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Modeling of drug diffusion in a solid tumor leading to tumor cell death

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(Dated:)

It has been shown recently that changing the fluidic properties of a drug can improve its efficacy in ablating solid tumors. We develop a modeling framework for tumor ablation, and present the simplest possible model for drug diffusion in a porous spherical tumor with leaky boundaries and assuming cell death eventually leads to ablation of that cell effectively making the two quantities numerically equivalent. The death of a cell after a given exposure time depends on both the concentration of the drug and the amount of oxygen available to the cell, which we assume is the same throughout the tumor for further simplicity. It can be assumed that a minimum concentration is required for a cell to die, effectively connecting diffusion with efficacy. The concentration threshold decreases as exposure time increases, which allows us to compute dose-response curves. Furthermore, these curves can be plotted at much finer time intervals compared to that of experiments, which may possibly be used to produce a dose-threshold-response surface giving an observer a complete picture of the drug’s efficacy for an individual. In addition, the diffusion, leak coefficients, and the availability of oxygen is different for different individuals and tumors, we produce artificial replication data through bootstrapping to simulate error. While the usual data-driven model with Sigmoidal curves use 12 free parameters, our mechanistic model only has two free parameters, allowing it to be open to scrutiny rather than forcing agreement with data. Even so, the simplest model in our framework, derived here, shows close agreement with the bootstrapped curves, and reproduces well established relations, such as Haber’s rule.

I. INTRODUCTION

Historically, cancer has been treated using either generic-global drugs or by cutting away the infected cells via surgery. While success rates have increased, these types of treatments tend to have unwanted side-effects and are often quite expensive. This necessitates a new paradigm for cancer treatment. In recent years there has been a shift towards developing individualized and targeted drug therapy [1, 2]. However, there are still many technical and financial hurdles [3] to overcome before truly personalized medicine can be implemented in real-life situations. In an effort to alleviate these hurdles, data analysis techniques with statistical models have been employed [4]. With enough accurate data these models are able to make remarkable predictions, however one cannot expect the patient specific data to always be available and accurate. Furthermore, the mechanisms driving the phenomena are not captured in these statistical models. A reliable mechanistic modeling framework may alleviate the financial hurdles by simulating various treatment options, based on the physical rather than statistical properties of a patient’s tumor, before administering drugs. While data analysis has yielded promising results and should still be employed whenever possible, mechanistic models serve as another weapon in the fight against cancer.

In the seminal work of Sugiura et al. [5] it was shown that ethanol can successfully deteriorate malignant tissue. In fact, Ryu et al. [6] showed a comparable survival rate to surgery in a statistical survey of hepatocellular carcinoma patients treated with ethanol injections. As with any type of treatment, safety supersedes efficacy. An early work on the safety of injecting ethanol into parathyroid tumors was conducted by Solbiati et al. [7]. For the past decade there have been numerous studies on the safety and efficacy of ethanol ablation for different types of tumors [8–12]. Some major drawbacks of the technique include the need for multiphase treatments, large amounts of fluid, and the rapid escape of ethanol in non-capsulated tumors [13]. Traditionally these drawbacks were greater for larger tumors, but more recently it has been shown by Kuang et al. [11] that a single-session high-dose ethanol injection can ablate hepatocellular carcinoma tumors of diameters up to 5 cm.

Recently, Morhard et al. [13] developed a new method employing ethyl cellulose for which single-phase small-volume treatments suffice to trigger ablation. Their goal is to maximize distribution of the ethanol throughout the tumor while minimizing the necessary solution volume. In essence, by changing the viscosity of the drug mixture, and hence its rate

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of diffusion, they are able to increase efficacy and decrease toxicity – the goal of cancer treatments. While this type of optimization of fluidic properties has only been used with ethanol injections, one may see the potential of using it for other drugs as well. After all, there is a wide variety of drug injection therapies [14–16].

Although it may seem obvious that drug transport occurs as a diffusive process, it is still worth noting that there is an abundance of drug diffusion models. Recent review articles have discussed diffusion-based models of drug distribution from the blood stream into solid tumors [17, 18]. In [18], Kim et al. advocate integrating mechanistic models with clinical data to improve understanding and prediction. In addition, quite sophisticated transport models have been developed for penetration into a solid tumor [19–21]. However, it is evident that drug distribution from an internal injection has not been investigated mathematically. Further, it is often the case that mathematical models describe either drug transport or cell death, but seldom both. In this work we endeavor to build a new modeling framework for injected drug distribution in solid tumors and its effect on tumor cell death, which we hope will facilitate individualized treatment. This is done by producing the simplest possible model in this framework, and laying bare the simplifying assumptions in order to open it to scrutiny and improvement.

The remainder of the paper is organized as follows: we start by discussing the modeling procedure, the focus of the article, in Sec. II and III. Section II contains the diffusion model for drug distribution, and Sec. III relates the concentration profile from diffusion to cell death. While the analytical solutions to the diffusion equation would generally be a linear combination of Bessel functions, since our initial condition is not a Bessel function a numerical method is required to match the coefficients of the Bessel functions. Instead we use finite difference methods in Sec. IV to numerically solve the equation from the start, and using this solution for the concentration profile the cell death is computed. In Sec. V we compare our dose-response curves with data points from experiments different from that of Morhard et al. [13] since they did not include dose-response curves in their article. Furthermore, using a different experiment shows the robustness of this type of modeling. Finally, we conclude with a discussion about applying our modeling framework to individualized treatment in Sec. VI.

II. RADIAL SYMMETRIC
CONCENTRATION DIFFUSION MODEL

In this section we derive the simplest possible model for drug diffusion in a tumor with leaky boundaries where injection occurs at a much faster timescale than diffusion. First, let us assume a spherical tumor with constant density. The drug concentration is given as the ratio of the volume of the drug to that of the tumor. Since this is a diffusive process the initial condition is taken as a compact Gaussian. Then the leak at the boundary is modeled as “Newton’s law of cooling” [22]. Finally, we nondimensionalize the equation and conditions to further simplify the model.

