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Modeling of Anticancer Drug Delivery By Temperature-Sensitive Liposomes

Vera Franziska Loeser
University of Wisconsin-Milwaukee

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MODELING OF ANTICANCER DRUG DELIVERY BY TEMPERATURE-SENSITIVE LIPOSOMES

by

Vera Franziska Loeser

A Thesis Submitted in Partial Fulfillment of the Requirements for the Degree of

MASTER OF SCIENCE in MATHEMATICS

at

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May 2017
Cytotoxic anticancer drugs are used to treat cancer, particularly tumors. These drugs themselves do not distinguish between healthy and tumor cells and attack all of them. Consequently physicians and chemists investigate safer ways of delivery that minimize damage to healthy cells. One of these ways are liposomal formulations of the anticancer drugs. Liposomes are vesicles that encapsulate the drug to shield the healthy parts of the body from the toxicity of the drugs. Due to the abnormal structure of tumors, especially their leaky vasculature, these macromolecules are able to diffuse into the tumor tissue whereas the normal vasculature prevents them to move into the tissue. After reaching the diseased tissue, the liposomes release their cargo triggered by a physical or chemical process.

In this thesis, we will investigate those liposomes that are activated upon entering regions of increased tissue temperature. Developing temperature-sensitive liposomes for targeted anticancer drug delivery is difficult to realize. The liposomes must be able to move far into the tumor against the interstitial pressure. The release of the drug must be triggered precisely in the tumor. We will show that the drug can move further into the tissue by diffusion.

The aim of the mathematical modeling is to improve our understanding of the release and transport processes involved and (in the future) to optimize the scheduling between tissue heating and the liposome administration.
To my family.

Thank you for all your support.

I love you.
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Chapter 1

Introduction

The field of modeling and simulating tumor growth and tumor treatment is sometimes called "Mathematical Oncology" and is a growing field combining mathematical sciences and biomedicine. It provides tools for quantification of key parameters from patient-specific clinical data as well as customization of cancer treatment using mathematical models (Kuang et al., 2016). For a better understanding of mathematical oncology a brief definition of cancer follows: Tumors or neoplasia can be seen as a kind of loss of tissue stability due to mutation, where the proliferation of cells is “uncontrolled”. Cancer is a malignant tumor, so that the tumor has the ability to invade the surrounded tissue and to build metastases. Metastasis describes the ability of a tumor to expand to other parts of the body (Kuang et al., 2016).

Mathematical modeling in general is a great tool for research and industry. The two major advantages of modeling are reduced costs and more precise results than those obtained by physical experiments. Therefore, mathematical modeling complements experimental work. Particular actions within a complex process can be simulated individually for a better understanding. In cancer research mathematical modeling of the medical treatment of tumor cells is a step to reduce the number of experiments in the petri dish or with animals. The drug and the tumor cells need to be developed first and then implanted into an animal. This procedure is time consuming and costly and does not guarantee that the setup of the experiments will succeed. Besides the economical point of view, also ethical issues arise.

On the other hand obtaining results from physical experiments is difficult and expensive as well, since methods like CT scans and coloring with radioactive elements are necessary. Some of these procedures are quite sensitive to errors.

Using mathematical modeling, the equations need to be found and parametrized and the
model needs to be implemented using programming languages. This can be time consuming and some sophisticated simulations need a lot of computational resources like high performance computing cluster. As soon as the simulations are running, multiple tests can be done without wasting material and the calculations are usually faster than physical experiments. The solutions of the simulations are obtained immediately without post-processing.

1.1 The Enhanced Permeability and Retention Effect

The enhanced permeability and retention (EPR) effect describes how molecules of certain sizes tend to accumulate preferentially in tumor tissue than in normal tissue. Hiroshi Maeda published as one of the first researchers new findings about this effect (Matsumura and Maeda, 1986; Maeda et al., 2003; Maeda, 2012). Various factors reinforce this effect and will be described in this chapter.

The EPR effect is a result of the rapid tumor growth and suppressed lymphatic drainage (Maeda et al., 2003). The tumor requires large amounts of oxygen and nutrients for growth. This results in the need to recruit the body’s vasculature to the tumor. Tumor vasculature is different from healthy vasculature in that the endothelial cells have wide fenestrations (Taurin et al., 2012). The fenestrations result from accelerated growth since the cells cannot grow and align appropriately. Sustenance is faster through the bigger pores between the endothelial cells than in healthy tissue because more particles can diffuse into the tumor tissue at once. On the other hand, the lymphatic system within the tumor is insufficient since it is not built up appropriately. Thus, the uptake of the tumor tissue will not wash out as fast as in healthy tissue (Miao et al., 2015). The retention of micromolecular drugs inside normal tissue is usually a few minutes. In contrast, the retention time in malignant tissue can last days to weeks (Maeda, 2012).

Targeting drugs in general exploit the fact that a drug has a site of intended action and should be brought to ill tissue without having a side effect elsewhere. Therefore, a release of
Figure 1.1: Comparison of the vasculature in normal and in tumor tissue.

the drug in the bloodstream within the tumor would be targeted drug delivery as well. This also applies to other diseases. However, the EPR effect supports the usage of targeting drugs by facilitating the import of macromolecular drugs into the tumor tissue, which is unlikely in normal tissue (Maeda et al., 2003; Maeda, 2012; Wong et al., 2015). Since macromolecules are not able to diffuse passively into normal tissue, the EPR effect is exploited for deliberate treatment of tumor cells. Therefore, the likelihood for healthy cells to be killed by the cytotoxic drug decreases, while it is more likely that the drug treats the tumor instead of passing the tumor.

Disadvantages of the EPR effect are a higher interstitial fluid pressure (IFP) inside the tumor and thus enhanced diffusion of molecules out of the tumor (Wong et al., 2015; Minchinton and Tannock, 2006). Because a lot of nutrients and fluid can move into the tumor and remains inside, the interstitial pressure within the tumor is much higher than the vascular pressure outside of the tumor. The interstitial pressure gradient creates a outward directed velocity field within the tumor that drives the uptake from the core of the tumor back towards its edge (Baxter and Jain, 1989). As a result the EPR effect allows transport both into and out of the tumor.
1.2 Targeted Drug Delivery in solid tumors

Macromolecules with a molecular weight > 45 kDa will result in a selective uptake mainly within the tumor since the fenestrations of normal tissue are too small (Maeda et al., 2003). In normal tissue the endothelial cells are aligned close to each other, so that mainly small molecules diffuse through the fenestrations. There are also transporters in the endothelial cells for active transport that can even work against a gradient (Saxena et al., 1991), but in this thesis we focus on the passive transport by diffusion through the fenestrations. The macromolecules will mainly stay within the vasculature without passing the barrier into the tissue. The movement of the macromolecules through the fenestrations into the tumor tissue is depicted in Fig. 1.1.

The encapsulation of the cytotoxic anticancer drugs within liposomes primarily shields the drug but as a side effect it increases its size. One representative of these anticancer drugs is Doxorubicin (DOX), whose interference with DNA leads to cell death. Their adverse side effects include cardiomyopathy which can lead to heart failure and suppression of white blood cell production in the bone marrow (Hinow et al., 2016). These side effects can be reduced by targeted drug delivery. The liposomal formulation of DOX became increasingly common, for example as Doxil® or ThermoDox®. The structure of these liposomes is depicted in Fig. 1.2. They are spherical vesicles having the drug surrounded by at least one lipid bilayer. This layer is again surrounded by a coat which protects the liposome against destruction by
the immune system. The size as well as the shape of the liposomes affect the biodistribution and uptake, in addition to circulation time of molecules (Arnida et al., 2011). Therefore, targeted drugs are designed as bigger liposomes carrying the actual drug into the tumor cell, where they release their cargo.

The release is triggered by a chemical or physical reaction such as heat (Gasselhuber et al., 2010; Kneidl et al., 2014), ultrasound (Staruch et al., 2011; Rizzitelli et al., 2015) or the pH difference between healthy tissue and the more acidic malignant tissue (Liu et al., 2014). For example, the low temperature-sensitive liposomal formulation of Doxorubicin (LTLD) releases its cargo in a high concentration when it is heated to temperatures between 39.5 °C and 42 °C (Zagar et al., 2014). The tissue can be heated via radio frequency tumor ablation (RFA) (Gasselhuber et al., 2010) or ultrasound mediated hyperthermia (Gasselhuber et al., 2012a).

