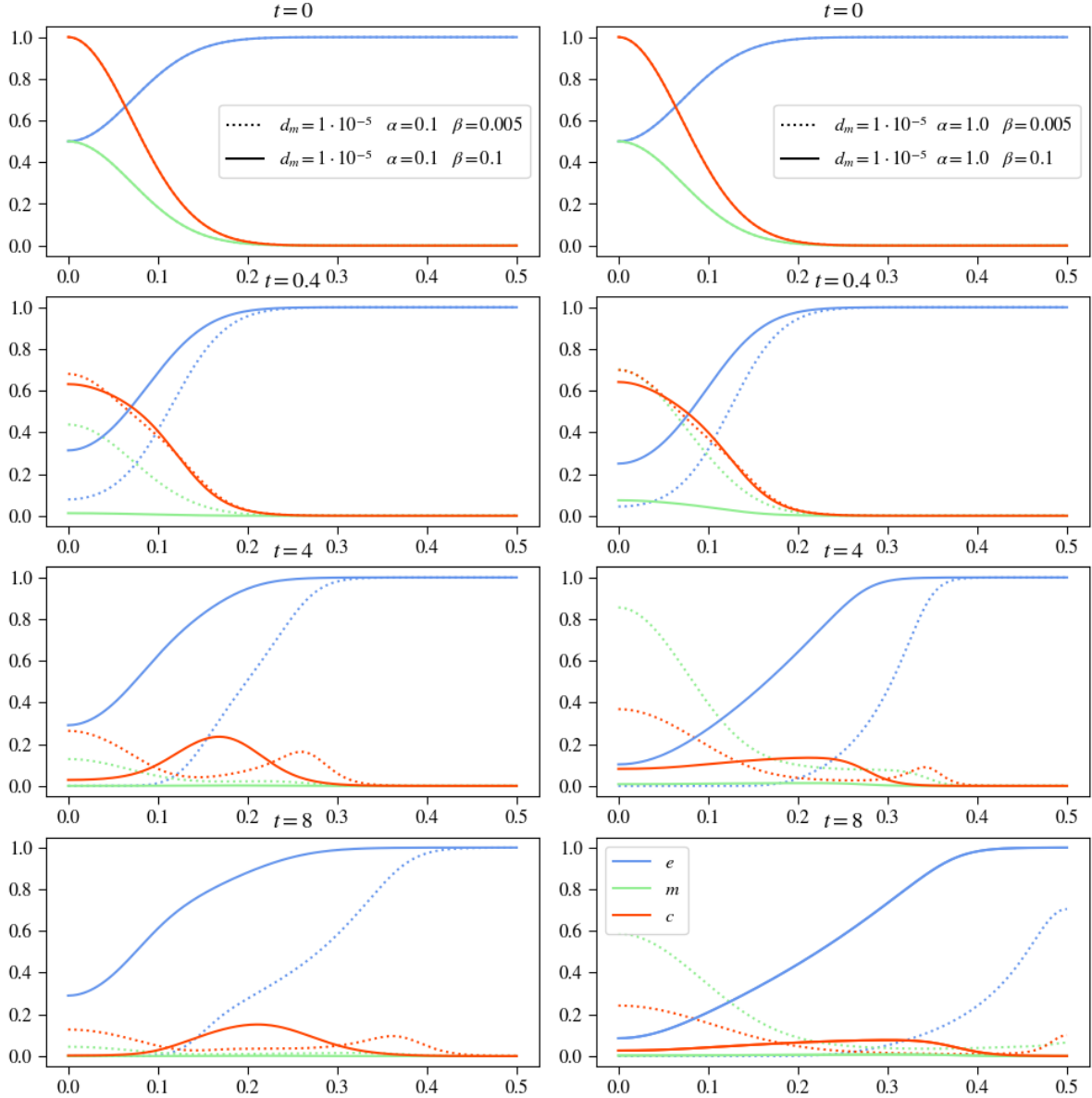
$d_m - \alpha - \beta$  Variation

FIGURE 22. Plots show results for varying both  $\alpha$  and  $\beta$  whilst keeping the other parameters constant, in the images on the left  $\alpha = 0.1$  with the solid line showing  $\beta = 0.005$  and the dotted line  $\beta = 0.1$  on the right  $\alpha = 1.0$  with the solid line showing  $\beta = 0.005$  and the dotted line  $\beta = 0.1$ .

Varying all parameters regarding the equation for the matrix-degrading enzymes required to split the results into two figures, 22 and 23.

Looking at the results, we see that for the same  $\alpha$  and  $\beta$  values, the curve describing the tumor cell density shares the behavior across varying  $d_m$ . For low  $\beta$  values, we see in every experiment that the tumor cells develop the secondary lump that detaches from the primary lump and invades the surrounding tissue separately. Considering how  $\alpha$

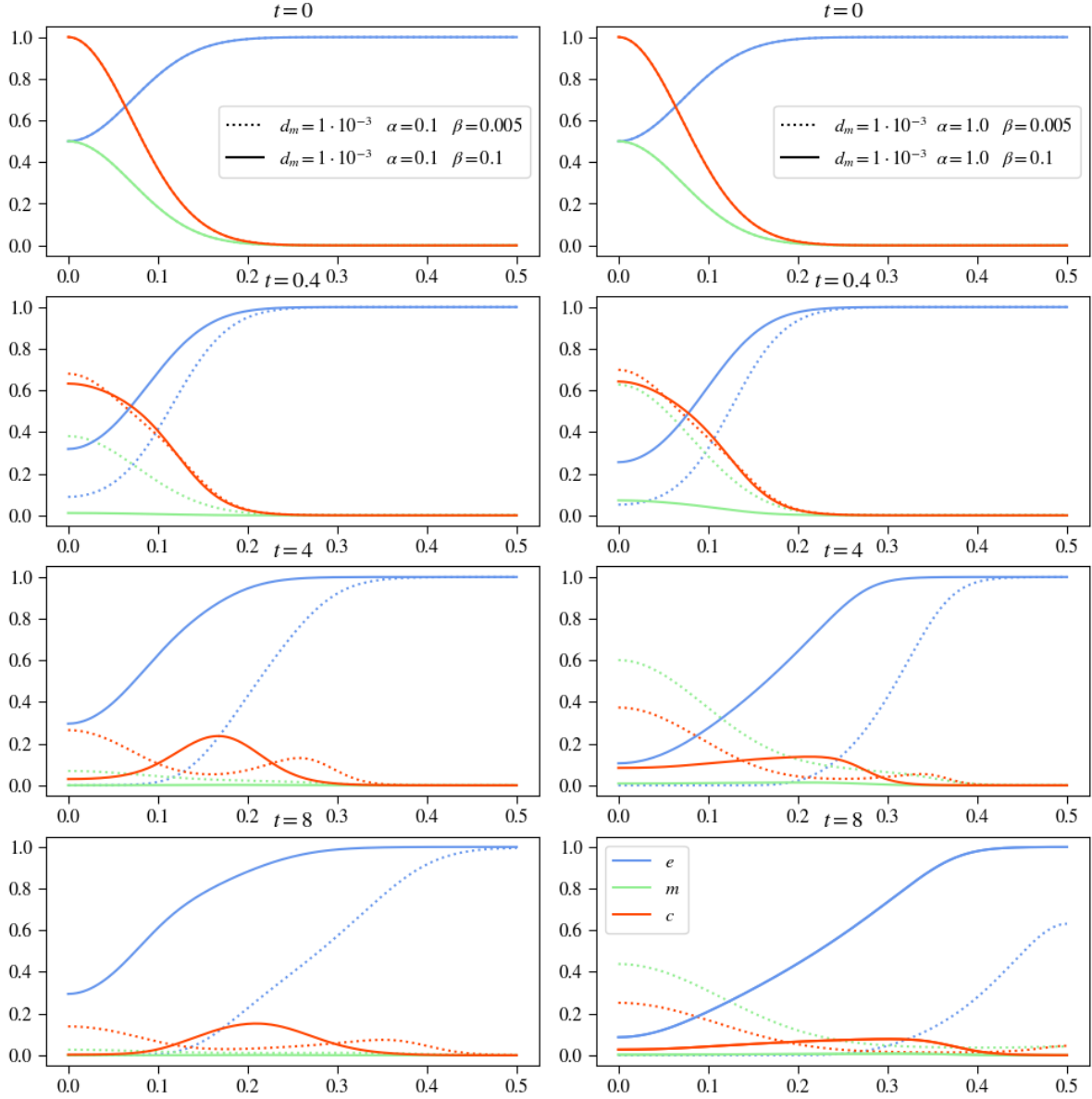


FIGURE 23. Plots show results for varying both  $\alpha$  and  $\beta$  whilst keeping the other parameters constant, in the images on the left  $\alpha = 0.1$  with the solid line showing  $\beta = 0.005$  and the dotted line  $\beta = 0.1$  on the right  $\alpha = 1.0$  with the solid line showing  $\beta = 0.005$  and the dotted line  $\beta = 0.1$ .

influences this lump, we observe that with increasing  $\alpha$ , the lump gets smaller and more of the tumor cells stay near the origin. Taking the experiments with high  $\beta$  values in regard, we observe that this curve fails to develop a secondary lump of tumor cells to invade separately. As seen in all the experiments varying  $\eta$ , a slowed degradation rate of the extracellular matrix leads to the extended exposition of the tumor cells at the origin to more substantial haptotaxis effects. This leads to all tumor cells being pulled outward, leaving little or no near the origin. This results in only one lump of cells that invades, with its maximum moving into the surrounding tissue. In the experiments with high values

for  $\beta$ , we see that the diffusion of the matrix-degrading enzymes has nearly no effect, as there are no apparent changes in the plot images.

Taking the curve of the extracellular matrix into account now, we can also see minimal deviation when varying  $d_m$ . We see no apparent changes in figures 22 and 23 regarding the ECM concentration which supports our claim from above that the diffusion of the matrix-degrading enzymes plays only a minor role in degrading the ECM, but also for the whole system in total.

The MDE curve can explain the effects mentioned above on ECM and tumor cell density curves. We see that for every experiment where  $\beta$  was set to  $\beta = 0.1$ , the curve of the MDEs was after already  $t = 0.4$  zero. Increasing the production rate  $\alpha$  was visibly insufficient to counter the matrix-degrading enzymes' fast decay process, resulting in the drastically slowed ECM degradation process, which in turn is responsible for the haptotaxis effects on the tumor cells. This explains why we saw no visible changes in the experiments varying  $d_m$  and  $\alpha$ . This again shows the significance of the decay parameter of the matrix-degrading enzymes.

### 4.3 2D Results with Proliferation and Renewal - Homogenous ECM

This section will examine how introducing tumor cell proliferation and extracellular matrix renewal influences the system. Modeled as logistical growth terms, with a limiting factor of spacial occupation, the parameters  $\mu_1$  for tumor cell proliferation and  $\mu_2$  for extracellular matrix renewal describe their influences. Those parameters were also incorporated in the previous experiments, though they were set to zero. Instead of treating them as parameters to vary, we are inspecting the whole system as new since they can drastically change the resulting simulations.