Consider the radially symmetric diffusion equation in spherical coordinates with constant diffusivity

\[
\frac{\partial u}{\partial t} = D \left( \frac{\partial^2 u}{\partial r^2} + \frac{2 \partial u}{r \partial r} \right)
\]

where \(u\) is the concentration of the drug and \(D\) is assumed to be the constant effective diffusivity of transport through porous media, which is inversely proportional to the viscosity of the drug mixture. Since we are injecting a volume into the center of the tumor and the diffusive timescale is fast, it is reasonable to assume an initial compact Gaussian distribution (i.e., a bump function) where the tail just reaches the boundary

\[
u(r, t = 0) = \frac{V_0}{V_b} \begin{cases} \exp \left(1 - \frac{R^2 - r^2}{\gamma}ight) & \text{for } r < R, \\ 0 & \text{for } r \geq R; \end{cases}
\]

where \(R\) is the tumor radius, \(V_0\) is the ratio of the injected drug volume to the tumor volume, and \(V_b\) is the volume of the bump function in spherical coordinates used for the purpose of normalization; that is,

\[V_b = \int_0^\pi \int_0^{2\pi} \int_0^R \exp \left(1 - \frac{R^2}{R^2 - r^2} \right) drd\theta d\phi\]

The generic bump function takes the 1-dimensional shape shown in Fig. 1, and by radial symmetry this can be extended to 3-dimensions. At the boundary we know the drug leaks due to the porosity of the tumor. However, diffusion in the tumor is different from diffusion outside of the tumor. In fact, the tumor is far denser than healthy cells, and therefore will hold onto the drug and let it accumulate as it leaks at a much slower rate. We assume the drugs start leaking at the boundary of the tumor according to “Newton’s law of cooling” [22],

\[
D \frac{\partial u}{\partial r} \bigg|_{r=R} = -\gamma u(r = R, t),
\]
where $\gamma$ is the leak coefficient.

### A. Nondimensionalization

In order to simplify the model and reduce the number of parameters into only the essential relations we nondimensionalize the problem. The concentration $u$ (ratio of drug volume to tumor cell volume) is by definition nondimensional. Therefore, we need only do a change of variables on $t$ and $r$. Let $\hat{r} = r/R$ and $\hat{t} = t/\tau$, which gives us the derivatives

$$
\frac{\partial}{\partial \hat{t}} = \frac{1}{\hat{r}} \frac{\partial}{\partial \hat{r}}; \quad \frac{\partial}{\partial r} = \frac{1}{R} \frac{\partial}{\partial \hat{r}} \Rightarrow \frac{\partial^2}{\partial r^2} = \frac{1}{R^2} \frac{\partial^2}{\partial \hat{r}^2}.
$$

Then our PDE becomes

$$
\frac{\partial u}{\partial \hat{t}} = D \tau \left( \frac{\partial^2 u}{\partial \hat{r}^2} + \frac{2 \partial u}{r \partial \hat{r}} \right),
$$

where $\tau = R^2/D$ is the diffusive timescale, which is directly proportional to the viscosity of the fluid mixture. Our initial condition becomes

$$
u(\hat{r}, t = 0) = \begin{cases} 
\hat{V}_0 & \text{for } \hat{r} < R/R, \\
0 & \text{for } \hat{r} \geq R/R;
\end{cases}
$$

where $\hat{V}_0$ is the injected nondimensional volume and $\hat{V}_b$ is the volume of the nondimensional bump function. Furthermore, the boundary condition becomes

$$
\left. \frac{D \partial u}{R \partial \hat{r}} \right|_{\hat{r}=1} = -\gamma u(\hat{r} = 1, t) \\
\Rightarrow \left. \frac{\partial u}{\partial \hat{r}} \right|_{\hat{r}=1} = -\frac{R \gamma}{D} u(\hat{r} = 1, t) = -\varepsilon u(\hat{r} = 1, t),
$$

where $\varepsilon = R \gamma/D$ is the nondimensional leak coefficient, which is directly proportional to the viscosity of the drug mixture.

Removing the hats gives us the full nondimensional problem

$$
(4a) \quad \frac{\partial u}{\partial \hat{t}} = \frac{\partial^2 u}{\partial \hat{r}^2} + \frac{2 \partial u}{r \partial \hat{r}}; \\
(4b) \quad \frac{\partial u}{\partial \hat{r}} \bigg|_{\hat{r}=1} = -\varepsilon u(\hat{r} = 1, t); \\
(4c) \quad u(\hat{r}, 0) = \begin{cases} 
\frac{\hat{V}_0}{\hat{V}_b} \exp \left(1 - \frac{1}{1-r^2} \right) & \text{for } r < 1, \\
0 & \text{for } r \geq 1;
\end{cases}
$$

### III. RELATING CONCENTRATION TO CELL DEATH

Once we have a model for drug distribution, we may use it to analyze cell death. In biological experiments this is done through dose-response curves (amount of drug administered vs percent cell death at a specific time after administration) [23, 24]. The dose-response is recorded by measuring the effect of a drug at various doses. A mathematical model allows us to do the same, but for far more finely spaced doses and exposure times, thereby allowing an observer to find an optimal treatment strategy for a specific individual.

It should be noted that diffusion happens on a very fast timescale, on the order of seconds, in contrast to the exposure timescale, which is on the order of days. Diffusion provides the amount of exposure the tumor has to the drug, which then triggers the mechanism for cell death (e.g., apoptosis) [25]. If the concentration of a drug in a certain region of the tumor is high enough to trigger cell death, most of the cells in that region will most likely die after some exposure time $\eta$, but often this is not instantaneous and may take days.