Our goal is to create an easily expandable model that can handle several targeted drug delivery methods. Therefore, we base on the work of Stapleton et al. (2013) and extend their model with the drug release using an adjustable “transfer function”. They proposed a model for the liposome transport in solid tumors based on biophysical transport equations. These equations describe the pressure driven fluid flow across blood vessels and through the tumor interstitium. Their parametrization results from three preclinical animal tumor models. Within this thesis, the release of the drug in our mathematical model is simulated using temperature-sensitive liposomes based on the results of Gasselhuber et al. (2010). They proposed a spatio-temporal multicompartmental pharmacokinetic model to describe the transport and release of the drug from the low temperature-sensitive liposomal formulation of Doxorubicin into the tumor plasma space of a human. They parametrized their model based on data for human lung cancer cells.
1.3 Structure of this Thesis

In this thesis we study the anticancer drug delivery using temperature-sensitive liposomes. The relevant properties of tumors and anticancer drugs were explained in detail in Ch. 1 to create a solid background knowledge and to aid in the understanding of the mathematical model. In Ch. 2 we derive a mathematical model to improve the understanding of the ongoing processing during targeted drug delivery. In Ch. 3 we touch upon the numerical methods and the results of the numerical simulations will follow in Ch. 4. In Ch. 5 we will summarize the work in a discussion and prospects will be given.
Chapter 2

The Mathematical Model

We consider an isolated model tumor with spherical symmetry that is homogeneously perfused by blood vessels and lacks lymphatic drainage (Baxter and Jain, 1989; Stapleton et al., 2013). So the microscopic elements as blood vessels, interstitial tumor matrix and individual cells are not modeled explicitly. After a suitable normalization by the tumor radius $R$, the spatial variable is in the interval $r \in [0, 1]$.

A schematic depiction of the model of drug release of temperature-sensitive liposomes is given in Fig. 2.1. Liposomes are delivered from the blood plasma compartment. We assume that the liposome injection rapidly equilibrates in the plasma, so an ordinary differential equation is sufficient to describe their concentration within the blood plasma. Their concentration $c(t)$ follows simple first order kinetics

$$\frac{dc}{dt} = -kc, \quad c(0) = c_0. \tag{2.1}$$

Here $k$ denotes the rate of clearance from the plasma (Ishida et al., 2002) and results from the assumption that the liposomes have a half-life of 24 hours. $c_0$ is the initial concentration. As long as the vascular pressure $p_v$ is higher than the interstitial pressure $p_i$ the liposomes are able to move from the blood plasma into the tumor tissue. Since the tumor volume is sufficiently small in comparison to the blood volume the passing of the liposomes into the tumor does not impact $c$ significantly. As long as the temperature surrounding the liposomes is sufficiently high the liposomes can release their cytotoxic cargo. After the liposomes release their cargo they are no longer considered. The concentration of the drug inside the tumor is denoted as $w(r, t)$.

Baxter and Jain (1989) derived an analytic steady-state solution for the IFP for an
isolated tumor without extravascular binding. Based on their aforementioned assumptions the interstitial fluid pressure is given by

$$p_i(r) = p_{i,\text{max}} \left( 1 - \frac{\sinh(\alpha r)}{r \sinh(\alpha)} \right), \quad (2.2)$$

where $p_{i,\text{max}}$ is the maximal IFP. Stapleton et al. (2013) report values between 3.5 - 5.7 mmHg, 1.7 - 4.7 mmHg and 4.56 - 42.3 mmHg from experiments of ME180, H520 and VX2 tumors, respectively. These experiments were performed in mice (ME180, H520) and rabbits (VX2). $\alpha$ is a dimensionless parameter that is defined as the ratio of vascular to interstitial permeability. A smaller value of $\alpha$ means that the interstitial permeability is relatively high, whereas a larger value for $\alpha$ corresponds to a relatively low interstitial permeability.
permeability. Therefore, a smaller value of $\alpha$ means that macromolecules can move more easily into the tumor tissue.

The interstitial fluid velocity is obtained from Darcy’s law,

$$v(r) = -K \frac{\partial p_i}{\partial r} = \frac{K p_{i,\text{max}}}{R} \frac{\alpha r \cosh(\alpha r) - \sinh(\alpha r)}{r^2 \sinh(\alpha)} ,$$

where $K$ denotes the hydraulic conductivity. This physical constant describes the permeability of the interstitium. This velocity field drives the transport of liposomes in the interstitial region of the tumor. Therefore $v(R) > 0$ on the edge of the tumor.

We assume a unidirectional flow of the liposomes such that the rate of transvascular exchange is proportional to the difference between vascular pressure $p_v$ and $p_i$. Since the IFP is increasing towards the center of the tumor whereas $p_v$ is assumed constant throughout the tissue, the exchange is only possible as long as the interstitial pressure is smaller than the vascular pressure. The concentration of the liposomes $u(r, t) \in (0, 1) \times (0, t_e)$, with $t_e$ as the end of the time interval, is given by

$$\frac{\partial u}{\partial t} = \frac{1}{V_r} c f_c (1 - \sigma) (p_v - p_i) \cdot H(p_v - p_i) - \theta_l \frac{\partial}{\partial r} (uv) - h(T(r, t)) u \quad \text{in} \quad (0, 1) \times (0, t_e).$$

(2.4)

The initial condition and Neumann boundary condition, respectively, are given by

$$u(r, 0) = u_0(r) \quad \text{for} \quad r \in \Omega := (0, 1),$$  

(2.5)

$$\frac{\partial u}{\partial r} = 0 \quad \text{on} \quad \Gamma := \partial \Omega \times (0, t_e).$$  

(2.6)

If the IFP is higher than $p_v$, no transvascular exchange can occur. Therefore, the positivity is modeled by a Heaviside step function $H$. The only source of liposomes in the tumor is the plasma perfusing the tumor. Here $T(r, t)$ denotes the tissue temperature (which is also assumed to be spherically symmetric), and $h$ is called a “transfer function”, similar to that in Hinow et al. (2016). $V_r$ is the ratio of the tumor volume to the blood volume. It ensures conservation of mass as the liposomes move between compartments of different volumes. The blood volume is 5 L whereas the tumor volume is 1 cm$^3$ and therefore significantly smaller.
$V_r$ is given by

$$V_r = \frac{3 \cdot 5}{4\pi \cdot 10^3} \approx 10^{-3}. \quad (2.7)$$

The constant $f_c$ denotes the capillary filtration coefficient, which is a constant of proportionality between capillary surface area and capillary hydraulic conductivity, and describes the permeability of the capillaries. The constant $\sigma$ denotes the filtration reflection coefficient, which describes the fact that molecules bigger than a specific threshold are retained by the semipermeable membrane. Although the permeability is enhanced within tumors not every molecule will diffuse into the tumor tissue.

Stapleton et al. (2013) describe the fractional rate of liposome transport through interstitium to fluid flow as $\theta_l = 0.5$. This results from the inertia of the relatively large liposomes within the fluid. As the liposomes are rather large particles of molecular weights in the range of $100\, MDa$ (Stapleton et al., 2013), they are not subject to diffusion. This is different for the drug that is released from them. The concentration of the drug $w(r, t) \in (0, 1) \times (0, t_e)$ within the tumor is governed

$$\frac{\partial w}{\partial t} = \frac{\partial^2}{\partial r^2} (Dw) - \theta_d \frac{\partial}{\partial r} (wv) + \beta h(T(r, t))u. \quad (2.8)$$

The initial condition and Neumann boundary condition, respectively, are given by

$$w(r, 0) = w_0(r) \quad \text{for} \quad r \in \Omega, \quad (2.9)$$

$$\frac{\partial w}{\partial r} = 0 \quad \text{on} \quad \Gamma := \partial \Omega \times (0, t_e). \quad (2.10)$$

The drug is subject to transport and diffusion, and is supplied by the liposomes as they enter the heated region. The diffusion coefficient $D$, as found in Fick’s second law of diffusion, describes the proportionality between the molar flux due to molecular diffusion and the gradient in the concentration. Since the tissue is heated up the diffusivity depends on the status of the tissue and therefore on the temperature: If the tissue is already ablated, the diffusivity decreases significantly within the necrotic tissue (Gasselhuber et al., 2010). For simplicity, we still assume the $D$ to be constant. Suggestions for improvement can be found
in the discussion in Ch. 5.