Tumor cell proliferation describes the natural ability of cells to proliferate. However, in the case of tumor cells, the production signaling comes from themselves, surpassing the control chain that limits normal cells to proliferate uncontrollably.

The extracellular matrix is a naturally dynamic structure that undergoes continuous degradation and remodeling. It does this in order to secure tissue development, wound repair tasks, regulate cellular functions and many more tasks. In a model describing the ECM, it is crucial to introduce a factor that incorporates these remodeling processes.

As in section 4.2, we use the same initial conditions for all three variables, assuming we have a homogenous ECM structure. Figure 3 depicts these initial conditions on the three variables  $c, e, m$  at dimensionless time  $t = 0$ .

Figure	Linestyle	$d_c$	$\gamma$	$\mu_1$	$\eta$	$\mu_2$	$d_m$	$\alpha$	$\beta$
24	.....	$5 \cdot 10^{-4}$	0.0055	0	10	0	$1 \cdot 10^{-3}$	0.3564	0
24	—	$5 \cdot 10^{-4}$	0.0055	0.1	10	0.5	$1 \cdot 10^{-3}$	0.3564	0
25	.....	$1 \cdot 10^{-3}$	0.0055	0.1	10	0.5	$1 \cdot 10^{-3}$	0.3564	0
25	—	$1 \cdot 10^{-4}$	0.0055	0.1	10	0.5	$1 \cdot 10^{-3}$	0.3564	0

Figure	Linestyle	$d_c$	$\gamma$	$\mu_1$	$\eta$	$\mu_2$	$d_m$	$\alpha$	$\beta$
25	---	$5 \cdot 10^{-5}$	0.0055	0.1	10	0.5	$1 \cdot 10^{-3}$	0.3564	0
26	.....	$5 \cdot 10^{-4}$	0.002	0.1	10	0.5	$1 \cdot 10^{-3}$	0.3564	0
26	—	$5 \cdot 10^{-4}$	0.008	0.1	10	0.5	$1 \cdot 10^{-3}$	0.3564	0
26	----	$5 \cdot 10^{-4}$	0.01	0.1	10	0.5	$1 \cdot 10^{-3}$	0.3564	0
27	.....	$5 \cdot 10^{-4}$	0.0055	0	10	0.5	$1 \cdot 10^{-3}$	0.3564	0
27	—	$5 \cdot 10^{-4}$	0.0055	0.5	10	0.5	$1 \cdot 10^{-3}$	0.3564	0
27	----	$5 \cdot 10^{-4}$	0.0055	1.0	10	0.5	$1 \cdot 10^{-3}$	0.3564	0
28	.....	$5 \cdot 10^{-4}$	0.0055	0.1	2	0.5	$1 \cdot 10^{-3}$	0.3564	0
28	—	$5 \cdot 10^{-4}$	0.0055	0.1	12	0.5	$1 \cdot 10^{-3}$	0.3564	0
28	----	$5 \cdot 10^{-4}$	0.0055	0.1	20	0.5	$1 \cdot 10^{-3}$	0.3564	0
29	.....	$5 \cdot 10^{-4}$	0.0055	0.1	10	0.1	$1 \cdot 10^{-3}$	0.3564	0
29	—	$5 \cdot 10^{-4}$	0.0055	0.1	10	0.6	$1 \cdot 10^{-3}$	0.3564	0
29	----	$5 \cdot 10^{-4}$	0.0055	0.1	10	1.0	$1 \cdot 10^{-3}$	0.3564	0
30	.....	$5 \cdot 10^{-4}$	0.0055	0.1	10	0.5	$1 \cdot 10^{-3}$	0.3564	0
30	—	$5 \cdot 10^{-4}$	0.0055	0.1	10	0.5	$1 \cdot 10^{-4}$	0.3564	0
30	----	$5 \cdot 10^{-4}$	0.0055	0.1	10	0.5	$1 \cdot 10^{-5}$	0.3564	0
31	.....	$5 \cdot 10^{-4}$	0.0055	0.1	10	0.5	$1 \cdot 10^{-3}$	0	0
31	—	$5 \cdot 10^{-4}$	0.0055	0.1	10	0.5	$1 \cdot 10^{-3}$	0.6	0
31	----	$5 \cdot 10^{-4}$	0.0055	0.1	10	0.5	$1 \cdot 10^{-3}$	1.0	0
32	.....	$5 \cdot 10^{-4}$	0.0055	0.1	10	0.5	$1 \cdot 10^{-3}$	0.3564	0.1
32	—	$5 \cdot 10^{-4}$	0.0055	0.1	10	0.5	$1 \cdot 10^{-3}$	0.3564	0.01
32	----	$5 \cdot 10^{-4}$	0.0055	0.1	10	0.5	$1 \cdot 10^{-3}$	0.3564	0.005
33 - left	.....	$5 \cdot 10^{-4}$	0.0055	0.1	10	0.1	$1 \cdot 10^{-3}$	0.3564	0
33 - left	—	$5 \cdot 10^{-4}$	0.0055	0.1	10	1.0	$1 \cdot 10^{-3}$	0.3564	0
33 - right	.....	$5 \cdot 10^{-4}$	0.0055	1.0	10	0.1	$1 \cdot 10^{-3}$	0.3564	0
33 - right	—	$5 \cdot 10^{-4}$	0.0055	1.0	10	1.0	$1 \cdot 10^{-3}$	0.3564	0
34 - left	.....	$1 \cdot 10^{-5}$	0.001	0.1	10	0.5	$1 \cdot 10^{-3}$	0.3564	0
34 - left	—	$1 \cdot 10^{-5}$	0.001	1.0	10	0.5	$1 \cdot 10^{-3}$	0.3564	0
34 - right	.....	$1 \cdot 10^{-5}$	0.01	0.1	10	0.5	$1 \cdot 10^{-3}$	0.3564	0
34 - right	—	$1 \cdot 10^{-5}$	0.01	1.0	10	0.5	$1 \cdot 10^{-3}$	0.3564	0
35 - left	.....	$1 \cdot 10^{-3}$	0.001	0.1	10	0.5	$1 \cdot 10^{-3}$	0.3564	0
35 - left	—	$1 \cdot 10^{-3}$	0.001	1.0	10	0.5	$1 \cdot 10^{-3}$	0.3564	0
35 - right	.....	$1 \cdot 10^{-3}$	0.01	0.1	10	0.5	$1 \cdot 10^{-3}$	0.3564	0
35 - right	—	$1 \cdot 10^{-3}$	0.01	1.0	10	0.5	$1 \cdot 10^{-3}$	0.3564	0
36 - left	.....	$5 \cdot 10^{-4}$	0.0055	0.1	2	0.1	$1 \cdot 10^{-3}$	0.3564	0
36 - left	—	$5 \cdot 10^{-4}$	0.0055	0.1	2	1.0	$1 \cdot 10^{-3}$	0.3564	0
36 - right	.....	$5 \cdot 10^{-4}$	0.0055	0.1	20	0.1	$1 \cdot 10^{-3}$	0.3564	0
36 - right	—	$5 \cdot 10^{-4}$	0.0055	0.1	20	1.0	$1 \cdot 10^{-3}$	0.3564	0

TABLE 4. Overview of all experiments conducted for the model with proliferation and renewal producing 2D output

As in section 4.2, we see in table 4 a detailed overview of all the experiments done in this section and the parameters used to produce the results. As before, most figures describe multiple experiments; the linestyle of the curve in the figure determines which experiment is precisely described by the set of parameters.

#### 4.3.1 Basecase Analysis

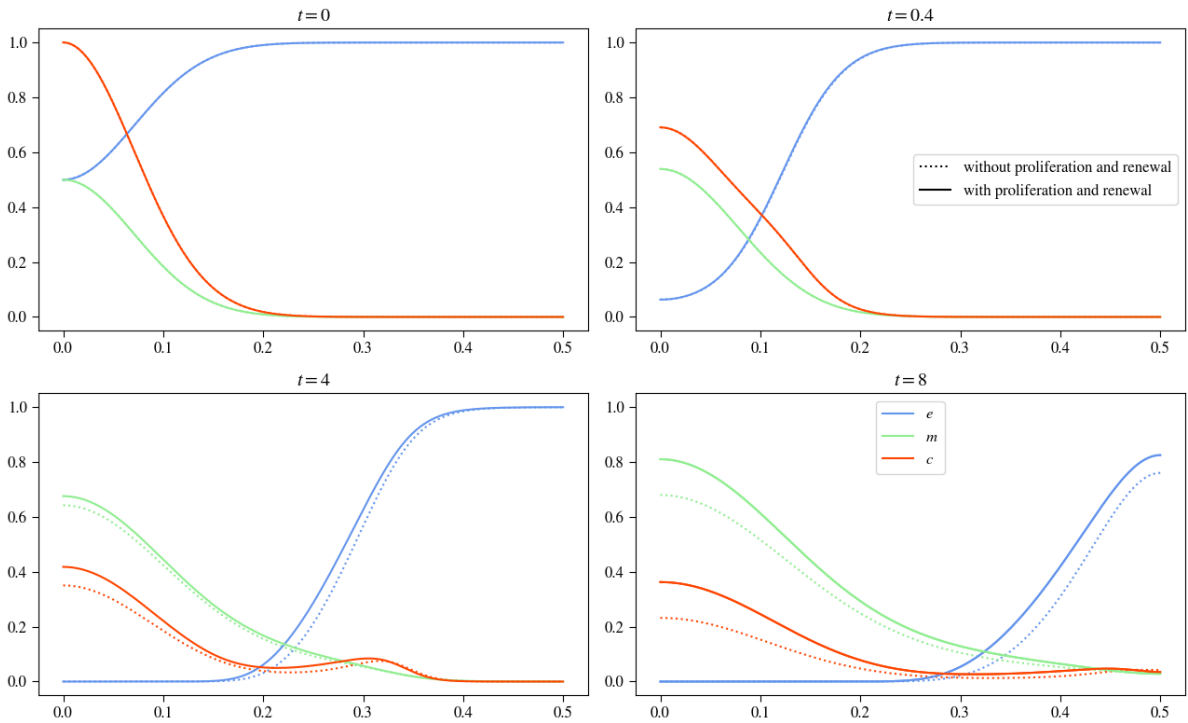


FIGURE 24. Describing the updated basecase, in the image above only the updated basecase is plotted, below it is compared to the initial basecase.

Establishing a basecase to compare the following parameter analysis results against it makes sense. In figure 24, you can see how introducing tumor cell proliferation and extracellular matrix renewal changes the simulation outcome. For this, we used the values  $\mu_1 = 0.1$  and  $\mu_2 = 0.5$  according to the only experiment found for this system of equations in the paper of Kolev et al. [13].

Comparing this new basecase to our initial model's basecase, we can see the influences of both  $\mu_1$  and  $\mu_2$ , as the tumor cell density curve is visibly higher than without proliferation, causing a higher production of matrix-degrading enzymes, which would lead to faster ECM degradation. However, this is countered by the renewal factor  $\mu_2$  causing the ECM concentration to be higher at the end, at  $t = 8$ , than in the initial basecase experiment.