While varying oxygen levels are a concern [26–29], as a first approximation and “proof of concept” model, we assume that the oxygen concentration throughout the tumor is constant for a given time. Therefore, the same amount of drug will kill a tumor cell no matter where in the tumor it is. We define a drug concentration threshold that is required to kill a cell for a given time, which is dependent on dose strength.

**Definition 1.** The minimum concentration $u_T(\eta, d)$ required to kill one cell after an exposure time $\eta$ hours at a given dose $d$ is said to be the concentration threshold of the tumor at time $\eta$. 
Variable oxygen availability may be implemented by prescribing a threshold \( u_T \) with radial dependence in addition to temporal and dose dependence, which is the focus of a future study.

Now we must relate the threshold \( u_T \) to time at a given dosage explicitly in order to compare against experimental data. Since log time is often used for time-response curves and time-dose-response surfaces, we assume the threshold to be related to time in a negative exponential manner,

\[
 u_T(\eta, d) = a - b \exp(-c \eta).
\]

Employing the three most used time points in oncology, \( \eta = 24, 48, 72 \) hours, we solve for the constants at a fixed initial concentration to get

\[
 u_T(\eta) = a - be^{-c\eta};
 a = \frac{u_T(24)u_T(72) - u_T(48)^2}{u_T(24) + u_T(72) - 2u_T(48)};
 b = \frac{(a - u_T(24))^2}{a - u_T(48)};
 c = \frac{1}{24} \ln \left( \frac{a - u_T(24)}{a - u_T(48)} \right).
\]

The plot of (5) is shown in Fig. 2. To produce Fig. 2, we use the specific data set in Sec. V, but this can be done in general by using three dose points at a given time. Once the three dose points are identified we numerically approximate the threshold \( u_T \) required for the model to match the response given as apoptosis fraction at the experimental dose points, which is accomplished by simply varying \( u_T \) until an error of 0.01 is reached between the numerical and experimental apoptosis fraction.

In addition to tumor cell death, most drugs also kill healthy cells, which contributes to toxicity. While the current format does not handle toxicity as the leakage equation (3) assumes the drug leaks into a vacuum. However, in real life the drug would leak into healthy cells that have completely different physical properties from tumor cells. This would require analyzing what is effectively two-phase diffusion, which is the subject of a future work. Regardless, one naive approach to handling toxicity as a function of injected volume is to define it as the maximum distance at the boundary between the concentration profile of the drug \( u(\eta, r = 1) \), and a prescribed toxicity threshold \( u_{\text{toxicity}} \). That is,

\[
 \text{Toxicity}(V_0) := \max(u(\eta, r = 1) - u_{\text{toxicity}}),
\]

whenever \( u(\eta, r = 1) > u_{\text{toxicity}} \), and zero otherwise. Again, since the drug is leaking into a vacuum, the leakage is a linear process, and hence we expect toxicity to be linear with \( V_0 \) as shown in Fig. 3.

IV. NUMERICAL SIMULATION OF SINGLE-TIME DOSE-RESPONSE CURVE

In order to simulate the dose-response curves one must first solve (4). Since our problem is radially symmetric, we may solve it on the spacial domain \( r \in [0, 1] \) by adding an additional “boundary” condition at the origin, namely \( \frac{\partial}{\partial r} u(0, t) = 0 \). Although a Bessel function solution can be found, matching it with the initial conditions in the model would not yield analytical solutions, and it would be necessary to solve the integrals from the inner products numerically. Therefore, we implement finite difference schemes to solve (4) instead of solving for the constants in the Bessel function solution.
We compare the solutions from three stencils: Forward in Time Centered in Space (FTCS), Backwards in Time Centered in Space (BTCS), and Crank-Nicolson. For diffusion problems, Crank-Nicolson is often used, however diffusion is only the base of our model, and we often need to run diffusion $10^5$ to $10^8$ times for one plot, which renders Crank-Nicolson inefficient. This application requires a fast, accurate, and stable solver. FTCS is certainly fast, but it is at the mercy of the CFL condition [30]; i.e., we require $\Delta t/(\Delta x)^2 \leq 1/2$. For the videos illustrating the scheme we do not need high accuracy, but we do need a fast scheme for high spatial resolution, so FTCS is employed. To produce the dose-response curves in Sec. V we employ BTCS since it is not much slower than FTCS and almost as accurate as Crank-Nicolson. This way we get high temporal resolution, which has the added benefit of providing accurate dose-response curves, without sacrificing stability.

Once a solution for $u$ from the finite difference schemes is obtained, the dose-response curves can be plotted. As outlined in Sec. III, in biological experiments the dose-response is recorded by measuring the effect of a drug at various doses. In our model we implement a threshold $u_T$ defined as the drug concentration required to kill a single cell. When the concentration $u$ is above this threshold on some interval $r \in [0,r_{death}]$, we may calculate the volume of tumor death, which we assume eventually ablates. Dividing this volume by the total nondimensional volume of $4\pi/3$ outputs the fraction of dead tumor cells, namely

$$P_{\text{dead}} := \frac{\text{Vol. Dead}}{\text{Total Vol.}} = \frac{4\pi r_{\text{death}}^3/3}{4\pi (r = 1)^{3}/3} = r_{\text{death}}^3. \quad (7)$$

This is demonstrated in Fig. 4. Where the concentration profile (blue curve) intersects the threshold $u_T(\eta, d)$ is the radius of cells killed after exposure time $\eta$, which we denoted as $r_{\text{death}}$.

There are three possible cases for the percent of the tumor that is killed, and hence eventually ablated, at an exposure time of $\eta$: no ablation, partial ablation, and full ablation. These are shown in Figs. 5-8.