The fractional rate of the transport of drug molecules through interstitium due to fluid flow is described by $\theta_d = 1$. This is an initial guess and needs to be parametrized more precisely. The constant $\beta > 0$ is the yield of drug molecules supplied by one liposome.

At the moment, Eq. 2.8 models only the release of the drug and its motion through the tumor. In the future, the metabolism and uptake of the drug need to be included. Ch. 5 will give more details on the improvement of the model.

The total amount of liposomes within the tumor is given by the integral

$$\bar{u}(t) = \int_V u(r, t) dV = \int_0^R u(r, t) \cdot 4\pi r^2 dr. \quad (2.11)$$

The parameters and their values used within the simulations are summed up in Table 2.1.
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Unit</th>
<th>Interpretation</th>
<th>Value(s)</th>
<th>Reference</th>
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</thead>
<tbody>
<tr>
<td>$p_i,max$</td>
<td>mmHg</td>
<td>maximal value of IFP</td>
<td>19.8</td>
<td>(Stapleton et al., 2013)</td>
</tr>
<tr>
<td>$p_v$</td>
<td>mmHg</td>
<td>vascular pressure</td>
<td>10</td>
<td>-</td>
</tr>
<tr>
<td>$R$</td>
<td>cm</td>
<td>radius of tumor</td>
<td>1</td>
<td>-</td>
</tr>
<tr>
<td>$\alpha$</td>
<td>-</td>
<td>ratio of vascular to interstitial permeability</td>
<td>1; 3; 5; 10; 25</td>
<td>(Stapleton et al., 2013)</td>
</tr>
<tr>
<td>$k$</td>
<td>$s^{-1}$</td>
<td>rate of liposome clearance from the plasma</td>
<td>$8.3 \times 10^{-6}$</td>
<td>(Ishida et al., 2002)</td>
</tr>
<tr>
<td>$c_0$</td>
<td>$pmol \cdot cm^{-3}$</td>
<td>Initial plasma concentration of liposomes</td>
<td>0.5</td>
<td>(Hinow et al., 2016)</td>
</tr>
<tr>
<td>$K$</td>
<td>$cm^2 \cdot mmHg^{-1} \cdot s^{-1}$</td>
<td>hydraulic conductivity or interstitial permeability</td>
<td>$662 \times 10^{-7}$</td>
<td>(Stapleton et al., 2013)</td>
</tr>
<tr>
<td>$\theta_l$</td>
<td>-</td>
<td>fractional rate of liposome transport through interstitium to fluid flow</td>
<td>0.5</td>
<td>(Stapleton et al., 2013)</td>
</tr>
<tr>
<td>$\sigma$</td>
<td>-</td>
<td>filtration reflection coefficient</td>
<td>0.19</td>
<td>(Stapleton et al., 2013)</td>
</tr>
<tr>
<td>$f_c$</td>
<td>$mmHg^{-1} \cdot s^{-1}$</td>
<td>capillary filtration coefficient</td>
<td>$1088 \times 10^{-7}$</td>
<td>(Stapleton et al., 2013)</td>
</tr>
<tr>
<td>$V_r$</td>
<td>-</td>
<td>relation of tissue to blood volume</td>
<td>$10^{-3}$</td>
<td>-</td>
</tr>
<tr>
<td>$D$</td>
<td>$cm^2 \cdot s^{-1}$</td>
<td>Diffusion coefficient of DOX</td>
<td>$6.7 \times 10^{-7}$ (non ablated)</td>
<td>(Gasselhuber et al., 2010)</td>
</tr>
<tr>
<td>$\theta_d$</td>
<td>-</td>
<td>fractional rate of DOX transport through interstitium to fluid flow</td>
<td>1</td>
<td>-</td>
</tr>
<tr>
<td>$\beta$</td>
<td>-</td>
<td>drug loading coefficient of DOX</td>
<td>$1.3 \times 10^{-7}$</td>
<td>(Hinow et al., 2016)</td>
</tr>
</tbody>
</table>

Table 2.1: Summary of all parameters for numerical approximation
Chapter 3

Numerical Methods

Eq. (2.4) is an example of an Advection-Reaction Equation (see Chapter 3.1) and Eq. (2.8) is an example of an Advection-Reaction-Diffusion Equation (see Chapter 3.2). In this thesis, we use the Strang Splitting method for the numerical approximation of these equations, which is a representative of an Operator Splitting method (Strang, 1968; Hundsdorfer, 2000). In these methods the equations are broken up in their physical components and then these components are solved sequentially. This method can be applied when one component is stiffer than the other or to separate several dimensions. Stiffness describes equations that cannot be solved with an explicit method since at least one component of the equation decays much faster than the other components.

This chapter analyses the numerical methods used for each equation generated by the Operator Splitting Method (see Appendix A).

3.1 The Advection-Reaction Equation

Eq. (2.4) is an example of an Advection-Reaction equation. Generally speaking the reaction term in this equation creates sources or sinks of the quantity, whereas the advection term describes the transport of this quantity by bulk motion.

Applying the Strang splitting described in Appendix A, we can write

\[ \frac{\partial u}{\partial t} = \frac{1}{V_r} c f_c (1 - \sigma) (p_v - p_i) \cdot H(p_v - p_i) - h(T) u - \theta I \frac{\partial}{\partial r} (uv) \]

**reaction term**

**advection term**
Figure 3.1: Scheme for the Strang Splitting of $u$

\[ \frac{\partial u^*}{\partial \tau} = \frac{1}{V_r} c f_c (1 - \sigma) (p_v - p_i) \cdot H(p_v - p_i) - h(T) u^* \quad \text{for} \quad t_n \leq \tau \leq t_n + \frac{1}{2} \]  

(3.1)

\[ \frac{\partial u^{**}}{\partial \tau} = -\theta_l \frac{\partial}{\partial r} (u^{**} v) \quad \text{for} \quad t_n \leq \tau \leq t_n + 1 \]  

(3.2)

\[ \frac{\partial u^{***}}{\partial \tau} = \frac{1}{V_r} c f_c (1 - \sigma) (p_v - p_i) \cdot H(p_v - p_i) - h(T) u^{***} \quad \text{for} \quad t_n + \frac{1}{2} \leq \tau \leq t_n + 1 \]  

(3.3)

with

\[ u^*(t_n) = u_n, \quad u^{**}(t_n) = u^*(t_n + \frac{1}{2}), \quad u^{***}(t_n + \frac{1}{2}) = u^{**}(t_n + 1). \]

This results in $u(t_{n+1}) = u^{***}(t_{n+1})$. Here the equation was split in a physical manner. First the reaction term will be evaluated and in the next step the transport will be calculated. The scheme is depicted in Fig. 3.1.

Eq. 3.1 and 3.3 are solved by the first representative of an L-stable method (definitions on stability can be found in Appendix B), the implicit Euler method

\[ u_{\tau+1} = u_\tau + \Delta \tau g(r, \tau + \Delta \tau, u_{\tau+1}) \]  

(3.4)

\[ g(r, \tau + \Delta \tau, u_{\tau+1}) = \frac{1}{V_r} c(\tau + \Delta \tau) f_c (1 - \sigma) (p_v - p_i) \cdot H(p_v - p_i) - h(T) u_{\tau+1}. \]  

(3.5)

The implicit Euler method was implemented by using a fixpoint iteration with an initial step using the explicit Euler method. With this method no system of equation needs to be solved. The implicit Euler method has order $O(\Delta t)$.