### 4.3.2 Parameter Analysis

For the Parameter Analysis of the model with proliferation and renewal, we are focusing on comparing the updated model's results with those produced by the model without proliferation and renewal. This will point out again the influence of  $\mu_1$  and  $\mu_2$  on the system.

#### $d_c$ Variation

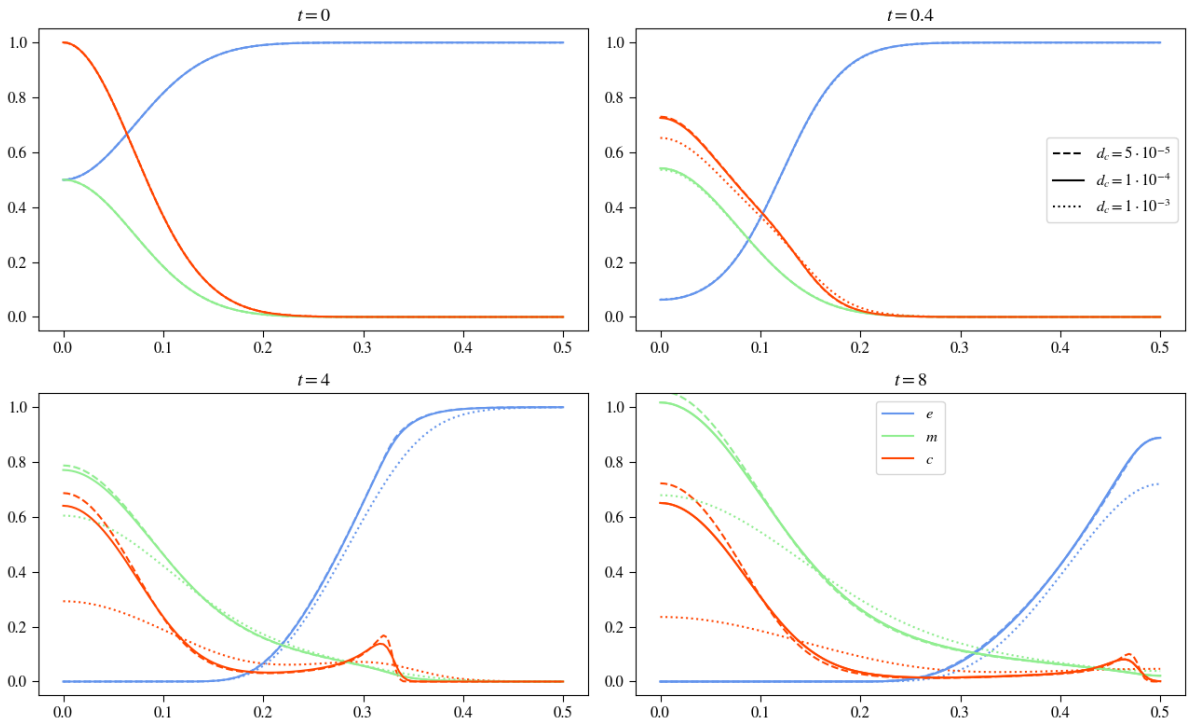


FIGURE 25. Plots show results for varying  $d_c$  whilst keeping the other parameters constant

Varying  $d_c$  with proliferation terms, we see the same effects as without proliferation. Higher values for  $d_c$  cause a more substantial influence of diffusion and a weaker one for the haptotaxis. This leads to a curve with less or no secondary lump of tumor cells invading the space separately but to a faster distribution process of tumor cells throughout space. The MDE concentration follows this behavior, depending on its production of the tumor cell density distribution in space. The ECM decays faster and the faster the tissue is invaded, the higher the diffusion factor is. Comparing them, we see little differences; only tumor cell density and ECM are raised a little in each plot due to the renewal and proliferation factors, which also causes a higher MDE concentration due to higher tumor cell densities.

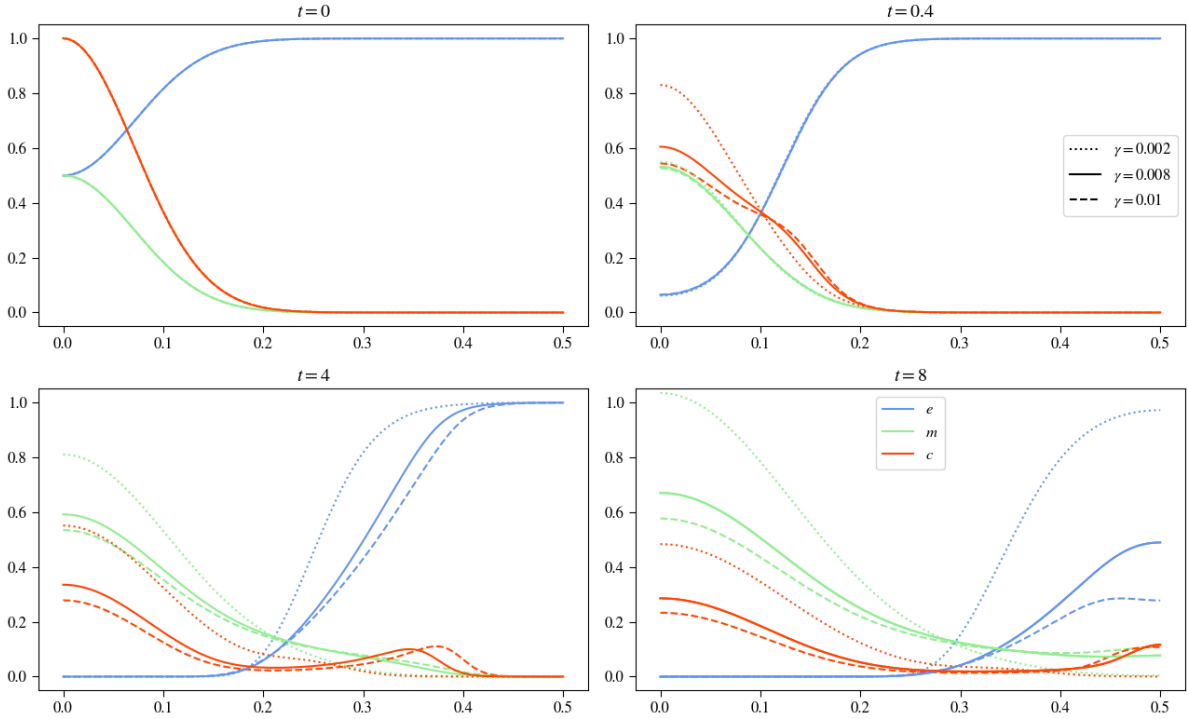


FIGURE 26. Plots show results for varying  $\gamma$  whilst keeping the other parameters constant.

### $\gamma$ Variation

When we look at  $\gamma$ , we also can see the same effects as the model without proliferation shows, with the adjustments varying  $d_c$ , with raised curves for all variables. Increasing  $\gamma$  means increasing haptotaxis effects, pulling the tumor cells stronger towards the extracellular matrix molecules. This causes a faster invasion pace and a higher density of tumor cells invading the tissue, but a lower staying at the center at  $x = 0$ . This also means that the ECM degrading process happens faster and the MDEs are more evenly distributed through space the higher  $\gamma$  is. As mentioned above, the same effects come in this experiment, introducing proliferation and renewal, with higher values for tumor cell density, MDE and ECM concentration, especially at the later points in time, clearly depictable. Interestingly, by introducing a renewal factor for the extracellular matrix, the proliferation of the tumor cells causes a faster production of matrix-degrading enzymes, which makes the system produce nearly the same results without proliferation and renewal concerning the ECM concentration. Still, introducing the renewal of the ECM results in slightly higher concentrations overall.

### $\mu_1$ Variation

The parameter  $\mu_1$  describes the proliferation of the tumor cells, using Kolev et al.'s estimate in [13] we assume an even distribution with  $\mu_1 \sim U[0.1, 1.0]$ . Introducing this factor, we can expect a faster production of the matrix-degrading enzymes and a faster degradation of the ECM with an increase in the total amount of tumor cells. Though

the effect of the faster ECM degradation might be damped by also introducing a renewal term, we can, especially for higher  $\mu_1$  values, expect to dominate the simulations and accelerate ECM degradation.

Since the basecase, using Kolve et al.'s values, sets the proliferation rate of the tumor cells to  $\mu_1 = 0.1$ ; we consider in the dotted experiment the effects of only introducing the renewal of the ECM.

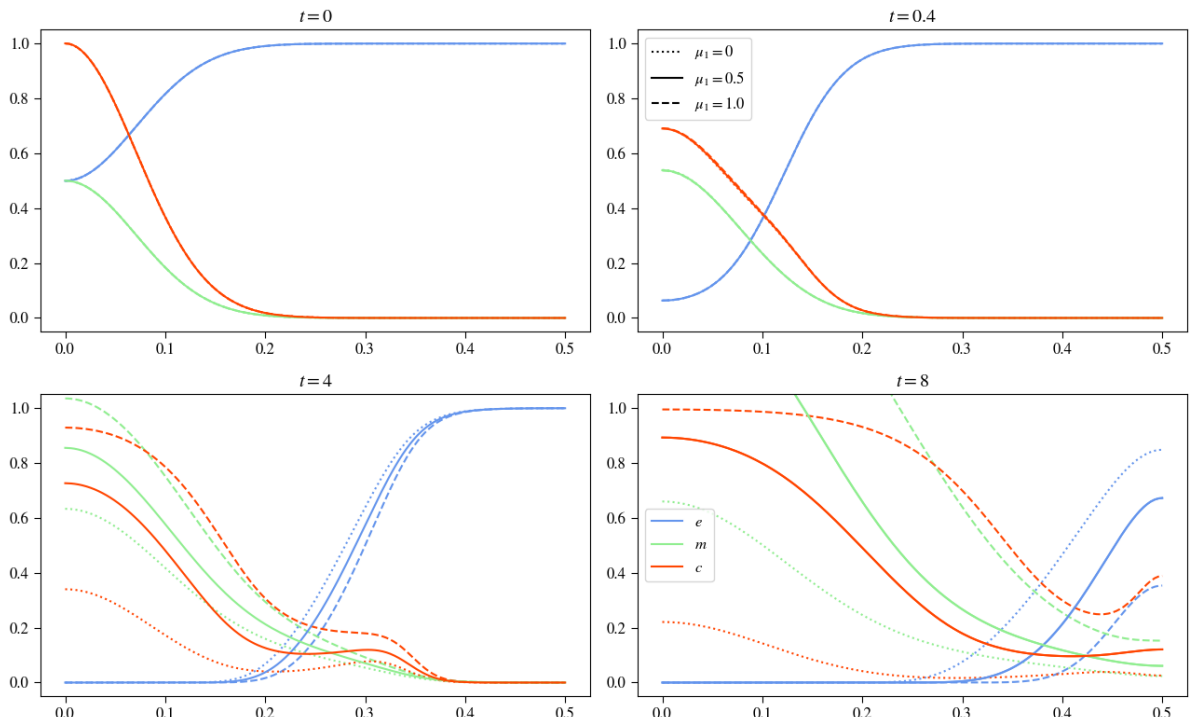


FIGURE 27. Plots show results for varying  $\mu_1$  whilst keeping the other parameters constant.