In Fig. 5, the initial volume is so low that the height of the bump function does not exceed the threshold. This provides an opportunity to calculate the least effective dose at a given exposure time; i.e. when the height of the bump function is equivalent to the threshold (illustrated in Fig. 6a). Notice that the height of the bump function (4c) at the center $r = 0$ is $V_0/V_b$, where $V_0$ is the initial volume scaled as a fraction of the tumor volume and $V_b$ is the volume of the bump function. If we let the height be $u_T$ then the least effective dose is

$$u_{\text{LED}} = u_T \cdot V_b = u_T \int_0^{\pi} \sin \phi d\phi \int_0^{2\pi} d\theta \int_0^1 r^2 \exp \left( 1 - \frac{1}{1 - r^2} \right) r^2 dr$$

$$= 3u_T \int_0^1 \exp \left( 1 - \frac{1}{1 - r^2} \right) r^2 dr = 3 \left( a - be^{-cn} \right) \int_0^1 \exp \left( 1 - \frac{1}{1 - r^2} \right) r^2 dr. \quad (8)$$

From this analytical formula we may plot the least effective dose as a function of exposure time in Fig. 6b. Furthermore, we observe that the steady-state least effective does is given by

$$u_{\text{LED}} = u_\ast \cdot V_b = 3 \cdot a \cdot V_b = \frac{3(u_T(24)u_T(72) - u_T(48)^2)}{u_T(24) + u_T(72) - 2u_T(48)} \int_0^1 e^{-1} \frac{1}{1 - r^2} r^2 dr. \quad (9)$$

Figure 7 shows the death of a fraction of the tumor cells, and hence, by our initial assumption, partial ablation. An exaggerated initial condition (Fig. 7a) is used to illustrate the diffusion of the drug. In order to present a profile similar to that of (4c) (Fig. 7b), we let the volume diffuse until the concentration at the boundary is above a small, but appreciable, amount of $u = 0.01$. This is due to the discrepancy between diffusion in real life, which happens in finite time, and mathematical diffusion, which occurs infinitely fast. After a maximum radius of death is attained (Fig. 7c), the percent cell death is calculated using the volume filled at this radius. After this maximum, the drug leaks out (Fig. 7d) and
FIG. 4. Hypothetical concentration profile used to demonstrate percent tumor death calculation. The blue (solid) curve represents the concentration profile of the drug, and the red (dashed) line represents the concentration threshold required to kill a cell. The radius \( r_{\text{death}} \) is taken at the intersection of the two curves and is used to calculate the percent volume and hence percent cell death.

FIG. 5. No ablation occurs, at time \( \eta \), when the height of the initial profile is lower than the threshold \( u_T \). This leads to an explicit formula for least effective dose shown in (8).

Eventually contributes to toxicity, which is left for a future study.

Finally, Fig. 8 shows complete tumor death for a threshold \( u_T \) at an exposure time \( \eta \). As in Fig. 7, an exaggerated initial condition (Fig. 8a) is used, and later a snapshot of a profile similar to (4c) (Fig. 8b) is shown. Since the concentration in the entire tumor exceeds the threshold, the entire tumor dies after a time \( \eta \).

Eventually all of the drug leaks out, but for the full ablation case the toxicity would be higher than with partial ablation. Then we have a balancing act between efficacy and toxicity. We may calculate the smallest dose required to kill the entire tumor at an exposure time \( \eta \) using the threshold \( u_T \). Since the diffusion and leak are linear, we would expect the relation between the threshold and smallest dose to be linear as well, which is precisely what we observe. If we let \( \Gamma_i \) be the smallest dose required to kill the tumor after a time \( \eta_i \) we get the following relation between the ratios of the smallest dose and the respective threshold

\[
\frac{\Gamma_i}{\Gamma_j} = \frac{u_T(\eta_i)}{u_T(\eta_j)}.
\]  

(10)

Since \( u_T \to u_\ast \) as \( T \to \infty \), we may define an optimal steady-state dose as the smallest dose required to kill the tumor assuming we wait infinitely long and the tumor growth is completely inhibited. By this definition we notice that we may calculate \( \Gamma \) for any
FIG. 7. Partial ablation at an exposure time \( \eta \). The left column represents the diffusion viewed as a projection onto \( \mathbb{R}^2 \). The right column represents the concentration profile (blue solid curve) compared to the threshold (red dashed line). An exaggerated initial condition was used in order to illustrate the diffusion of the drug in (a), but the initial condition from the model (4) is more like the profile in (b). (a) Exaggerated initial condition to illustrate the diffusion of the drug. (b) Profile similar to that used for the initial condition in (4c). (c) Maximum radius of effective concentration is attained. (d) The drug eventually leaks out contributing to toxicity. See ref. [31] for the corresponding animation.

\[
\Gamma = \left( \frac{u_T}{u_*} \right) \Gamma_* \Rightarrow \Gamma = \frac{u_T}{u_*} \frac{\Gamma_*}{u} = \left[ a - be^{-c\eta} \right] \frac{\Gamma_*}{u_*},
\]

which is precisely the modified Haber’s rule \([33]\) because \( \Gamma_*/u_* \) is constant. In addition, we may fix a threshold \( u_T \), and observe how the optimal dose for that threshold varies with the nondimensional leak coefficient \( \varepsilon \), which we recall is directly proportional to the viscosity of the drug mixture. We may calculate the “optimal dose” for different leak coefficients, which represents different tumors and drugs, by using a bisection method as the leak coefficient \( \varepsilon \) is varied. We observe in Fig. 9 that this relation, represented by the blue curve, is not linear although it may seem linear without the presence of the red dashed line from the lowest to the highest point of the blue curve in that plot.