The term $O(\Delta t)$ denotes a quantity whose size is proportional to $\Delta t$ or even smaller.
Eq. 3.2 was solved by using the first order upwind differencing scheme, which takes the flow direction into account. This discretization method creates the following equation which needs to be implemented:

\[
    u_{r+1}^{\tau} = \left(1 - \frac{\theta \Delta t}{\Delta r} v_r \right) u_r^\tau + \frac{\theta \Delta t}{\Delta r} v_{r-1} u_{r-1}^\tau \quad \text{if } \theta_l \geq 0. \tag{3.6}
\]

The advantages of this upwind differencing method are the properties of positivity and local preservation of mass and uniform stability. The method does not prevent a loss of mass due to outflow out of the domain. The disadvantage is artificial diffusion within the advection term, which could be eliminated by implementing a more complex method (Quarteroni et al., 2010, Remark 12.6). The artificial diffusion develops, because the mass within one grid cell moves only partly into the next grid cell. A method which attacks this issue will be discussed in Ch. 5.

The upwind differencing scheme is stable as long as the Courant-Friedrichs-Lewy condition is fulfilled. This condition describes the convergence behavior of \( x_i \) in its neighborhood. In this case the condition says that \( x_i - \theta \Delta t \Delta r \leq x_{i+1} \), otherwise the method is not convergent. Since \( x_i = i \Delta r \), the condition for the upwind method is

\[
    0 \leq \frac{\theta \Delta t}{\Delta r} \leq 1 \quad \text{for } \theta_l \geq 0 \tag{3.7}
\]

### 3.2 The Advection-Reaction-Diffusion Equation

Eq. (2.8) is an example of an Advection-Reaction-Diffusion equation. In addition to the structure described in the previous Chapter 3.1 a diffusion term occurs in this type of equation. Applying the Strang splitting described in Appendix A, we can write

\[
    \frac{\partial w}{\partial t} = \beta h(T)u - \theta_d \frac{\partial}{\partial r} (wv) + \frac{\partial^2}{\partial r^2}(Dw) \quad \text{(reaction term)}
\]

\[
    \text{---} \quad \text{(advection term)}
\]

\[
    \text{---} \quad \text{(diffusion term)}
\]

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\[ \frac{\partial w^{(1)}}{\partial \tau} = \beta h(T) u_t \quad \text{for} \quad t_n \leq \tau \leq t_{n+\frac{1}{2}} \]  
\[ \frac{\partial w^{(2)}}{\partial \tau} = -\theta_d \frac{\partial}{\partial r} (w^{(2)} v) \quad \text{for} \quad t_n \leq \tau \leq t_{n+\frac{1}{2}} \]  
\[ \frac{\partial w^{(3)}}{\partial \tau} = \frac{\partial^2}{\partial r^2} (Dw^{(3)}) \quad \text{for} \quad t_n \leq \tau \leq t_{n+1} \]  
\[ \frac{\partial w^{(4)}}{\partial \tau} = -\theta_d \frac{\partial}{\partial r} (w^{(4)} v) \quad \text{for} \quad t_{n+\frac{1}{2}} \leq \tau \leq t_{n+1} \]  
\[ \frac{\partial w^{(5)}}{\partial \tau} = \beta h(T) u \quad \text{for} \quad t_{n+\frac{1}{2}} \leq \tau \leq t_{n+1} \]

with

\[ w^{(1)}(t_n) = w_n, \quad w^{(2)}(t_n) = w^{(1)}(t_{n+\frac{1}{2}}), \quad w^{(3)}(t_n) = w^{(2)}(t_{n+\frac{1}{2}}), \]
\[ w^{(4)}(t_{n+\frac{1}{2}}) = w^{(3)}(t_{n+1}), \quad w^{(5)}(t_{n+\frac{1}{2}}) = u^{(4)}(t_{n+1}). \]

The solution of the reaction equation for the drug can be found analytically. Since the right hand side does not depend on \( w \), the equation is linear and can therefore just be multiplied by \( \Delta \tau \) to evaluate the reaction. The reaction equation can also be calculated by the implicit Euler method as described in Ch. 3.1. This becomes important if the reaction equation would be extended by other terms like the uptake of the drug or metabolism. Then the reaction term would also depend on \( w \) itself and may not be solved by simple integration. 

For the advection term the upwind method is used as described in Ch. 3.1. The diffusion term is approximated by the Crank-Nicolson method (Grossmann et al., 2007, Section 2.6.1). This method is a combination of a half implicit and a half explicit Euler step and can be derived as follows:

\[ \frac{\partial w}{\partial t} = \frac{1}{2} \left( \frac{\partial^2 Dw}{\partial t^2} \bigg|_t + \frac{\partial^2 Dw}{\partial t^2} \bigg|_{t+1} \right) = \frac{1}{2} \left( \frac{Dw^{t+1}_{r+1} - 2Dw^{t}_{r} + Dw^{t}_{r-1}}{\Delta r^2} + \frac{Dw^{t}_{r+1} - 2Dw^{t}_{r} + Dw^{t}_{r-1}}{\Delta r^2} \right). \]  

Reordering with all \( w^{t+1} \) on the left hand side we get the final form of the Crank-Nicolson
\[- \kappa D w_{r+1}^{t+1} + (1 + 2\kappa D) w_{r}^{t+1} - \kappa D w_{r-1}^{t+1} = \kappa D w_{r+1}^{t} + (1 - 2\kappa D) w_{r}^{t} + \kappa D w_{r-1}^{t}, \]  
(3.14)

with

\[ \kappa = \frac{\Delta t}{2\Delta r^2}. \]  
(3.15)

This implicit method results in a tridiagonal matrix, which needs to be calculated in each time step. The advantage of this method is that the system of equations can be solved within \( O(r) \) work. This makes the Crank-Nicolson method basically as efficient as an explicit method.

With a series expansion in space and in time it can be shown that the Crank-Nicolson method is second order accurate in space and in time (Hundsdorfer, 2000). For stability the Courant-Friedrichs-Lewy condition is fulfilled if

\[ \frac{\Delta t}{\Delta r^2} \leq \frac{1}{2}. \]  
(3.16)

If this condition is violated, oscillations may occur.
Chapter 4

Results

The numerical methods were implemented using the Open Source computing language Python (Python Software Foundation, 2017). The source code is attached in Appendix C.

4.1 Transfer function

The transfer function was determined from Gasselhuber et al. (2010, Figure 2). They experimentally determined the release of Doxorubicin by heating human blood plasma in single cuvettes to 25°C and 37 – 47°C for at least 5 minutes. A liposome suspension (10 µL, 2 mg/mL DOX) of ThermoDox® (Celsion, Columbia, MD) was added to 750 µL of the heated plasma. The fluorescence intensity caused by the DOX release was monitored by a spectrofluorometer. A more precise description of the methods and materials can be found in Gasselhuber et al. (2010). We focused on the results depicted in Gasselhuber et al. (2010, Figure 2).

We are especially interested in the approximately linear slope within the first second (see Fig. 4.1). Therefore the release in Gasselhuber et al. (2010, Figure 2) is evaluated after one second with the slope taken from the gradient triangle. We consider only those temperatures which force the liposomes to release at least 50 % of their cargo, so the gradients were fitted to a piecewise linear function (see Fig. 4.1) dependent on the temperature.

\[
h(T) = \begin{cases} 
0 & T \leq 38 \\
0.3789(T - 38) & 38 < T < 39 \\
0.0176T - 0.3107 & 39 \leq T. 
\end{cases}
\] (4.1)
4.2 Pressure and Velocity

We assume a constant vascular pressure so that the transvascular exchange is driven by the IFP. This relation is depicted in Fig. 4.2. The liposomes cannot pass the point where the IFP outweighs the vascular pressure. However, the drug can diffuse further into the tumor.
4.3 Temperature distribution

We assume a temperature field that has the value 50 °C in the center of the tumor and decreases to 36 °C body temperature at the edge of the tumor. We approximate the decay of the temperature similar to the interstitial pressure so that we use a hyperbolic sine, whereas the temporal aspect is taken from Gasslhuber et al. (2010, Figure 9). The exponential decay of the temperature was taken into account

\[
T(r,t) = T_r \cdot T_t + T_0
= \left( 20 - \frac{20 \sinh(3r)}{r \sinh(3)} \right) e^{-\lambda t} + 36,
\]

where \( r \) is the normalized spacial component and \( t \) is the time. The exponential decay constant \( \lambda = 8.32 \ h^{-1} \) was estimated from the half-life of ca. 5 minutes (Gasslhuber et al., 2010, Figure 9). The function is depicted in Fig. 4.4.
Figure 4.5: Concentration of liposomes $u$ in a steady-state temperature field in absence of transport. The transition zone of the drug release is marked by dashed lines.