The effects of  $\mu_1$  take some time to act, as we can see no deviations for all the experiments in figure 27 at  $t = 0.4$ .

At the next point at  $t = 4$ , the differences are striking concerning all variables. With a higher proliferation factor of the tumor cells, the MDE concentration also rises clearly and the ECM degradation is accelerated. However, at this point, the deviations regarding the curve for the ECM concentration are still subtle.

It looks different at the last point in time, at  $t = 8$ . As mentioned above, an increase in total tumor cells drastically increases the MDE concentration and accelerates the ECM degradation.

Introducing  $\mu_1$  means introducing one of the hallmarks of cancer, sustaining proliferative signaling, which makes the tumor cells themselves responsible for producing them instead of requiring other hormones or enzymes to trigger growth signaling. Setting  $\mu_1 = 0$  could result from a drug working, hindering growth signaling.

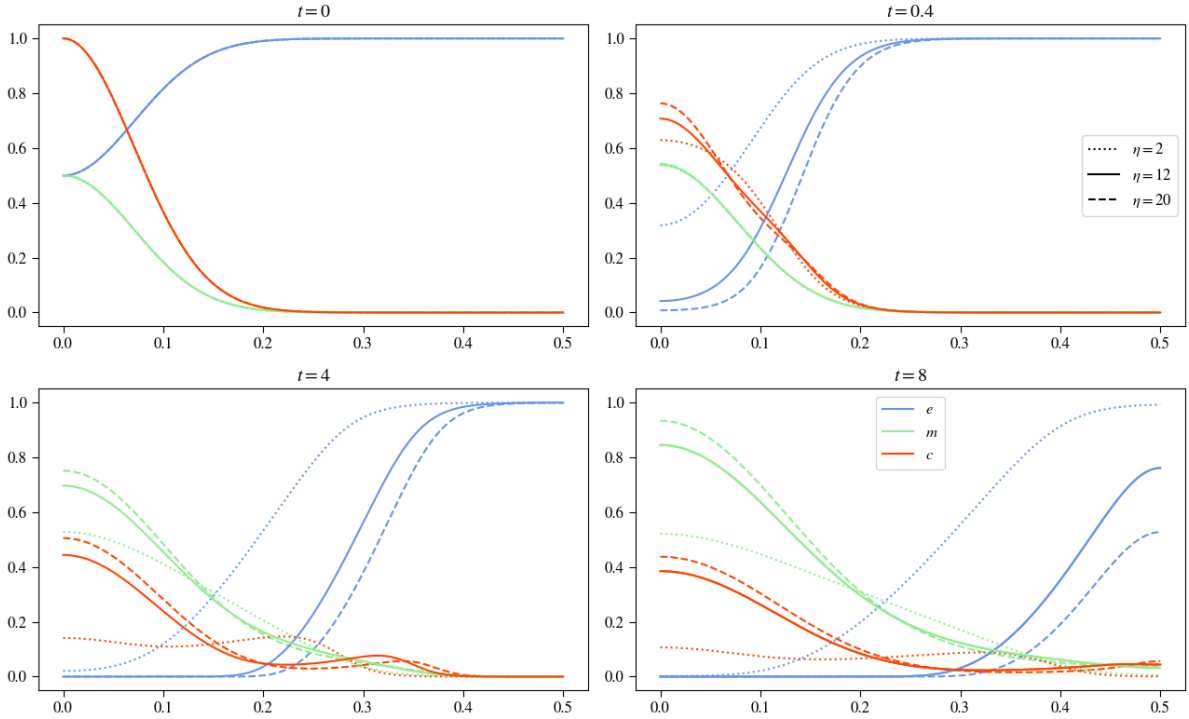


FIGURE 28. Plots show results for varying  $\eta$  whilst keeping the other parameters constant.

### $\eta$ Variation

We see mostly the same effects as we compare the  $\eta$  variation between with and without proliferation and renewal models. We see little for the solid and dashed curves, though the new model's curves are all slightly raised. Looking at  $\eta = 0$ , we see some interesting deviations. At the time point  $t = 0.4$ , the plots still look somewhat similar, but looking at  $t = 4$ , we see that the curve of the tumor cells has a more even distribution along the x-axis and also its maximum is visibly lower with the value of about 0.2 at  $x = 1.4$  instead of 0.25 at  $x = 1.3$ . This behavior is due to the renewal of the ECM, where without proliferation, this curve stayed constant throughout the experiment. Here, it can increase, which it does, altering the slope of the curve and, therefore, influencing the haptotactic pull for the tumor cells. Additionally, the other two experiments showed a visible increase in the tumor cell density and the matrix-degrading enzyme concentration. However, there is only a slight increase in the ECM concentration; here, we can see no increase in area for the tumor cell density. The renewal of the ECM counters the proliferation of the tumor cells and the slowed ECM degrading process. The ECM has visibly increased at the last two points, with both curves of ECM and tumor cells almost mirroring each other. Summing up the areas of both variables, they occupy the space wholly needed for the logistical growth terms. This means that proliferation and renewal will play no more critical role in continuing this experiment as they have reached an equilibrium state and cancel each other out.

### $\mu_2$ Variation

The parameter  $\mu_2$  describes the renewal processes of the extracellular matrix molecules. Also, using Kolev et al.'s estimate in [13], we can assume an even distribution with  $\mu_2 \sim U[0.1, 1.0]$ . In natural processes, the renewal of the extracellular matrix is essential to regulate cell differentiation and wound repair, for example [14]. As we see in figure 29,

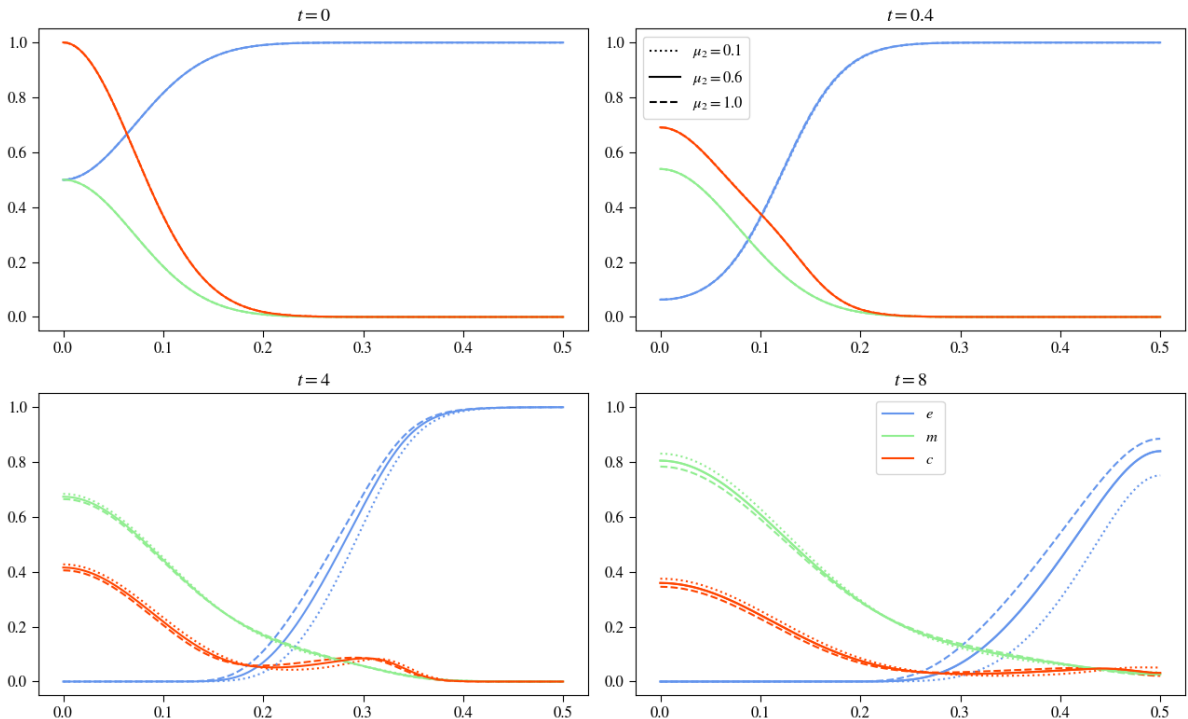


FIGURE 29. Plots show results for varying  $\mu_2$  whilst keeping the other parameters constant.

this renewal process takes some time to show effects. There are no deviations for the experiments after  $t = 0.4$ .

The differences are still subtle when looking at the results later at  $t = 4$ . Increasing  $\mu_2$  slows down the extracellular matrix degradation process and affects the motility of the tumor cells, pulling less of them outwards into the surrounding tissue. The MDE concentration nearly overlays completely, with a minimal higher concentration for the lowest  $\mu_2$  experiment at the origin due to the slightly higher tumor cell density in this area.

The differences at the last point in time at  $t = 8$  are still minor compared to varying the other parameters. As mentioned before, the slowed ECM degradation diminishes haptotactic effects, which leads to slightly less matrix-degrading enzyme concentration at the origin.

Varying the ECM renewal rate at this magnitude shows little influence on the resulting simulations.

At this point, it is important to say that the renewal rates for tissue in the human body strongly vary. Comparing, for example, bone tissue with connective tissue, we see

these processes at highly different time scales. Because we did not have experimental data for this parameter, we only investigated Kolev et al.'s estimates, but considering realistic cases, it is important to have better measures for this parameter.