V. COMPARISON WITH CLINICAL DATA

While there was no data on the rate of cell death in the study by Morhard et al. [13] because it focused on tumor shrinkage, we may use other clinical/experimental data sets to show qualitative agreement. This would indicate the modeling framework is reasonable, and then the models may be modified for different applications. In this light, a “proof of concept” comparison is made between the model and a sample data set obtained from the LINCS database (http://lincs.hms.harvard.edu/db/) which, among other things, offers cellular responses to chemical and genetic perturbations \([34]\). In this article we consider apoptosis fraction as the cellular response observed under different dosage of different compounds. So, our experimental data consists of a set of dose-response pairs observed for several cell-lines and compounds at 24, 48 and 72 hours.

Since the experiments are so different and don’t report all the necessary physical properties, calibration between the two types of dosage and the model is necessary. The Morhard et al. [13] experiments vary the injection volume (as a percentage of tumor volume), and the LINCS database \([34]\) varies the molarity. Morhard et al. also run the experiments for a variety of viscosities, but the LINCS database does not include information about the fluidic properties of the drug. As we wish to have as few free parameters as possible, let us keep the viscosity, and hence the leak coefficient constant. Then we simulate dose as an injected volume in Sec. VA and molarity in Sec. VB.
FIG. 8. Full ablation at exposure time \( \eta \). The left column represents the diffusion viewed as a projection onto \( \mathbb{R}^2 \). The right column represents the concentration profile (blue solid curve) compared to the threshold (red dashed line).

(a) An exaggerated initial condition was used in order to illustrate the diffusion of the drug in (a), but the initial condition from the model (4) is more like the profile in (b). (a) Exaggerated initial condition to illustrate the diffusion of the drug. (b) Profile similar to that used for the initial condition in (4c). (c) Full ablation is attained, but just as in Fig. 7, the remainder of the drug will leak out and contribute to toxicity. See ref. [32] for the corresponding animation.

A. Simulating dose as injected volume

In the experiments of Morhard et al. [13], for a given viscosity, the volume of the solution is varied. Since no information is given about the molarity in the Morhard et al. experiments and no information about the volume is given in the LINCS data [34], we need to assume a correlation between molarity and volume. In order to simulate dose as injected volume we match the 1\( \mu \)M dose from the LINCS data [34] with an injected volume of 1/12 tumor volume. By employing (5) we find the concentration thresholds at the three time points for this initial injected volume. Furthermore, we match two other doses by varying the initial volume until the distance between two out of the three time points is minimized. We choose to match two time points since it is evident that the data set is quite noisy and it has outliers that do not obey the Hill equation [23]. One concern may be the nonuniformity of the initial volume - dose relation, but this is to be expected since the relation is dependent on the tumor size and concentration of the drug in the solution. The dose-response curves from the model and the observed data points are shown in Fig. 10.

In this section, using a leak coefficient of \( \varepsilon = 5 \), we illustrate the performance of the model for cell line “C32” and drug “Selumetinib” (the performance for the remaining cell lines are illustrated in Appendix A). Since the data is quite noisy for low doses a truncated data set, shown in Table I, is much more reliable for comparisons with the model. The responses for a dose of 1\( \mu \)M from the pair is matched with 1/12 tumor volume from the model. Equation 5 gives us the relation between the three time points (24, 48, and 72 hours) and the thresholds \( u_T(24) \), \( u_T(48) \), and \( u_T(72) \). These thresholds are then used to produce the simulated dose-response curves in Fig. 10. The data from the 1\( \mu \)M dose will match up exactly with the curves. In order to match the data from the 0.316\( \mu \)M and 3.16\( \mu \)M doses with the respective initial concentration in the model we identify the concentration for which the difference between the response in the data and the model of two time points is minimized. For example, in Fig. 10, the 0.316\( \mu \)M
TABLE I. Truncated data set for cell line “C32” with drug “Selumetinib”

<table>
<thead>
<tr>
<th>Dose (micromole)</th>
<th>Time (hours)</th>
<th>Response (apoptosis fraction)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.316</td>
<td>24</td>
<td>0.2148</td>
</tr>
<tr>
<td></td>
<td>48</td>
<td>0.262</td>
</tr>
<tr>
<td></td>
<td>72</td>
<td>0.4026</td>
</tr>
<tr>
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<td>24</td>
<td>0.2365</td>
</tr>
<tr>
<td></td>
<td>48</td>
<td>0.5982</td>
</tr>
<tr>
<td></td>
<td>72</td>
<td>0.7591</td>
</tr>
<tr>
<td>3.160</td>
<td>24</td>
<td>0.2716</td>
</tr>
<tr>
<td></td>
<td>48</td>
<td>0.8303</td>
</tr>
<tr>
<td></td>
<td>72</td>
<td>0.8621</td>
</tr>
</tbody>
</table>

FIG. 10. Simulated dose-response curves for cell line “C32” and drug “Selumetinib” compared to experimental data points (stars). The blue curves and markers are at 24 hours (bottom), red is 48 hours (middle), and black is 72 hours (top). Molarities from the LINCS data [34] have been converted to initial volumes.

Dose corresponds to an initial concentration of approximately $1/20$ of the tumor volume since the 72 hour and 48 hour time points are closest to the curve at that dose, and similarly the $3.16\mu M$ corresponds to approximately $7/50$ of the tumor volume since the 72 hour and 24 hour time points are closest. Since 72 hours is near the steady state of this process, the curves often match the data at that time point. For the 24 and 48 hour time points, the data becomes quite noisy with outliers and responses that would not be possible to achieve, indicating possibly large errors. If the distance between all three points were minimized, these outliers would have a stronger effect.