Figure 4.6: Concentration of liposomes $u$ in a time dependent temperature field in absence of transport.

4.4 Liposome concentration

4.4.1 Reaction term

At first, we a look at the reaction equation on its own (Fig. 4.5) in the steady state of the temperature field, such that $T(r, t) = T(r, 0)$. Although the assumption of the steady state
is not realistic, the results from that calculation help to understand the impact of both terms within the reaction term.

The temperature increases towards the center of the tumor and the transition zone for the drug release (38 – 39 °C) is marked by a dashed line in Fig. 4.5. In the beginning, the liposomes enter the tumor at a rate proportional to the pressure difference. As soon as they reach the transition zone the liposomes start to release their cargo such that the concentration of liposomes decreases towards the center of the tumor. This explains the steep slope at approximately \( r = 0.9 \). With \( \alpha = 3 \) this structure of the individual terms is still visible, but the more \( \alpha \) increases, the more the slope becomes uniformly and steep.

Since the temperature of the tissue cools down quickly with time, the liposomes decay in reality at a lower rate (Fig. 4.6). So in the realistic simulation the reaction term will be dominated by the exponential decay of the temperature and of the liposome concentration in the plasma.

### 4.4.2 Advection term

To examine the advection term, an initial block of higher concentration of liposomes was set on the domain in the interval \([0.225, 0.275]\) and \([0.725, 0.775]\). These locations are chosen for analytic purposes and are not quite appropriate. The pure advection becomes visible without the differences in concentration due to the reaction terms (Fig. 4.7). This part of the analysis is important to show the impact of the interstitial velocity \( v \) (Fig. 4.3) on Eq. 2.4. Additionally the significant artificial diffusion resulting from the upwind differencing scheme becomes visible.

Since the velocity is increasing towards the edge of the tumor, the peaks of concentration are not symmetric but faster moving at the right end than at the left end. This increases the artificial diffusion of the upwind differencing method even more. The material tends to stream out of the tumor, although the likelihood depends on the distance, which it can pass during that amount of time. Therefore the likelihood of liposomes streaming out of the
tumor depends on $\alpha$.

Notable is the peak of advection in the plot $\alpha = 3$ (Fig. 4.7b). If $\alpha$ has a smaller value, then the velocity is too weak to move the material far away. With higher values of $\alpha$ the magnitude of velocity decreases to some extent in comparison to those values with lower $\alpha$, but towards the edge of the tumor it increases more rapidly.

### 4.4.3 Full equation

The impact of $\alpha$ is even more visible when we have a look at how far the liposomes penetrate the tumor. Since the IFP depends on $\alpha$, so does the reaction term. The liposomes cannot
pass the point where the IFP outweighs the vascular pressure (Fig 4.2 and Fig. 4.8).

The total amount of liposomes within the tumor can be calculated by integrating $u$ as described in Eq. 2.11 and is depicted in Fig. 4.9. This relatively high concentration in comparison to the initial concentration $c_0$ within the plasma results from the change of volume. The blood volume is ca. 5 $L$ whereas the tumor volume is only 1 $cm^3$. The concentration is plotted on a logarithmic scale such that the decay is in fact exponential, resulting from Eq. 2.1.

## 4.5 Drug concentration

### 4.5.1 Diffusion term

Drug molecules are subject to diffusion. The diffusion acts autonomously since it does not depend on the liposome concentration $u$ or the ratio of vascular to interstitial permeability $\alpha$. Due to the concentration within $w$ the drug molecules diffuse toward areas with less concentration as depicted in Fig. 4.10. This motion is independent of the flow directions such that the drug can move further into the tumor due to diffusion and especially pass the point where the IFP outweighs the vascular pressure.
4.5.2 Full equation

For the temperature distribution we assume that the tumor is heated by a probe located in the center of the tumor. If the liposomes cannot move far enough into the tumor such that their temperature reaches a specific threshold, no drug can be released. This is the reason why in the case of $\alpha \geq 25$ no drug concentration can be found in the tumor. Looking at the distance the drug moves into the tumor, the impact of the diffusion becomes clearly visible. The reaction term just generates the drug in place, whereas the advection term moves the drug outward. However, integrating over the time (Fig 4.11), the drug moves further inward than the liposomes (Fig. 4.8) due to the diffusion.
The distribution of the total amount of drug within the tumor can be calculated by integrating $w$ as described in Eq. 2.11 and is depicted in Fig. 4.12. On the one hand, the amount of drug in the tumor is impacted by the amount of liposomes in the tumor. If less liposomes are present then less drug molecules can be released. On the other hand, the release of the drug is impacted by the temperature since the release rate $h$ depends on the temperature. Thus, the release depends on the time. When the tumor cools down, less drug can be released from the liposomes. Since $\alpha$ controls the depth of the import of liposomes and the velocity and consequently the advection it has also an impact on the retention time of the drug within the tumor. An increasing $\alpha$ leads to a shorter retention time.

Notable is the decay of $w$ with increasing $\alpha$ as described in the beginning of this section. The influence of $\alpha$ on the concentration of the drug becomes obvious, when $w$ is plotted three-dimensional with respect to time and space (Fig. 4.13). Although the difference of $\alpha$ is not big, the difference of the drug concentration is clearly affected.
Figure 4.13: Concentration of the drug with respect to space and time

(a) $\alpha = 1$

(b) $\alpha = 3$
Chapter 5

Conclusion

5.1 Summary

The simulation of tumor growth and tumor treatment is one way for a better understanding of the particular components in these processes. Precise measurements of drug concentration in tumors requires a lot of pre- and post-processing steps, whereas mathematical simulation gives immediate results. On the other hand, mathematical simulations are only as reliable as the model. The more physical processes and side effects are neglected the less precise the simulation results. After setting up a mathematical model, the impact of single terms on the system can be analyzed.

We have set up a mathematical model that represents the delivery of temperature-sensitive liposomes and the release of their cargo within tumor tissue. These liposomes are spherical vesicles containing the drug and release their cargo in regions of increased temperature. We have modeled the elevated interstitial pressure and the interstitial velocity based on the model of Baxter and Jain (1989). For the concentration of the liposomes we extended the model given in Stapleton et al. (2013) and coupled that concentration with the concentration of the drug itself. The model for the drug release follows the idea of Hinow et al. (2016), where the drug is released by a transfer function. Therefore, we implemented some results for thermo-sensitive drug delivery referring to Gasselhuber et al. (2010). Our current model can be improved or extended as described in Ch. 5.2.

The parametrization of the mathematical model relies on parameters from available literature. A specific threshold exists for temperature-sensitive liposomes causing them to release their cargo as shown in Fig. 4.1 (Gasselhuber et al., 2010, Figure 2). This threshold is usu-
ally achieved by heating such as radio frequency tumor ablation. The temperature-sensitive liposomes are used to complement the tumor treatment by RFA. Around the area of ablated tissue the tumor will be treated additionally by the drug released from the liposomes (Gasselhuber et al., 2010).

The release of the drug depends on the temperature which itself has a temporal and a spatial component. We varied the ratio of vascular to interstitial permeability $\alpha$ in the simulation. As $\alpha$ increases the retention time of the drug within the tumor decreases due to a faster advection. An improvement of the efficiency of the treatment could be an incremental heating of the tumor, such that the liposomes have a better chance to release their cargo.

Developing temperature-sensitive liposomes for targeted anticancer drug delivery is difficult to realize. The liposomes must be able to move far into the tumor although the interstitial pressure counteracts the transport of the liposomes. The release of the drug must be triggered easily enough so that the tumor can be treated sufficiently. However, liposomes outside the tumor should tightly seal the drug to prevent toxic side effects. The diffusion of the drug molecules improves the penetration depth of the tumor.