### $d_m$ Variation

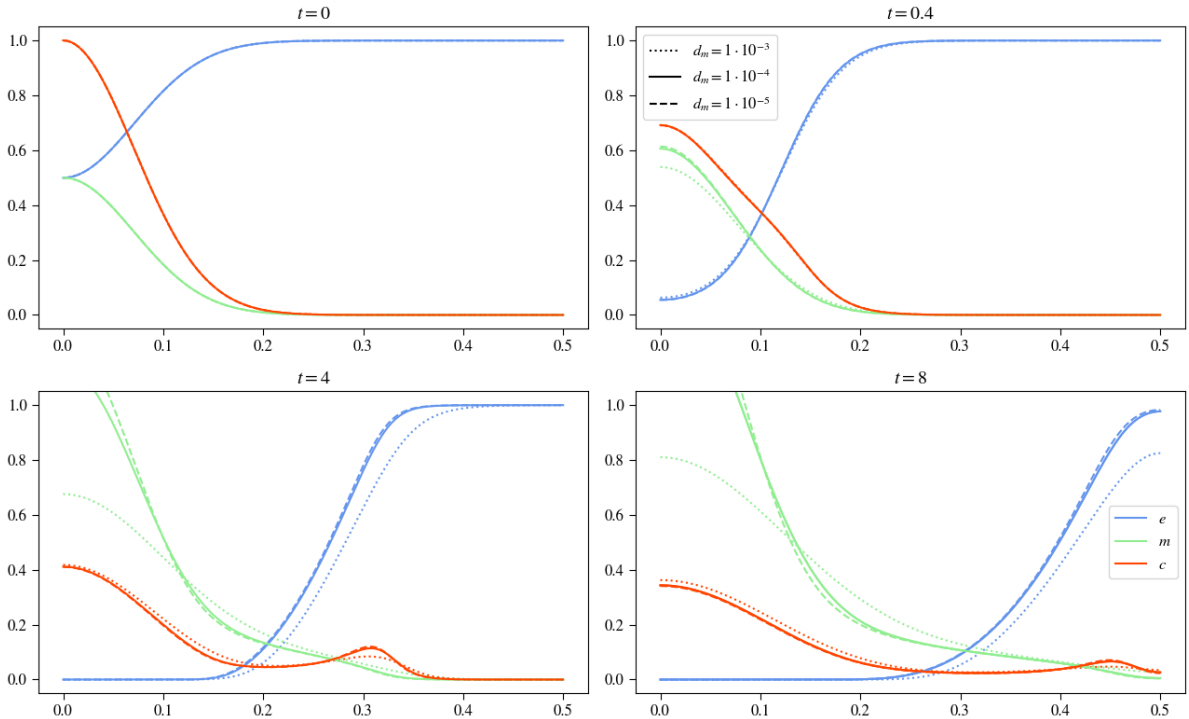


FIGURE 30. Plots show results for varying  $d_m$  whilst keeping the other parameters constant.

Comparing the results varying the diffusion factor of the matrix-degrading enzymes does, as before, yield only minor differences between the initial and updated model. As observed before, the tumor cell density curve and the MDE concentration curves are slightly raised due to the proliferation of the tumor cells. However, the ECM curves for the two lower values of varying  $d_m$  seem to be subject only to minimal change. We can see that it is raised only for very high values of  $d_m$  compared to the model without renewal. The other two curves take off at the same point along the x-axis and finish at the same values for their ECM concentration. Looking at the tumor cell density curves for those  $d_m$  values, we see that towards  $x = 0.5$ , they do not describe an as steep bump as the initial model. This also causes a little less MDE concentration, which is responsible for the seemingly unchanged behavior of the extracellular matrix concentration.

### $\alpha$ Variation

Taking a look at comparing the  $\alpha$ -variation yields more exciting results since  $\mu_1$  acts as a secondary MDE production effect by producing tumor cells, which in turn produce

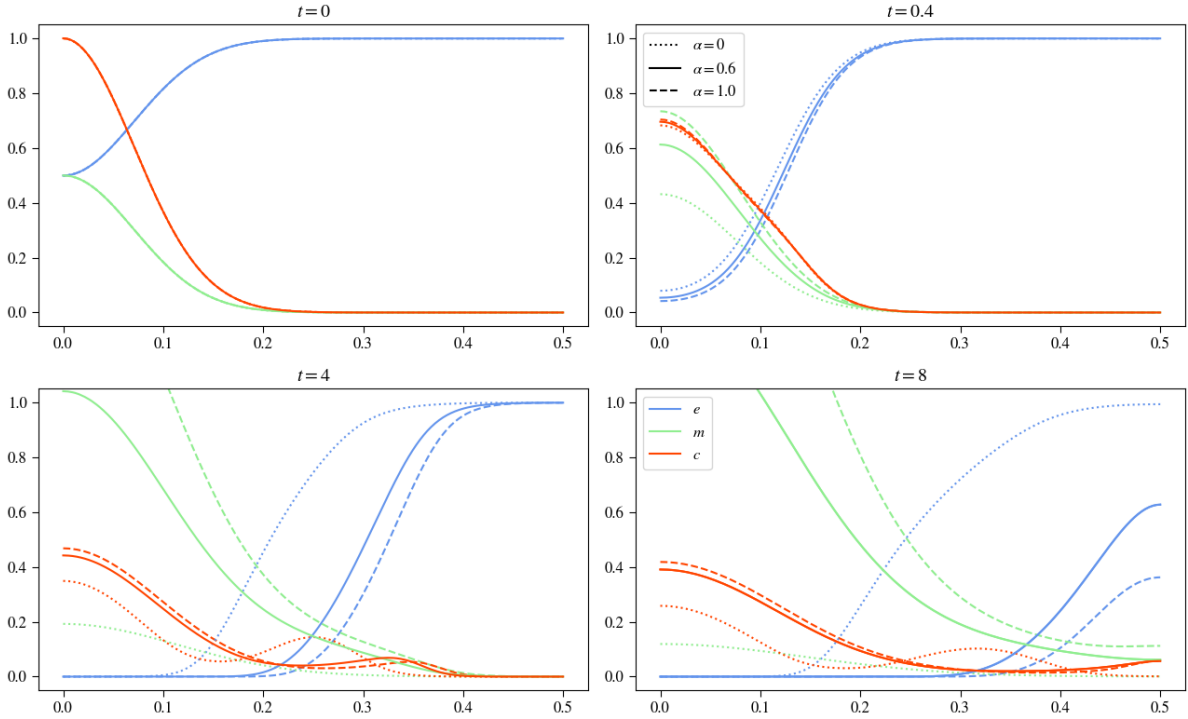


FIGURE 31. Plots show results for varying  $\alpha$  whilst keeping the other parameters constant.

the matrix-degrading enzymes. Though the overall shape and effects to be observed are the same, after  $t = 4$ , the model with proliferation exceeds one at the origin for the MDE concentration for the two higher  $\alpha$  experiments; in contrast, for the model without proliferation, only the one with the highest  $\alpha$  value did so. The tumor cell density curve is slightly raised, which allows the MDE concentration. However, the higher values for the MDEs leave the ECM degrading process untouched, with no visible difference between the initial model and the updated one.

### $\beta$ Variation

Considering  $\beta$ , we can expect that with the introduction of  $\mu_2$ , the ECM degradation will be slowed considerably with rising  $\beta$  since this reduces the MDE concentration and renews the ECM. Looking at the plots, we can see this behavior in the dotted line, which shows the experiment results for the highest  $\beta$  value of 0.1. Though even at the end, it has an overall area that is slightly less than the initial condition, we can see going from timestep  $t = 0.4$  to  $t = 4$  that MDE decay and ECM renewal were sufficiently strong to restore the ECM and going from  $t = 4$  to  $t = 8$  we see this behavior again, renewing the ECM. The other two experiments for  $\beta$  showed no effects as strong as with  $\beta = 0.1$ . However, we can still see the effects of proliferation and renewal especially clear in the solid line,  $\beta = 0.01$ , at the last point in time, where we can observe a visible increase of both ECM and tumor cell density. In this experiment, we see that  $\beta$  is a little too low to counter the effects of ECM degradation; going from  $t = 4$  to  $t = 8$ , we see an apparent

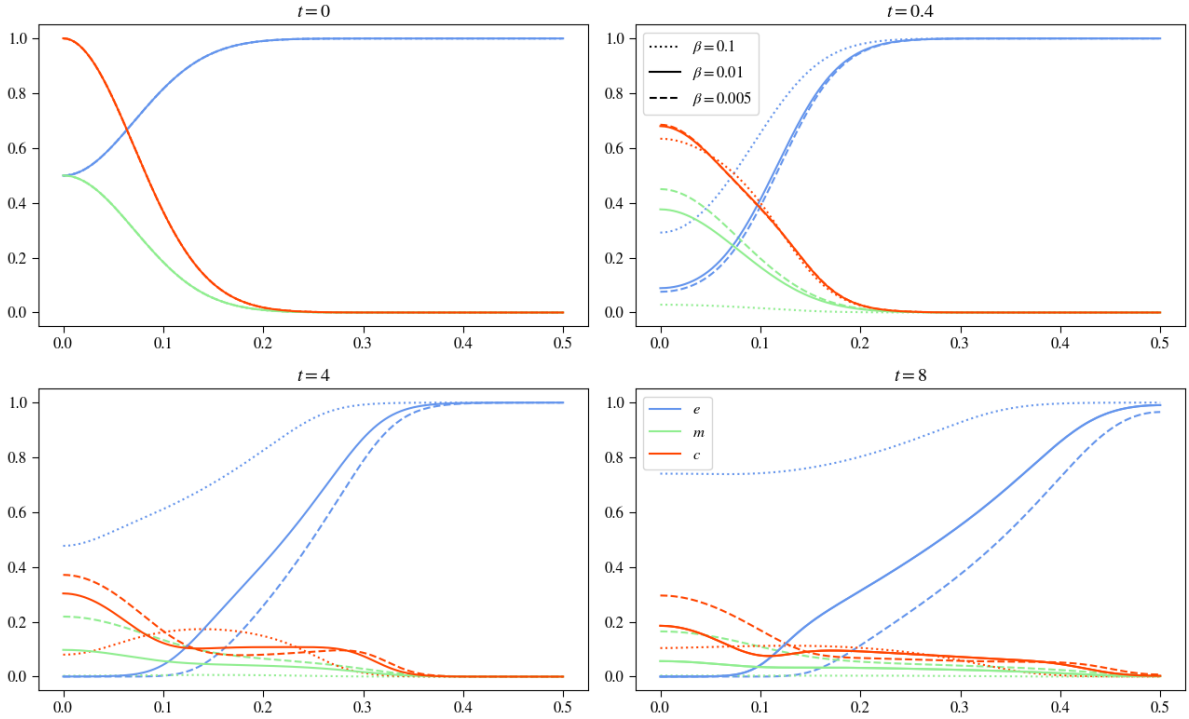


FIGURE 32. Plots show results for varying  $\beta$  whilst keeping the other parameters constant.

decline of ECM concentration, though it is not as striking as for  $\beta = 0.005$ .

## Cross-Variation

### $\mu_1 - \mu_2$ Variation

The effects of observing this cross variation take some time, as did the separate variations of both  $\mu_1$  and  $\mu_2$ . For both  $\mu_1 = \mu_2 = 0.1$ , we see that slower ECM renewal and slower tumor cell proliferation increase the degrading process of the extracellular matrix and, with this, affect the haptotaxis effect to increase slightly. At the center, a lump remains that has a maximum higher than for the experiment with  $\mu_2 = 1.0$  and the invasion of the tissue has proceeded faster. Increasing  $\mu_2$ , as previously mentioned, results in slower ECM degradation due to the increased renewal term and therefore, the tumor cells are stretched out more evenly along the x-axis. Looking at the results when increasing  $\mu_1$ , we also see the effects only after  $t = 4$ . For  $\mu_2 = 0.1$ , we see that the tumor cell density at  $x = 0$  is slightly more significant and at  $x \approx 2.9$ , the curve for  $\mu_2 = 0.1$  is also slightly larger, being a little below  $\mu_2 = 1.0$  in between  $x \approx 0.2$  and  $x \approx 0.29$ . The curve for the MDEs looks very similar in both cases for  $\mu_2$  due to the very similar tumor cell density curve,  $c$ , though the eECM has visibly faster degraded for  $\mu_2 = 0.1$  due to the slower renewal.