Besides the foregoing three dose points we assume that at a dose of zero the response should be around zero. Once we match the original concentration levels with the tumor volumes, we empirically fit a sigmoidal curve to the experimental dose-response data observed at 72 hours. The empirical model is given by

\[
\text{Apoptosis fraction} = a + \frac{\Delta}{1 + \left(\frac{\delta}{\text{concentration}}\right)^\theta} \quad (12)
\]

where $a$ is the lower asymptote of the response curve, $\Delta$ is the range of response, $\delta$ is typically interpreted as $EC_{50}$ and $\theta$ is the Hill coefficient. The parameters of the Hill equation (12) are estimated using a non-linear least squares fit. We then generate 50 replicates from the fitted sigmoid curve using a parametric bootstrap procedure with errors drawn from a Gaussian process with an exponential correlation function. The noise variance is comparable to the residual variance obtained from the least square fit (12) and the range parameter is chosen to ensure smoothness and monotonicity of each replicate. Figure 11 shows 50 bootstrap replicates for cell line “C32” and drug “Selumetinib”. It should be noted that molarities from the LINCS data [34] have been converted to initial volumes.

FIG. 11. Fifty sigmoidal curves, defined as (12), fitting the empirical data, which is given a Gaussian spread to simulate error at each dose point, for cell line “C32” and drug “Selumetinib” at 72 hours of exposure time. Molarities from the LINCS data [34] have been converted to initial volumes.

From these bootstrap replicates of sigmoidal
curves we compute the point-wise 95% confidence intervals at each matched dose, then join them into a piece-wise linear approximate confidence band. An example of this is shown in Fig. 12 for the foregoing illustrative case. The dose-response curve produced by our model is represented as the black curve, and the point-wise empirical quantiles are connected by the piece-wise linear dashed curves. Observe that the curve produced by our model is contained within the 95% pointwise confidence interval, indicating statistical adequacy of the the deterministic model. The remainder of the data pairs are relegated to Appendix B.

Figure 12. Theoretical dose-response curve for cell line “C32” and drug “Selumetinib” within a 95% confidence band at 72 hours of exposure time. The black (solid) curve represents the dose-response from the model. The blue (bottom) and red (top) dashed lines represent the piece-wise linear curve connecting the lower and upper quantiles of each of the 95% confidence intervals. Molarities from the LINCS data [34] have been converted to initial volumes.

We would like to point out that since we are computing pointwise confidence intervals, there could be situations where for some dose levels our model output lies outside the 95% confidence interval computed at those doses. Rare occurrences (less than 5% of the times) of such instances do not necessarily indicate statistical inadequacy of the posited model. However, frequent occurrences of such events (more than 10% of the times) would indicate statistical inadequacy.

Given the empirical evidence that our model is statistically adequate to generate dose-response curves from experimental data, we proceed to compute the theoretical EC50s in terms of initial volumes for this dataset. The model is cut off at the maximum empirical response, and the EC50s, by definition, are computed as the dose to achieve half that response. We use a bisection scheme similar to that of the optimal dose. EC50s are then approximated within a maximum response error of $|u_{\text{approx}} - u(\text{EC50})| < 0.0025$. The spatial resolution of the diffusion scheme will limit the error bound, so this must be chosen to be equivalent or less than the desired error. Table II shows the theoretical EC50s and the associated errors.

B. Simulating dose as molarity

In contrast to the previous section, we may also produce dose response curves for the molarity as opposed to the injected volume by making the assumption that the amount of solution required to kill a cell decreases as molarity increases. In other words, if we increase molarity we lower the concentration threshold. To calibrate this from the LINCS data set [34], reliable data at a particular time point is necessary, which for most drugs will be that of 72 hours. For that time point the required threshold to match the data at three doses, $u_T(\eta = 72, d_1)$, $u_T(\eta = 72, d_2)$, $u_T(\eta = 72, d_3)$, can then be numerically approximated. It is assumed that the threshold will obey the Hill equation [23],

$$u_T(\eta = 72, d) = u_\infty + \frac{h - u_\infty}{1 + (\sigma/d)^\theta}, \quad (13)$$

where $d$ is the dose and $h$ is the height of the profile of the initial condition (4c). The three free parameters to be fitted using the three dose points $u_T(\eta = 72, d_i)$ are $u_\infty$, which is the lower asymptote for the threshold as $d \to \infty$, $\sigma$, and $\theta$. The lower asymptote, $u_\infty$, is approximated by first assuming $u_\infty = u_T(\eta = 72, d = 3.16)$, but if this were the lower asymptote, $u(\eta = 72, d = 3.16)$ would never match with the data. So $u_T$ is decreased until $u(\eta = 72, d = 3.16)$ is close to the data within the chosen error tolerance. Further, the other terms are given by,

$$\sigma = \exp \left( F_i \ln d_j - F_j \ln d_i \right) / F_i - F_j,$$

$$\theta = \frac{F_i}{\ln \sigma/d_j},$$

$$F_i := \ln \frac{h - u_T(\eta = 72, d_i)}{u_T(\eta = 72, d_i) - u_\infty},$$

where $i$ and $j$ denote two of the other dose points. The function for (13) can be computed for cell line “C32” and drug “Selumetinib” using data from Table I in Sec. VA, which is illustrated in Fig. 13.