5.2 Discussion

The mathematical model as well as the numerical approximation can be improved in the future. The most important step in the future is the improvement of the model of the drug. On the one hand, the drug uptake from the cells is not considered yet. This may impact the amount of concentration as well as the behavior of the drug molecules within the tumor in view of the advection and diffusion. The actual intention of the drug is the treatment of tumor cells. The decomposition and metabolism of the drug is not considered yet. An example for the metabolism can be found in Hinow et al. (2016). Those drug molecules which are not taken up from the tumor cells will start to dissociate after some time. After that they cannot treat the tumor cells any longer and should be neglected in the further
calculations.

Currently this model simulates the drug delivery within an isolated tumor. In the future, we can extend this model for tumors surrounded by healthy tissue. This will impact the interstitial pressure and the interstitial velocity at the edge of the tumor as well as the temperature field since the transition between the tissues has to be continuous.

The Operator Splitting method has an order of $O(\Delta t^2)$. However, this order can only be maintained if the numerical methods used in combination with the splitting are of at least the same order. Therefore the implicit Euler method and the upwind differencing method (both explained in Ch. 3.1) need to be replaced by more accurate methods. For sufficiently precise results the time step size and the grid resolution have to be very small which lead to long simulation times. By more precise numerical methods the number of calculations can be reduced since bigger resolutions will result in likewise precise simulation results.

The L-stable implicit Euler method is the first representative of the backward differentiation formulas and is sometimes abbreviated as BDF1. It is the first representative since it only uses one previous step. A suggestion for an improvement is using the second backward differentiation formula (BDF2), which has an order of $O(\Delta t^2)$ and uses the two previous steps for the calculation of the next step. So instead of Eq. 3.4 we receive the following differentiation:

$$u_{\tau+1} = \frac{4u_\tau - u_{\tau-1} + 2\Delta \tau g(r, \tau + \Delta \tau, u_{\tau+1})}{3},$$

(5.1)

where $g(r, \tau + \Delta \tau, u_{\tau+1})$ is still defined by Eq. 3.5.

Many numerical methods exist to approximate the advection equation. Therefore, the upwind differencing method could be replaced by a more accurate method. One of these methods was introduced by Egan and Mahoney (1972). This method reduces the effects of the artificial diffusion by advecting the subgrid center of mass and the subgrid spread of the molecules along with the molecules themselves. This counteracts the error of the advection that the molecules just move partly into the next grid cell. The previously mentioned
upwind differencing method stores only the amount of concentration within one grid cell. The amount within one cell decreases while it advects into the next cell. The decrease of the concentration peak in this simulation due to numerical diffusion is showed in Fig. 4.7. Especially in the cases of a ratio of vascular to interstitial permeability of $\alpha = 3$ or $\alpha = 5$ the diffusion is significant. However, the peak concentration should not decrease and the width of the mass should not spread (Fig. 5.1). Because of the data storage of the method by Egan and Mahoney (1972) the previous shape of the concentration curve can be restored from the additional information. In a one-dimensional case with constant velocity this method is exact. The problem with this method is the complex data storage, since not only the concentration, but also the center of mass and the distribution of the molecules need to be stored. The reconstruction is complex and so the current source code of the simulation needs modification.

Instead of approximating the temperature by Eq. 4.2, thermodynamics, e. g. as found in Gasselhuber et al. (2010), could be implemented into the model. Consequently a new partial differential equation (PDE) has to be solved and the results are used for the calculation of Eq. 2.4 and Eq. 2.8. This will improve the accuracy of the reaction terms in both equations and therefore the whole mathematical model.

Also some constants may be improved by a sufficient modeling of their component. For
example, the diffusion coefficient of the drug is dependent on the status of the tissue. Gasselhuber et al. (2010) refer to two different diffusion constants. Within the ablated tissue, the drug diffusion constant is smaller than in the non-ablated tissue. Therefore, the diffusion coefficient $D$ can be extended as $D(T(r, t))$. The simulation itself can already calculate the diffusion with a continuous function as diffusion coefficient. In the future the function needs to be set up and parametrized. The value of the diffusion constant has to be a continuous function to avoid discontinuity in the solutions. Additionally, the fractional rate of liposome transport through the interstitium to fluid flow $\theta_l$, as well as $\theta_d$ for drug transport, require further investigations. $\theta_l$ was taken from Stapleton et al. (2013) although it is not obvious how they received this value. A value for $\theta_d$ (currently $\theta_d = 1$) needs to be estimated in the same way. Overall, our parametrization was taken from the literature. In a next step the parametrization could be verified or improved by physical experiments.

Although the assumption of radial symmetry is valid, the results would be more realistic if the mathematical model would be extended to all three dimensions. Therefore, the model itself needs some adjustments as well as the implementation for the simulation.


Appendix A

The Operator Splitting Method

The idea of operator splitting can be explained most easily by considering a linear ordinary differential equation (ODE) system like

\[ y'(t) = Ay(t), \quad (A.1) \]

with the analytical solution

\[ y(t_{n+1}) = e^{A(t_{n+1} - t_0)} y(t_0) = e^{\Delta t A} y(t_n). \quad (A.2) \]

Here \( A \) denotes a matrix containing the coefficients for the system of equations and \( \Delta t \) denotes the time step size.

If \( A \) can be split into \( A = \sum_{k=1}^{n} A_k \), then we can solve a system of sub-problems

\[ y'(t) = A_1 y(t) \]

\[ \vdots \]

\[ y'(t) = A_n y(t), \quad (A.3) \]

such that the analytical solution (Eq. (A.2)) can be approximated by

\[ y(t_{n+1}) = e^{\Delta t A_n} \ldots e^{\Delta t A_1} y(t_n). \quad (A.4) \]

This method is the simplest splitting method with an convergence order of \( \mathcal{O}(\Delta t) \). The term \( \mathcal{O}(\Delta t) \) denotes a quantity whose size is proportional to \( \Delta t \) or even smaller. Definitions are summarized in Appendix A on page 38.

To achieve more accuracy and symmetry, the order of the \( A_k \) can be interchanged (Strang, 1968). Therefore the sequences of the sub-problems will be reversed, such that

\[ y(t_{n+1}) = \left( e^{\frac{\Delta t}{2} A_n} \ldots e^{\frac{\Delta t}{2} A_1} \right) \left( e^{\frac{\Delta t}{2} A_1} \ldots e^{\frac{\Delta t}{2} A_n} \right) y(t_n) \]

\[ = e^{\frac{\Delta t}{2} A_n} \ldots e^{\frac{\Delta t}{2} A_2} e^{\Delta t A_1} e^{\frac{\Delta t}{2} A_2} \ldots e^{\frac{\Delta t}{2} A_n} y(t_n). \quad (A.5) \]

By applying a series expansion around \( y(t + \Delta t) \) or using the Baker-Campbell-Hausdorff formula (Hundsdorfer, 2000) and the Zassenhaus formula (Casas et al., 2012), the local and global truncation error can be obtained. Overall, the operator splitting using the Strang method has an accuracy of \( \mathcal{O}(\Delta t^2) \).
Appendix B

Definitions

To apply a numerical method within a simulation, the results gained by using that method should approximate the real solution closely. There exist some measurements for the quality of a numerical method, namely stability, consistence and convergence. Among others, the following definitions classify the methods used in Chapter 3.

Convergence of Numerical Methods

A fundamental theorem for the numerical solution of ODEs was developed by Peter Lax and Robert D. Richtmeyer and can be easily extended to PDEs. In the following the theorem itself and the necessary definitions will be given.

**Theorem.** Lax-Richtmeyer Theorem (Reißel, 2015, page 22)
A numerical method is convergent iff it is 0-stable and consistent. Furthermore, its order of convergence equals its order of consistence.

A proof of this theorem can be found for example in Quarteroni et al. (2010, page 41). Therefore, a consistent method is convergent if it is 0-stable and a 0-stable method is convergent if it is consistent. The zero-stability describes numerical methods that are not sensitive to noisy input data.