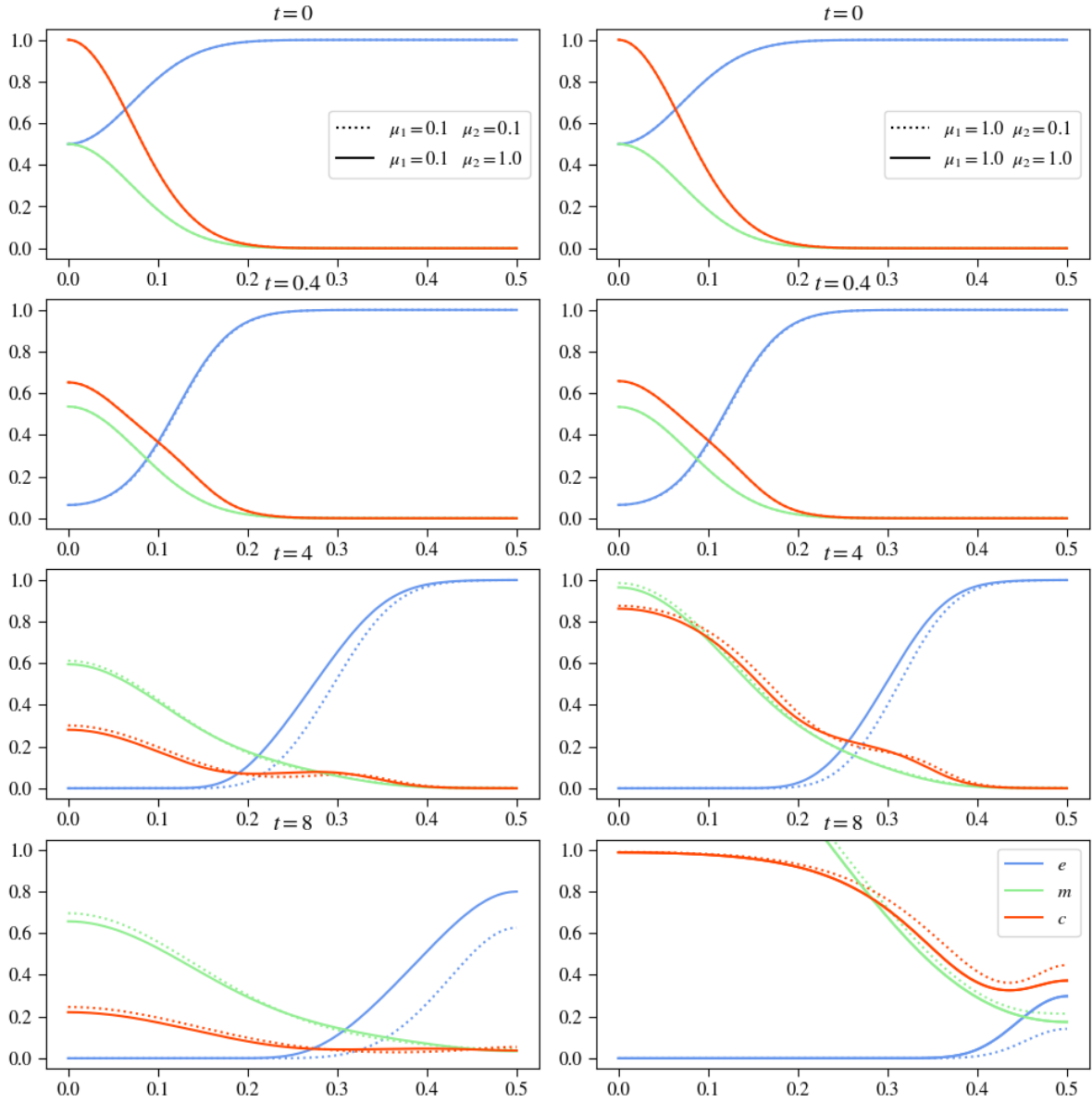


FIGURE 33. Plots show results for varying both  $\mu_1$  and  $\mu_2$  whilst keeping the other parameters constant.

### $d_c - \gamma - \mu_1$ Variation

First we are going to take a look at how changing  $\gamma$  and  $\mu_1$  affects the system whilst having lower diffusion values for the tumor cells as the basecase value with  $d_c = 5 \cdot 10^{-5}$ . Figure 34 shows the results of these experiments. Inspecting the dotted curves on the left side column, shows the results for all parameters set to their lower values, we see no division of the tumor cells, indicating that the haptotactic coefficient was set too low. Because of the high concentration of tumor cells near the origin, we see also high values for the MDE concentration. Due to the decreased motility of the tumor cells we see that the ECM is also slower degraded compared to the other parameters, even when  $\mu_1$  was

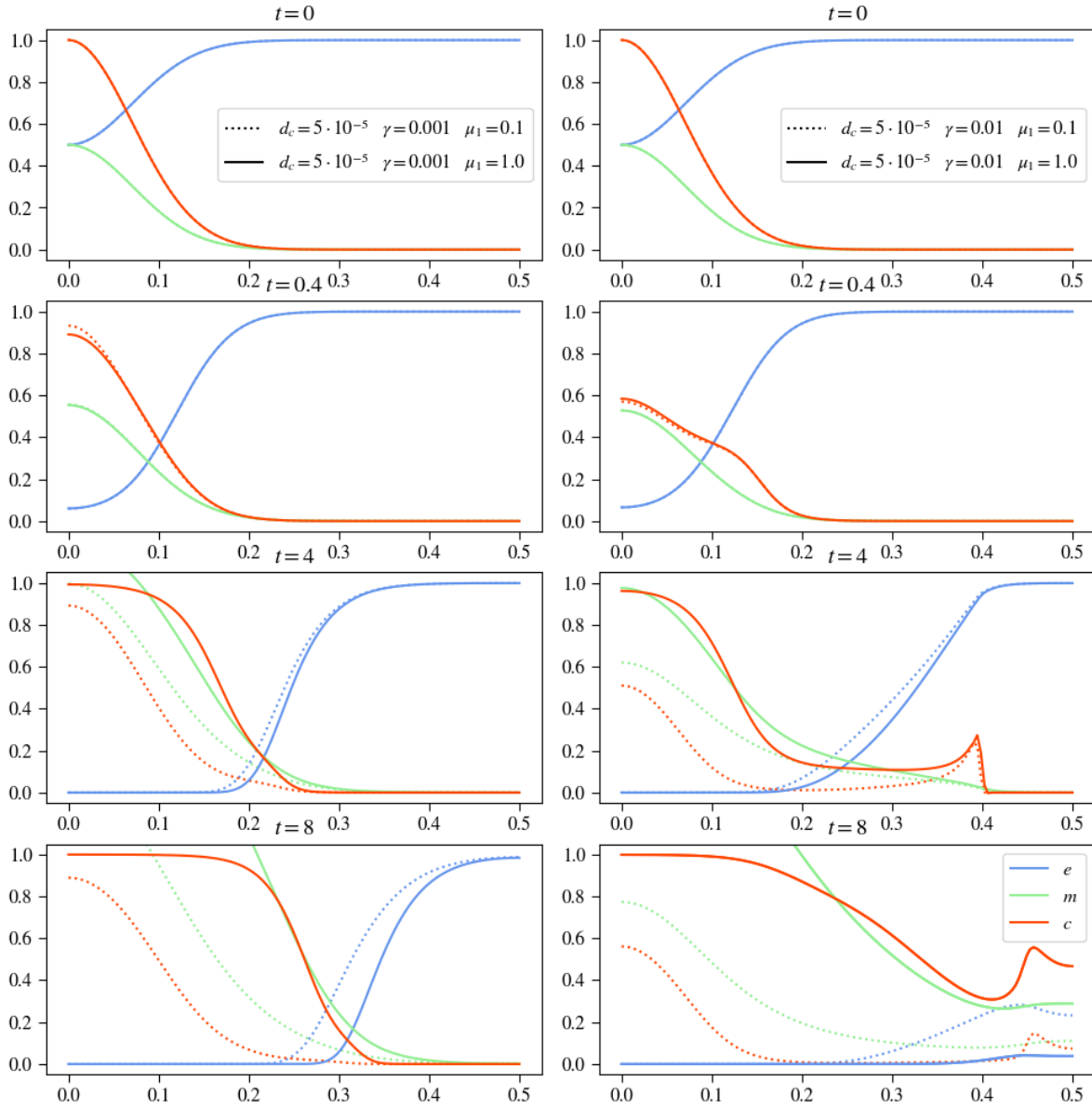


FIGURE 34. Plots show results for varying both  $d_c$ ,  $\gamma$  and  $\mu_1$  whilst keeping the other parameters constant. This plot is the first of two, with the same  $d_c$  value for every plot in this figure.

increased.

The right side of the same figure shows a clear division of tumor cells detaching from the main lump and invading faster. The split off lump of cells even forms for high  $\mu_1$  values a pregnant peak at  $t = 4$ . For both high and low values of  $\mu_1$  we see a considerably faster ECM degradation, though at the last point in time we see that with high proliferation and growth factor on the tumor cells the MDE production is distinctly higher which causes faster ECM degradation.

Taking a look at figure 35, showing results for increased diffusion on the tumor cells, we see no division on the tumor cell density curve with a low  $\gamma$  value. Regarding the two experiments with low  $\gamma$  and high  $\mu_1$  values we see the effect of diffusion clearly on