As in the previous section, the performance of the model for cell line “C32” and drug “Selumetinib” is illustrated using data from Table I (the performance for the remaining cell lines are illustrated
TABLE II. Theoretical EC50s with response errors

<table>
<thead>
<tr>
<th>Cell line</th>
<th>Drug</th>
<th>EC50</th>
<th>Response error</th>
</tr>
</thead>
<tbody>
<tr>
<td>C32</td>
<td>SB590885</td>
<td>0.0562</td>
<td>0.0006</td>
</tr>
<tr>
<td></td>
<td>PLX-4720</td>
<td>0.0684</td>
<td>0.0015</td>
</tr>
<tr>
<td></td>
<td>AZ-628</td>
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<td>0.0007</td>
</tr>
<tr>
<td></td>
<td>Selumetinib</td>
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<td>0.0007</td>
</tr>
<tr>
<td></td>
<td>Vemurafenib</td>
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<td>0.0017</td>
</tr>
<tr>
<td>KOLO 858</td>
<td>SB590885</td>
<td>0.1055</td>
<td>0.0015</td>
</tr>
<tr>
<td></td>
<td>PLX-4720</td>
<td>0.1260</td>
<td>0.0005</td>
</tr>
<tr>
<td></td>
<td>AZ-628</td>
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<td>0.0019</td>
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<tr>
<td></td>
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<td>0.0014</td>
</tr>
<tr>
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<td>AZ-628</td>
<td>0.1191</td>
<td>0.0013</td>
</tr>
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<td></td>
<td>Selumetinib</td>
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<td>0.0002</td>
</tr>
<tr>
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<td>SB590885</td>
<td>0.1074</td>
<td>0.0021</td>
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<td>Selumetinib</td>
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<tr>
<td></td>
<td>Vemurafenib</td>
<td>0.1230</td>
<td>0.0023</td>
</tr>
</tbody>
</table>

FIG. 13. Relation between dose and concentration threshold from (13) for parameter values $u_\infty = 0.045$, $\sigma = 0.1689$, and $\theta = -2.0476$. The 72 hour time point produces the parameter values $u_\infty = 0.045$, $\sigma = 0.1689$, and $\theta = -2.0476$. The dose-response curves may then be produced by using (13) for the 72 hour time point threshold values and multiplying it by the ratio of the other thresholds to the 72 hour threshold at the 1µM dose point from (5); that is,

$$u_T(\eta, d) = u_T(\eta = 72, d) \frac{u_T(\eta, d = 1)}{u_T(\eta = 72, d = 1)}. \quad (14)$$

The aforementioned dose-response curves are presented in Fig. 14.

Since the dosage has been changed from volume to molarity the sigmoidal curves will also be slightly different from those in Fig. 11. Using (12) we produce the fifty sigmoidal curves with Gaussian spread using molarities for the dosage for cell line “C32” and drug “Selumetinib” in Fig. 15.

Further, as was done in Sec. V A, we produce 95% confidence intervals at each relevant dose point from the sigmoidal curves of Fig. 15. These intervals are then connected in a piece-wise linear manner to produce approximate 95% confidence bands, which have less variance than a true 95% confidence band. Observe that in Fig. 16 the dose-response curve for cell line “C32” and drug “Selumetinib” is contained within the confidence band (dashed blue and red lines). The curves for the re-
FIG. 15. Fifty sigmoidal curves, defined as (12), fitting the empirical data, which is given a Gaussian spread to simulate error at each dose point, for cell line “C32” and drug “Selumetinib” at 72 hours of exposure time.

remaining cell line and drug pairs are relegated to Appendix B.

FIG. 16. Theoretical dose-response curve for cell line “C32” and drug “Selumetinib” within a 95% confidence band at 72 hours of exposure time. The black (solid) curve represents the dose-response from the model. The blue (bottom) and red (top) dashed lines represent the piece-wise linear curve connecting the lower and upper quantiles of each of the 95% confidence intervals.

In Sec. VA the EC50s were numerically approximated from the model due to the use of initial injected volumes of the drug as the initial condition. However, in this section we keep molarity for the dosage, and hence the EC50s can simply be approximated from the data directly without having to invoke the model.

C. Dose-threshold-response surface

While a dose-response curve may be easily fit to the empirical data, a reliable dose-time-response surface is much more difficult due to how noisy the data is. With a mechanistic model we get a full dose-threshold-response surface (Fig. 17) for a particular nondimensional leak coefficient (dependent on the density of the tumor and fluidic properties of the drug), which can be turned into an individual’s dose-time-response surface by using (5) in Sec. III to relate the threshold to time.

We observe that the surface in Fig. 17 is qualitatively similar to that of Miller et al. [33], where they fit empirical data to Haber’s rule. This further solidifies the agreement between our new model and previous studies.

FIG. 17. Simulated dose-threshold-response surface for a specific nondimensional leak coefficient, which is dependent on an individual tumor density and viscosity of the drug mixture. This can be converted into a dose-time-response surface by calibrating the time-threshold relation, (5) in Sec. III, for each cell line and drug.

VI. CONCLUSION

Cancer treatments have come a long way in terms of their increased efficacy and reduced toxicity. However, it is still very much a “trial and error” process. Since individuals contain such a variety of biological properties, and in fact separate tumors in the same individual are also quite different, we need to develop better individualized treatments. While there is an abundance of statistical models [4], there are few mechanistic models detailing the distribution of drugs inside a tumor leading to tumor cell death. There has been recent work on transport models for
drugs into solid tumors [17–21], but there are no models in the literature that describe the diffusion from an injection into a solid tumor causing it to ablate.

In this investigation we developed a radially symmetric concentration diffusion model in a solid porous spherical tumor with leaky boundaries in Sec. II, which is then nondimensionalized to absorb the constants (radius, diffusivity, leak coefficient) into the nondimensional leak coefficient, effectively making it a one parameter model. Then in Sec. III, the concentration from the diffusion model is related to cell death assuming a fixed constant concentration required to kill a cell after an exposure time $\eta$, which we call the concentration threshold, $u_T$. Three time points for a cell line, at a specific initial dose, are used to relate the threshold to time using (5). Numerical simulations used to produce dose-response curves are described in Sec. IV, and the curves themselves are illustrated in Sec. V, where they are tested against empirical data from the LINCS database [34].