**Definition.** Zero Stability (Reißel, 2015, page 21)
Given a general numerical method \( \phi(\Delta t, t, y) \) approximating \( y'(t) = f(t, y) \) we can calculate the solution and a noisy solution for the next time step by

\[
\begin{align*}
y_{i+1} &= \phi(\Delta t, t, y_i), & y_0 &= y_0, \\
\tilde{y}_{i+1} &= \phi(\Delta t, t, \tilde{y}_i) + \Delta t \delta_{i+1}, & y_0 &= y_0 + \delta_0.
\end{align*}
\]

For

\[ |\delta_i| \leq \varepsilon \quad \forall i \] (B.3)

a method is called zero-stable if

\[
\exists \Delta t_0 > 0 \quad \exists \varepsilon > 0 : |y_i - \tilde{y}_i| \leq c_\varepsilon \varepsilon \quad \forall i, \quad \forall \Delta t \leq \Delta t_0.
\] (B.4)

Consistence describes the precision of the solution of a numerical method. In other words, how accurate is the numerical approximation in comparison to the true solution. The order of consistence defines the order of convergence.
**Definition.** Consistence (Reißel, 2015, page 21)

Given the exact evolution \( \Phi(\Delta t, t, y) = y(t) + \int_{t}^{t+\Delta t} f(\tau, y(\tau))d\tau \) for \( y'(t) = f(t, y) \), a numerical approximation \( \phi(\Delta t, t, y) \) is called consistent if

\[
|\Phi(\Delta t, t, y) - \phi(\Delta t, t, y)| \xrightarrow{\Delta t \to 0} 0.
\]

Furthermore, the order of consistence \( p \) is given by

\[
|\Phi(\Delta t, t, y) - \phi(\Delta t, t, y)| = O(\Delta t^{p+1})
\]

**Stability of Numerical Methods**

Beside the previous mentioned zero stability there exist more precise definitions of stability.

**Definition.** Absolute Stability (Quarteroni et al., 2010, page 489)

We consider the linear Cauchy problem (or so called test problem)

\[
\begin{aligned}
  y'(t) &= \lambda y(t), \quad t > 0, \\
  y(0) &= 1
\end{aligned}
\]

with \( \lambda \in \mathbb{C} \), whose solution is \( y(t) = e^{\lambda t} \). As long as \( \Re(\lambda) < 0 \), \( \lim_{t \to \infty} |y(t)| = 0 \). A numerical method for approximating the test problem (B.7) is absolutely stable if

\[
|y_n| \xrightarrow{t_n \to \infty} 0.
\]

Let \( \Delta t \) be the discretization stepsize. Therefore, the numerical solution of \( y_n \) of (B.7) depends on \( \Delta t \) and \( \lambda \) and is not stable for all values of these parameters. The region of absolute stability of the numerical method is defines as a subset of the complex plane

\[
S = \{ z = \Delta t \lambda \in \mathbb{C} : (B.8) \text{ holds} \}.
\]

So \( S \) contains all values of the product \( \Delta t \lambda \) for which the solutions of the numerical method decay to zero as \( t_n \) tends to infinity.

**Remark.** A-Stability and L-Stability (Reißel, 2015, pages 64-68) A numerical method is called A-stable if

\[
\mathbb{C}^- = \{ z \in \mathbb{C} : \Re(z) \leq 0 \} \subset S,
\]

so if the stability region \( S \) (B.9) coincides with the left complex half plane. Furthermore, a numerical method is called L-stable if it is A-stable and the solutions decay sufficiently fast such that

\[
y_n \xrightarrow{\Re(\lambda) \to -\infty} 0.
\]
Another definition of the L-Stability is the boundedness of the parameter $\lambda$

\[ \Re(\lambda) < \infty, \quad \Delta t \to \infty. \] (B.12)

So the L-stable methods are suitable for stiff (or rapidly decreasing) problems whereas A-stable problems cannot represent the steep slope within the solution.
Appendix C

Source Code

```python
from numpy import sqrt, sinh, cosh, exp, pi
import logging, getopt, sys
from parameters import *

# Baxter1989, (8a)
# dimensionless interstitial pressure in isolated tumor
#
# Parameters:
# r : current radial position (cm)
# R : radius of tumor (cm)
# a : dimensionless ratio of vascular to interstitial permeability to fluid flow
def p_hat(r, a):
    if r==0:
        return 1 - a / sinh(a)
    r_hat = r/R
    return 1 - (sinh(a * r_hat) / (sinh(a) * r_hat))

# Baxter1989, (8a)
# interstitial pressure in isolated tumor (mmHg)
#
# Parameters:
# phat : dimensionless interstitial pressure in isolated tumor
# p_e : effective pressure (mmHg)
# p_inf : surrounding pressure (mmHg)
def p_i(phat):
    # Baxter1989, (8a)
    # return phat*(p_e - p_inf) + p_inf
    ### own choice
    return p_imax*phat
```

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# Baxter1989, (8b)
# dimensionless interstitial velocity in isolated tumor
#
# Parameters:
# r : current radial position (cm)
# R : radius of tumor (cm)
# a : dimensionless ratio of vascular to interstitial permeability to fluid flow

def v_hat(r, a):
    if r==0:
        return 0.
    r_hat = r/R
    return (a * r_hat * cosh(a * r_hat) - sinh(a * r_hat)) / ((
        r_hat ** 2) * sinh(a))

# Baxter1989, (8b)
# interstitial velocity in isolated tumor (cm/sec)
#
# Parameters:
# vhat : dimensionless interstitial velocity in isolated tumor
# R : radius of tumor (cm)
# K : hydraulic conductivity (cm**2/(mmHg * sec))
# p_e : effective pressure (mmHg)
# p_inf : surrounding pressure (mmHg)

def v_i(vhat):
    return p_imax * K_t * vhat / R

# Stapleton2013, alpha (as in (2))
# should be between 0.5 and 150
# dimensionless ratio of vascular to interstitial permeability to fluid flow
#
# Parameters:
# V : Volume
# fc : capillary filtration coeff (1/(mmHg * sec))
# K : hydraulic conductivity, interstitial permeability (cm**2/(mmHg * sec))

def alpha(V):
    return V**(1./3) * sqrt(fc_t/(K_t*V))
# first half of Stapleton2013, (1)
#
# Parameters:
# r : current radial position (cm)
# V : Volume
# c : plasma concentration of the nanoparticle
# fc : capillary filtration coeff
# p_v : vasculature pressure (mmHg)
# p_hat : dimensionless interstitial pressure in isolated tumor
# p_r : interstitial pressure in isolated tumor (mmHg)
# sigma: filtration reflection coefficient

def lambda(r, a, t):
    phat = p_hat(r, a)
    pr_i = p_i(phat)
    if (p_v - pr_i) <= 0.:
        return 0.
    return fc_t * (p_v - pr_i) * (1 - sigma) * c(t)

# Concentration clearance
#
# Parameters:
# k : liposome clearance rate
# Cp: initial liposome concentration
# t : time

def c(t):
    return Cp*exp(-k*t)

# Ablation coefficient (Gasselhuber2010)
#
# Parameters:
# currently constant

def D(r, t):
    return 3600*6.7e-7 #cm^2/s

# Temperature distribution
#
# Parameters:
# r: current normalized radial position (cm)

def T(r, t):
    if r==0:
        r=0.00000001
    return (20 - (20. * sinh(3 * r))/(r * sinh(3))) * exp(-k2*t)+36
# heat transfer function
#
# Parameters:
# T : temperature

def h(T):
    if T <= 38:
        return 0
    elif T < 39:
        return (37.6 * T - 1428.8)/100
    else:
        return (1.76842105 * T - 31.07368421)/100

# use Simpson rule

def integrate(u, t, r, Nr):
    h = 1.*R/Nr/3
    I = 0.0
    for i in range(1,int(Nr/2)):
        I += u[2*i-2,t]*r[2*i-2]**2 + 4*u[2*i-1,t]*r[2*i-1]**2 +
             u[2*i,t]*r[2*i]**2
    return I*h*4*pi