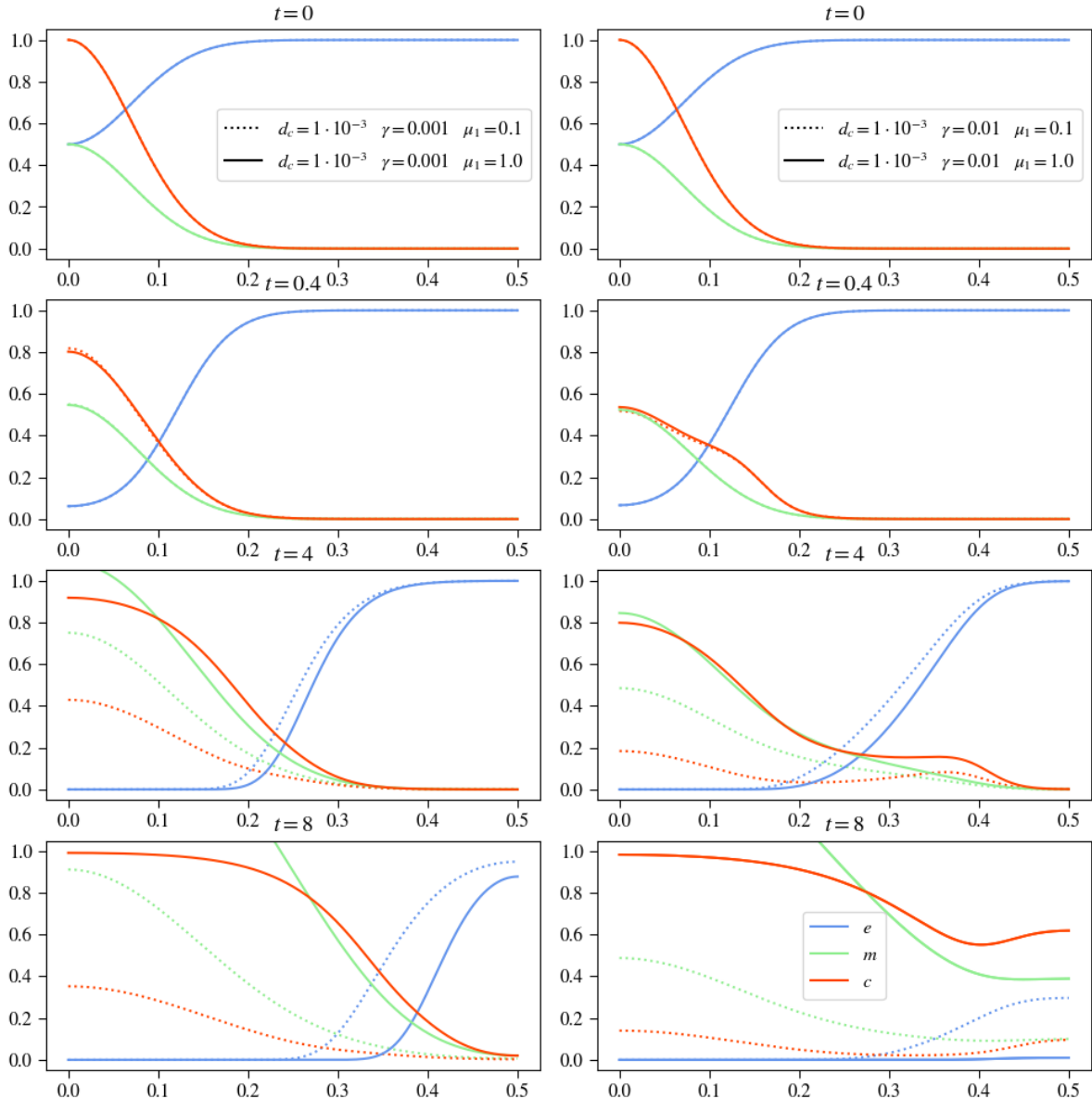


FIGURE 35. Plots show results for varying both  $d_c$ ,  $\gamma$  and  $\mu_1$  whilst keeping the other parameters constant. This plot is the second of two, with the same  $d_c$  value for every plot in this figure.

the tumor cells. Though the red curves look the same in shape the one with the higher diffusion value looks shifted to the right, the same goes for the MDE concentration, resulting in faster ECM degradation. Comparing the results of increasing  $\mu_1$  between diffusion values, we can also observe the effect of diffusion clearly. The curve for the tumor cells is more evenly distributed, which causes the same behavior in the MDE curve and also a lower concentration and density in total in this regard.

The right column of figure 35 shows results for increasing both  $d_c$  and  $\gamma$ . Here we see that the division forming on the tumor cells clearly, though not as pregnant as with low diffusive values. The fast and even spread of the tumor cells as well as their proliferation term, results in the highest rate of ECM degradation.

This variation is an excellent example of the relationship between haptotaxis and diffusion. We see with low values on the diffusion coefficient the changes in the haptotactic coefficient more striking, with a pointy peak in the tumor cells curve, but with also high diffusion values the peak is not as pregnant and not as striking.

Another observation is the influence of the logistical growth rate of the tumor cells. In the last experiment we see that with a fast and even spread we get the highest overall density of tumor cells and the fastest ECM degradation.

### $\eta - \mu_2$ Variation

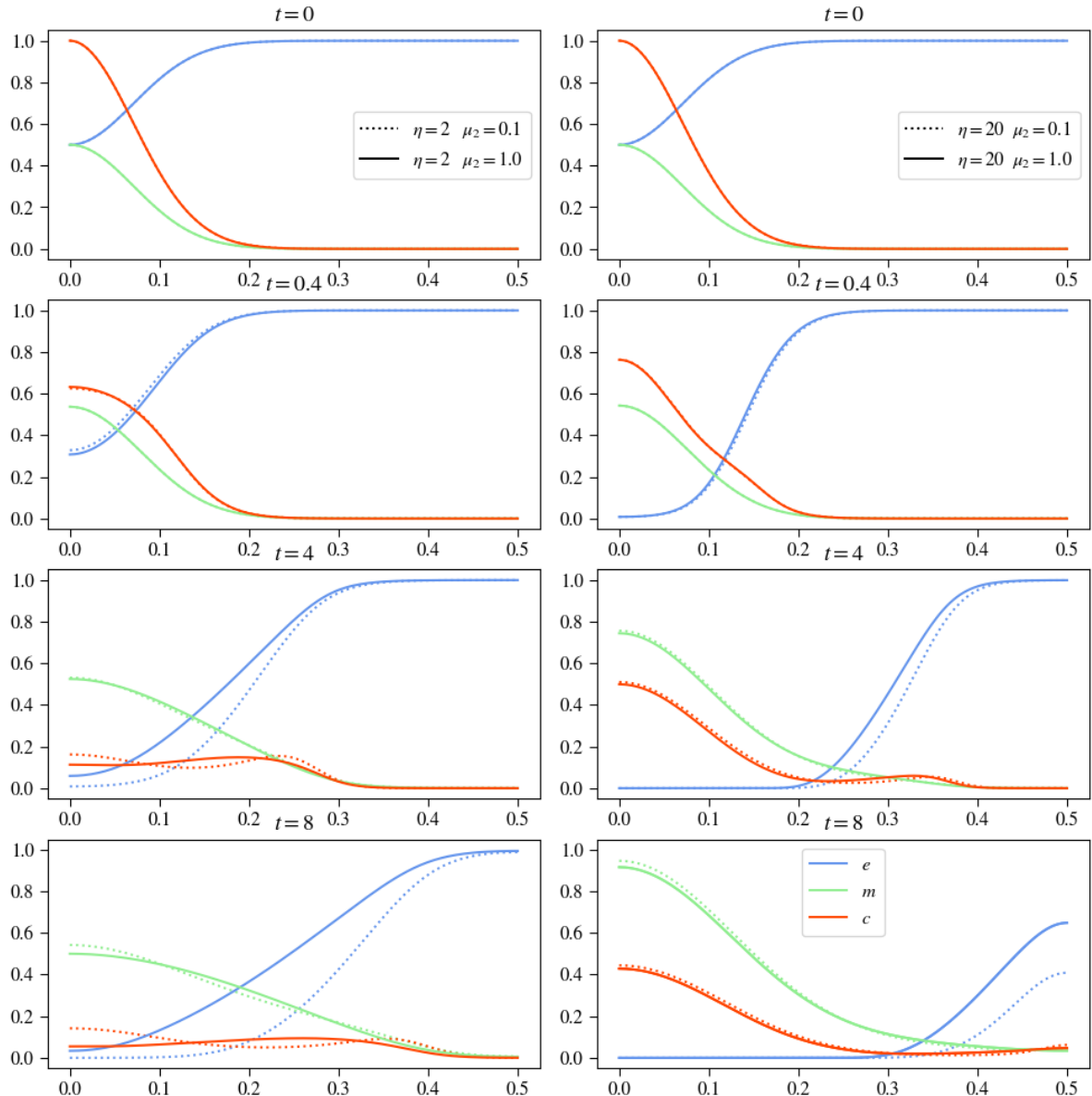


FIGURE 36. Plots show results for varying both  $\eta$  and  $\mu_2$  whilst keeping the other parameters constant.

Varying both  $\eta$  and  $\mu_2$ , we can expect apparent changes in the curve describing the ECM concentration. On the left side of figure 36, we see the two experiments for low  $\eta$  values and that increasing  $\mu_1$  has only a little effect. Where we could have expected to see an increase of the ECM, maybe even, we see that the ECM curves for both experiments verify that the renewal factor  $\mu_2$  was too low to counter the ECM degradation, even with a low degradation factor. Still, between  $\mu_2 = 0.1$  and  $\mu_2 = 1.0$ , there are visible differences in the degrading speed of the ECM. We can also observe that with the higher renewal term, the tumor cell density curve receives more of an effect of haptotaxis, resulting in a more stretched curve with only one long lump of tumor cells, whereas, for the lower renewal factor, we can still clearly see that there is a division that invades the tissue and one that stays at the origin. Concerning the MDE curves, we see little difference for higher  $\mu_2$ , which means more stretched tumor cell density. We can also observe a more stretched MDE curve with a lower maximum at the origin.

Looking at the experiments with raised  $\eta$  to accelerate the ECM degrading process, we can only see pregnant differences in the curve describing the ECM concentration; the other two look across the steps in time to be widely similar. For the ECM curve, we see that the experiment with the lower  $\mu_2$  value results in a faster degradation process.

#### 4.4 2D Results with Proliferation and Renewal - Heterogenous ECM

Figure 37 you can see the effects using a heterogenous extracellular matrix structure. The plots show the tumor cell density in the left column, the extracellular matrix concentration in the middle column and the matrix-degrading enzyme concentration on the right. For the parameters, we assumed the basecase studying the model with proliferation.

This experiment describes a more realistic biological scenario, as seen in figure ??, where the tumor cells are located at the basement membrane of neighboring tissue and have degraded this membrane to invade the surrounding tissue and degrade the extracellular matrix.

The first image in the middle column shows the experiment's initial distribution. The extracellular matrix molecule concentration is only on the right side of the plot, which indicates that neighboring tissue has been invaded. In the center of the same image, there is a hollow spot where the tumor cells of the initial distribution are located.

After four time steps, the ECM slowly degrades and the tumor cells are pulled further into the neighboring tissue by haptotaxis caused by the ECM structure. The concentration of the matrix-degrading enzymes shows little difference in their behavior compared to the experiments done with a homogenous ECM. They only depend indirectly on the ECM by being produced where the tumor cells are being pulled by the extracellular matrix concentration.