While [13] did not include dose-response curves in their study, we use a different empirical data set to show qualitative agreement and use it as a “proof of concept” comparison. The comparison with a completely different data set solidifies the argument that this is a modeling framework that can be applied to other experiments and not just a model for the Morhard et al. experiments [13]. In applications the nondimensional leak coefficient would be found empirically from the individual tumor’s porosity, density, and tortuosity, and the drug mixture’s viscosity. We envision that the effective diffusivity and dimensional leak coefficient may be empirically calculated by using a nontoxic fluid with the desired viscosity, effectively bypassing the need to directly analyze the tumor’s physical properties. Then the model can be used to simulate the affect of the drug at varying viscosity for that individual. In this study, due to the lack of availability of such information, we fix the nondimensional leak coefficient and approximate the thresholds necessary to match three dose points at a particular exposure time. This function for the threshold is then used to produce the curves at all other exposure times. Since this can theoretically be done at all exposure times, if an individual tumor/drug combination’s nondimensional leak coefficient is known, a dose-time-response surface can be created.

It is true that, due to its simplicity, the model presented here misses some of the behavior associated with the diffusion of a drug leading to cell death. Critics may even suggest that it is naive. It is completely legitimate to call into question the assumption that the tumor is a uniformly dense sphere (i.e., constant diffusivity and radial symmetry). One may also point to the use of a constant (in space, but not in time) concentration threshold for the death of a tumor cell since oxygen availability varies in the tumor [26–29]. However, scrutiny is precisely what we seek because this will lead to more sophisticated models, but these will still be in the framework of drug diffusion inside a solid tumor leading to tumor cell death, and eventually ablation. We shall endeavor to develop more models in this framework, and we hope that other researchers will as well.

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See supplemental material at [url will be inserted by publisher] for an animation of full ablation of a tumor.

See supplemental material at [url will be inserted by publisher] for an animation of partial ablation of a tumor.

See supplemental material at [url will be inserted by publisher] for the LINCS database excell sheet.
Appendix A: Dose-response curves

Here we present the remainder of the dose-response plots from Sec. VA and Sec. VB.

For each of the curves in Fig. 18, an initial dose of 1\(\mu\)M is mapped to an initial concentration of \(1/12\) tumor volume. Equation 5 gives us the relation between the three time points (24, 48, and 72 hours) and the thresholds \((u_T(24), u_T(48), \text{and } u_T(72))\). These thresholds are then used to produce the simulated dose-response curves in Fig. 18. Then the two other initial doses are matched by varying the dose until a minimum distance between two of the time points and the curves is achieved. We choose to match two time points since it is evident that the data set is quite noisy and it has outliers that do not obey the Hill equation [23].

In Fig. 19, the dose from the LINCS data [34] is kept as is. The dose-response curves may then be produced by using (13) for the 72 hour time point data values.

For both figures we need one extra point in order to fit a four parameter sigmoidal curve to the empirical data points. Notice that at an initial dose of zero the response must be around zero, and hence we use that point as well. Using a bootstrap scheme fifty sigmoidal curves fitting the data with a prescribed probability spread at each dose point, are produced (shown in Figs. 11 and 15 in Sec. VA). These curves give us an artificial replication study. From the curves we find the 95\% confidence intervals at each matched dose point, and connect a piecewise linear curve to the lower and upper quantiles of these intervals, which yields a confidence band. Then the dose-response curves from the model are plotted. Since the response from some data pairs do not fit the Hill equation, the parameters for the sigmoidal curve may not always be estimable, and hence these pairs are not included.

Appendix B: Confidence bands

In this appendix, we present the remainder of the confidence band plots from Sec. VA and Sec. VB.

For each of the curves in Fig. 20, an initial dose of 1\(\mu\)M is mapped to an initial concentration of \(1/12\) tumor volume. Equation 5 gives us the relation between the three time points (24, 48, and 72 hours) and the thresholds \((u_T(24), u_T(48), \text{and } u_T(72))\). Then three other initial doses are matched by varying the dose until a minimum distance between the 72 hour time point and the curves is achieved. We choose to study the 72 hour time points since the least amount of noise is observed at this exposure time and therefore more closely matches the Hill equation [23].

In Fig. 21, the dose from the LINCS data [34] is kept as is. The dose-response curves may then be produced by using (13) for the 72 hour time point data values.

For both figures we need one extra point in order to fit a four parameter sigmoidal curve to the empirical data points. Notice that at an initial dose of zero the response must be around zero, and hence we use that point as well. Using a bootstrap scheme fifty sigmoidal curves fitting the data with a prescribed probability spread at each dose point, are produced (shown in Figs. 11 and 15 in Sec. VA). These curves give us an artificial replication study. From the curves we find the 95\% confidence intervals at each matched dose point, and connect a piecewise linear curve to the lower and upper quantiles of these intervals, which yields a confidence band. Then the dose-response curves from the model are plotted. Since the response from some data pairs do not fit the Hill equation, the parameters for the sigmoidal curve may not always be estimable, and hence these pairs are not included.
FIG. 18. Matrix of dose-response curves from the entire data set. The columns denote the drugs and the rows denote the cell lines for the data pairs. Molarities from the LINCS data [34] have been converted to initial volumes. Dose-response curves are produced in accordance with the techniques delineated in Sec. V A.
FIG. 19. Matrix of dose-response curves from the entire data set. The columns denote the drugs and the rows denote the cell lines for the data pairs. Dose-response curves are produced in accordance with the techniques delineated in Sec. V B.
FIG. 20. Matrix of dose-response curves within 95% confidence bands. The columns denote the drugs and the rows denote the cell lines for the data pairs. Molarities from the LINCS data [34] have been converted to initial volumes. Dose-response curves are produced in accordance with the techniques delineated in Sec. V A.
FIG. 21. Matrix of dose-response curves within 95% confidence bands. The columns denote the drugs and the rows denote the cell lines for the data pairs. Dose-response curves are produced in accordance with the techniques delineated in Sec. VB.