# use Operator splitting as explained in the thesis to calculate
# u and w

def getU(C0, Nr, tmin, tmax, al):
    dr = 1.*R/(Nr)
    dt = dr
    Ntau = 100
    dtau = dt/Ntau

    logging.info("dt = "+str(dt))
    logging.info("Nt = "+str(Nt))
    logging.info("dr = "+str(dr))
    logging.info("Nr = "+str(Nr))
    logging.info("alpha = "+str(al))

    # divide r into N equidistant points
    r = numpy.linspace(0, R, Nr)
# create matrix
u = numpy.zeros((Nr, Nt), dtype=numpy.float64)
# initial matrix
u[ :, 0] = C0

# Upwind differencing
# for advection term: u_t + f*u_x = 0
n = dtau/(dr)
m = f*n
logging.info("m = "+str(m))

# speedup: calculate h and v
local_h = numpy.zeros(Nr)
local_v = numpy.zeros(Nr)

# Concentration Liposomes

for t in range(Nt-1):
    # in each time step solve System with Strang Splitting
    # Strang splitting: 0.5A + B + 0.5A
    un = numpy.zeros((Nr, Ntau), dtype=numpy.float64)
    un[ :, 0] = u[ :, t]
    logging.info("t="+str(t))
    logging.debug("Calculate u(r,t)"

    for x in range(Nr):
        local_h[x] = h(T(r[x], (t+1)*dt))
        local_v[x] = v_i(v_hat(r[x], al))

# 0.5 Euler

for tau in range(int(Ntau/2)-1):
    for x in range(Nr):

        it=100
        error=1

        # fix point iteration for backward euler
        # initial euler step
        un[x, tau+1] = un[x, tau] + dtau*(lambda(r[x], al, ~tau)/Vr - h(T(r[x], t*dt))*(un[x, tau]))
        while error > 1e-8:
\[ u_n = u[x, \tau] + d\tau \times \left( \frac{r[x]}{V_r - \text{local}_h[x]} \times (u[x, \tau + 1]) \right) \]

error_v = un[x, \tau + 1] - u_n

error = \text{numpy.sqrt}(\text{numpy.dot}(error_v, error_v))

\[ u_n[\tau + 1] = u_n \]

it = 1

if it < 0:
    print("Too many iterations")
    break

logging.debug("RK1_u")

# set last value as start value
un[:, 0] = un[:, \text{int}(Ntau/2) - 2]

for \tau in range(\text{int}(Ntau/2) - 1):
    for x in range(1, Nr):
        \# chain rule
        un[x, \tau + 1] = (1 - m * \text{local}_v[x]) \times un[x, \tau] + m \times \text{local}_v[x-1] \times un[x-1, \tau]

logging.debug("DC2_u")

for \tau in range(\text{int}(Ntau/2) - 1, Ntau - 1):
    for x in range(1, Nr):
        \# chain rule
        un[x, \tau + 1] = (1 - m * \text{local}_v[x]) \times un[x, \tau] + m \times \text{local}_v[x-1] \times un[x-1, \tau]

logging.debug("DC2.2_u")

# set last value as start value
un[:, \text{int}(Ntau/2) - 1] = un[:, Ntau - 1]
for tau in range(int(Ntau/2)-1, Ntau-1):
    for x in range(Nr):
        it=100
        error=1

        # fix point iteration for backward euler
        # initial euler step
        un[x,tau+1] = un[x,tau] + dtau*(lambda(r[x], al, tau)/Vr - h(T(r[x], t*dt))*(un[x,tau]))
        while error > 1e-8:
            u_n = un[x,tau] + dtau*(lambda(r[x], al, t+tau)/Vr - local_h[x]*(un[x,tau+1]))
            error_v = un[x, tau+1] - u_n
            error = numpy.sqrt(numpy.dot(error_v, error_v))
            un[x,tau+1] = u_n
            it-=1
            if it < 0:
                print("Too many iterations")
                break

        # set next value of u
        u[:,t+1] = un[:,1]

        logging.debug("RK1.2_u")

    # fix boundary
    u[0,:] = u[1,:]

return u

def getW(u, w0, Nr, tmin, tmax, al):
    dr = 1.*R/(Nr)
    dt = dr
    Ntau = 100
    dtau = dt/Ntau

    logging.info("dt = "+str(dt))
    logging.info("Nt = "+str(Nt))
    logging.info("dr = "+str(dr))
logging.info("Nr = " + str(Nr))
logging.info("alpha = " + str(al))

# divide r into N equidistant points
r = numpy.linspace(0, R, Nr)
# create matrix
w = numpy.zeros((Nr, Nt), dtype=numpy.float64)
# initialize matrix
w[:, 0] = w0
# Upwind differencing
# for advection term: \( w_t + w_x = 0 \)
n = dtau/(dr)
m = f*n
logging.info("m = " + str(m))

# speedup: calculate h, v and D
local_h = numpy.zeros(Nr)
local_v = numpy.zeros(Nr)
local_D = numpy.zeros(Nr)

for t in range(Nt-1):
    # in each time step solve System with Strang Splitting
    # Strang splitting: 0.5A + 0.5B + C + 0.5B + 0.5A
    wn = numpy.zeros((Nr, Ntau), dtype=numpy.float64)
    wn[:, 0] = w[:, t]

    for x in range(Nr):
        local_h[x] = h(T(r[x], t*dt))
        local_v[x] = v_i(v_hat(r[x], al))
        local_D[x] = D(r[x], t*dt)

        logging.info("t="+str(t))
        logging.debug("Calculate w(r,t)")

    # 0.5 INTEGRATION

for tau in range(int(Ntau/2)-1):
    for x in range(Nr):
        wn[x, tau+1] = wn[x, tau] + beta*local_h[x]*u[x, t]* (dtau)
logging.debug("RK1_w")
# set last value as start value
wn[:,0] = wn[:,int(Ntau/2)-2]

# 0.5 UPWIND
for tau in range(int(Ntau/2)-1):
    for x in range(1,Nr):
        # chain rule
        wn[x,tau+1] = (1 - n*local_v[x])*wn[x,tau] + n*
                      local_v[x-1]*wn[x-1, tau]

logging.debug("DC2_w")
# set last value as start value
wn[:,0] = wn[:,int(Ntau/2)-2]

kappa = dtau/(2.0*dr**2)

# CRANK NICOLSON
for tau in range(1,Ntau):
    temp = wn[:, tau-1] # get values of previous time
    b_r = numpy.zeros((Nr), dtype='float64')
    A = numpy.zeros((Nr,Nr), dtype='float64') # only
                      inner values
    for x in range(1,Nr-1):
        A[x,x] = ( 1 + 2*kappa*local_D[x] )
        A[x, x+1 ] = -kappa*local_D[x+1]
        A[x, x-1] = -kappa*local_D[x-1]
        b_r[x] = kappa*local_D[x+1]*temp[x+1] + (1 - 2* 
                      kappa*local_D[x])*temp[x] + kappa*local_D[x
                      -1]*temp[x-1]

    # use boundary values
    A[0,0] = ( 1 + 2*kappa*local_D[0])
    A[0, 1 ] = -kappa*local_D[1]
    A[Nr-1, Nr-1] = ( 1 + 2*kappa*local_D[Nr-1] )
# solve system of equations
wn_new = numpy.linalg.solve(A, b_r)

# save values from this time step
wn[ :, tau] = wn_new

logging.debug("CN3_w")
# set last value as start value
wn[ :, int(Ntau/2)-1] = wn[ :, Ntau-1]

# 0.5 UPWIND
for tau in range(int(Ntau/2)-1, Ntau-1):
    for x in range(1, Nr):
        # chain rule
        wn[x, tau+1] = (1 - n*local_v[x])*wn[x, tau] + n* 
        → local_v[x-1]*wn[x-1, tau]

logging.debug("DC2.2")
# set last value as start value
wn[ :, int(Ntau/2)-1] = wn[ :, Ntau-1]

# 0.5 INTEGRATION
for tau in range(int(Ntau/2)-1, Ntau-1):
    for x in range(Nr):
        wn[x, tau+1] = wn[x, tau] + beta*local_h[x]*u[x, t
        ↦ ]*(dtau)

# set next value of w
w[ :, t+1] = wn[ :, Ntau-1]

# end of time loop
w[0, :] = w[1, :]

return w