The next point shows increased ECM degradation, further invading tumor cells into the tissue. The tumor cells behave as a semicircular wave moving into the direction of the ECM; the primary lump remains at the center, with the edge of having containing a smaller amount of cells moving outwards. The image describing the matrix-degrading enzyme concentration still shows only minor effects; looking closely, we can see that from

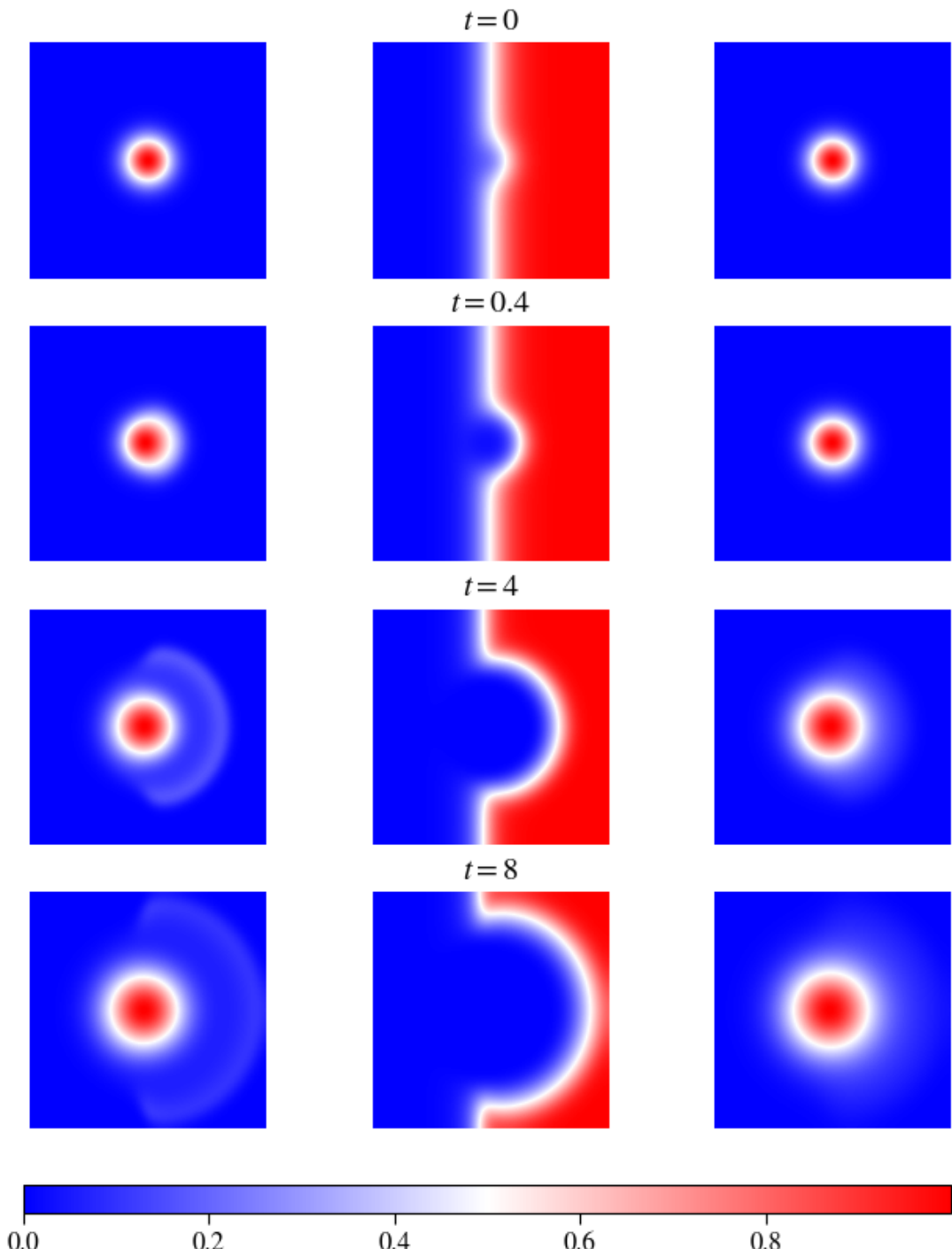


FIGURE 37. 2D Results using a heterogenous ECM with the parameter values:  $d_c = 5 \cdot 10^{-4}$ ,  $\gamma = 0.0055$ ,  $\mu_1 = 0.1$ ,  $\eta = 10$ ,  $\mu_2 = 0.5$ ,  $d_m = 1 \cdot 10^{-3}$ ,  $\alpha = 0.3564$ ,  $\beta = 0$ ; left: tumor cell density, middle: ECM concentration, right: MDE concentration.

the center moving to the right, there is a slightly higher concentration than in the other direction.

The last row depicts the experiment after  $t = 8$  timesteps and the above mentioned effects are propagated. The ECM degradation has continued, as has the invasion of the tumor cells. The wave moving in the direction of the remaining extracellular matrix molecules has spread through space, decreasing its strength. The MDEs still show little influence on the heterogeneous ECM structure, with the primary lump staying centered and only making it difficult to recognize more concentration towards the movement of the tumor cells and concentration of the extracellular matrix.

Considering different extracellular matrix molecules and structures or physical influences such as heat or radiation, the ECM structure and the system's behavior can be adjusted to simulate better real-life scenarios of cancer invading tissue.

## 5 Conclusion and Discussion

In this study, we conducted a parameter analysis for a numerical model in tumor development research. The aim was to investigate the impact of varying model parameters, dimensions and the structure of the extracellular matrix on simulation outcomes and provide insights into the model’s behavior under different conditions. This was done to facilitate implementing this model in a real-world biological or medical scenarios, facilitating an entry point for researchers to choose the parameters for their experiments.

The results of our analysis highlight the model’s sensitivity to changes in key parameters, such as the diffusion or haptotaxis coefficients on the tumor cells, the proliferation and growth rate of the tumor cells, the degradation rate of the extracellular matrix and the production and decay rates for the matrix-degrading enzymes. We observed that small variations in these parameters can lead to significant differences in simulation outcomes, indicating the importance of carefully selecting and calibrating model parameters for accurate representation of biological phenomena.

Furthermore, our study revealed the complex interplay between different parameters during cross-variating them and their effects on tumor invasion and extracellular matrix degradation. For example, we found that increasing the haptotactic flux coefficient of the tumor cells is the critical component to controlling how many cells of a tumor invade the surrounding tissue and how many stay at the center of the simulation causing drastic changes in the rate at which the extracellular matrix is degraded. Most of the parameters studied increased the invasion pace of the tumor cells and the degradation of the extracellular matrix, when they were increased.

Moreover, the influence of spatial dimensions on simulation outcomes was shown, with simulations in higher dimensions producing qualitatively different results compared to the initial one-dimensional model. This underscores the importance of considering the spatial complexity of biological systems when designing computational models for tumor development.

Additionally, our analysis sheds light on the limitations of the current model and areas for further improvement. The model could be extended in both discrete and continuous ways, for example, by implementing biochemical effects like chemotaxis, which is done in Kolev et al.’s work [13] or implement immune cell interactions or heterogeneity in tumor cell populations, to capture the complexity of the tumor microenvironment better.

In conclusion, our parameter analysis provides valuable insights into the behavior of the investigated numerical model in tumor development research and underscores the importance of parameter selection and model validation in Mathematical Oncology. By refining our understanding of the underlying mechanisms driving tumor progression, such models have the potential to inform therapeutic strategies and improve patient outcomes in the fight against cancer.

## References

1. Anderson, A. Continuous and Discrete Mathematical Models of Tumor-induced Angiogenesis. en. *Bulletin of Mathematical Biology* **60**, 857–899. ISSN: 00928240. <http://link.springer.com/10.1006/bulm.1998.0042> (2023) (Sept. 1998).
2. Anderson, A. R. A., Chaplain, M. A. J., Newman, E. L., Steele, R. J. C. & Thompson, A. M. Mathematical Modelling of Tumour Invasion and Metastasis. en. *Journal of Theoretical Medicine* **2**, 129–154. ISSN: 1027-3662, 1607-8578. <http://www.hindawi.com/journals/cmmm/2000/490902/abs/> (2023) (2000).
3. Bekisz, S. & Geris, L. Cancer modeling: From mechanistic to data-driven approaches, and from fundamental insights to clinical applications. *Journal of Computational Science* **46**. 20 years of computational science, 101198. ISSN: 1877-7503. <https://www.sciencedirect.com/science/article/pii/S1877750320304993> (2020).
4. Hanahan, D. Hallmarks of Cancer: New Dimensions. *Cancer Discovery* **12**, 31–46. ISSN: 2159-8274. eprint: <https://aacrjournals.org/cancerdiscovery/article-pdf/12/1/31/3052722/31.pdf>. <https://doi.org/10.1158/2159-8290.CD-21-1059> (Jan. 2022).
5. Franssen, L. C., Lorenzi, T., Burgess, A. E. F. & Chaplain, M. A. J. A Mathematical Framework for Modelling the Metastatic Spread of Cancer. en. *Bulletin of Mathematical Biology* **81**, 1965–2010. ISSN: 0092-8240, 1522-9602. <http://link.springer.com/10.1007/s11538-019-00597-x> (2023) (June 2019).
6. Yin, A., Moes, D. J. A. R., van Hasselt, J. G. C., Swen, J. J. & Guchelaar, H.-J. A review of mathematical models for tumor dynamics and treatment resistance evolution of solid tumors. en. *CPT Pharmacometrics Syst. Pharmacol.* **8**, 720–737 (Oct. 2019).
7. Chaplain, M., Lolas, G. & ,The SIMBIOS Centre, Division of Mathematics, University of Dundee, Dundee DD1 4HN. Mathematical modelling of cancer invasion of tissue: dynamic heterogeneity. en. *Networks & Heterogeneous Media* **1**, 399–439. ISSN: 1556-181X. <http://aimsciences.org//article/doi/10.3934/nhm.2006.1.399> (2023) (2006).
8. Merino-Casallo, F., Gomez-Benito, M. J., Hervas-Raluy, S. & Garcia-Aznar, J. M. Unravelling cell migration: defining movement from the cell surface. en. *Cell Adh. Migr.* **16**, 25–64 (Dec. 2022).
9. Klawonn, A., Köhler, M. & Rheinbach, O. HiFlow3 – A Hardware-Oriented Parallel Finite Element Package. *Innovative Parallel Computing, Proceedings of the 5th International Conference ParCo2001* **13**, 213–220 (2002).
10. Kitware, Inc. *ParaView* <https://www.paraview.org/>. Zuletzt abgerufen am [Datum]. 2022.

11. Stéphanou, A., McDougall, S., Anderson, A. & Chaplain, M. Mathematical modelling of the influence of blood rheological properties upon adaptative tumour-induced angiogenesis. *Mathematical and Computer Modelling* **44**. Advances in Business Modeling and Decision Technologies [pp. 1-95], 96–123. ISSN: 0895-7177. <https://www.sciencedirect.com/science/article/pii/S0895717705004565> (2006).
12. Aznavorian, S., Stracke, M., Krutzsch, H., Schiffmann, E. & Liotta, L. Signal transduction for chemotaxis and haptotaxis by matrix molecules in tumor cells. *The Journal of cell biology* **110**, 1427–38 (May 1990).
13. Kolev, M. & Zubik-Kowal, B. Numerical Solutions for a Model of Tissue Invasion and Migration of Tumour Cells. *Computational and Mathematical Methods in Medicine* **2011**, 452320. ISSN: 1748-670X. <https://doi.org/10.1155/2011/452320> (Dec. 2010).
14. Lu, P., Takai, K., Weaver, V. M. & Werb, Z. Extracellular matrix degradation and remodeling in development and disease. en. *Cold Spring Harb. Perspect. Biol.* **3**, a005058–a005058 (Dec. 2011).