Multiscale modelling of drug transport and metabolism in liver spheroids

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Abstract

In early preclinical drug development, potential candidates are tested in the laboratory using isolated cells. These *in-vitro* experiments traditionally involve cells cultured in a two-dimensional monolayer environment. However, cells cultured in three-dimensional spheroid systems have been shown to more closely resemble the functionality and morphology of cells *in-vivo*. While the increasing usage of hepatic spheroid cultures allows for more relevant experimentation in a more realistic biological environment, the underlying physical processes of drug transport, uptake and metabolism contributing to the spatial distribution of drugs in these spheroids remain poorly understood. The development of a multiscale mathematical modelling framework describing the spatiotemporal dynamics of drugs in multicellular environments enables mechanistic insight into the behaviour of these systems. Here, our analysis of cell membrane permeation and porosity throughout the spheroid reveals the impact of these properties on drug penetration, with maximal disparity between zonal metabolism rates occurring for drugs of intermediate lipophilicity. Our research shows how mathematical models can be used to simulate the activity and transport of drugs in hepatic spheroids, and in principle any organoid, with the ultimate aim of better informing experimentalists on how to regulate dosing and culture conditions to more effectively optimise drug delivery.

**Keywords:** Drug transport; Spheroid; Organoid; Hepatocytes; Systems pharmacology; Mathematical modelling.
1 Introduction

The discovery of potential toxicity *in-vitro* remains an important process in providing pre-clinical safety assurances during drug development. However, conventional 2D *in-vitro* experiments, such as monolayer cell culture, tend to be poorly predictive of toxicity, and emerging 3D systems are shown to be more physiologically relevant and predictive of the *in-vivo* environment (1, 2). Accordingly, 3D cell culture systems such as multicellular spheroids are increasingly being utilised in drug development and hepatic safety assessment (3, 4).

Although 3D spheroid systems offer improvements in terms of physiological relevance and *in-vivo*-like functionality, the mechanistic interaction between these systems and drugs is not yet fully understood.

Multiscale *in-silico* methods can improve the application of 3D spheroid models to assess the hepatotoxicity of drug candidates (5, 6). Indeed, mechanistic mathematical modelling of drug metabolism and transport in 3D microtissues is important for the pharmaceutical industry as it facilitates an improved platform for both preclinical drug development and *in-vivo* extrapolation (7). This utilisation of mathematical models, devised to tackle pharmacological research challenges in a systems biology approach, has become known as part of the evolving field(s) of systems pharmacology and/or systems toxicology (8, 9). This approach is a multiscale, multidisciplinary field that employs holistic, integrative methods in order to enhance the understanding and prediction of emergent system properties. Moreover, this methodology is strictly quantitative requiring the integration of quantitative data and modelling to develop mechanistic knowledge of the system and reveal pharmacological and toxicological properties. Consequently, systems pharmacology models are becoming an
increasingly important part of the toolkit to improve capabilities and drive innovation for in-vitro safety assessment (10-12).

In this study we have characterised the spatiotemporal dynamics of drugs in an *in-vitro* hepatic spheroid system by simulating relevant physical processes *in-silico*. A data-driven, multiscale, mathematical modelling framework combining mechanistic information relating to the diffusion, transport and metabolism of chemical species in a hepatocyte spheroid is presented. A microscale single-cell model is analysed to study different transport mechanisms by varying boundary conditions on the cell membrane. This model is then coupled to a multicellular model developed to evaluate the effects of cellular arrangement and density on the transport and penetration of drugs, simulating the *in-vitro* microtissue environment. Such effects include a non-linear relationship between drug lipophilicity and spheroid penetration whereby drug delivery to the spheroid core is minimised for drugs of intermediate lipophilicity. The integration of experimental data allows for the development of realistic geometries and parameterisation of the multiscale model for a range of drugs. Ultimately, by accurately simulating the processes of drug transport and metabolism we aim to enhance the understanding of underlying mechanisms and optimise the use of these systems *in-vitro*.

2 Methods

2.1 Microscale transport – crossing the cell membrane

To simulate the distribution of drugs throughout a 3D tissue comprised of multiple hepatocytes, it is necessary to determine how drugs penetrate and cross the cell membrane. This membrane comprises a phospholipid bilayer, providing a hydrophobic protective barrier
for the cell. Consequently, this chemical barrier property is a key determinant in the effective
permeability of any drug. Many factors affect drug permeability in tissues such as ionisation,
aqueous diffusion between lipid barriers, and protein binding, but the partition into the
membrane (determined by lipid solubility) is one of the most important (13). Highly
lipophilic substances can more readily penetrate the membrane via free diffusion, whilst
relatively hydrophilic substances, (highly soluble in polar solvents such as water or blood),
cannot enter the cell easily and require specific transporters (Figure 1A). The relative role of
transporter proteins in intracellular drug transport is still debated and there remain different
views as to whether passive diffusion or carrier-mediated transport is the major mechanism
(14-18). For the entirety of this study we refer to the two main types of transport: passive
diffusion – entering cells down a concentration gradient directly through the membrane
(passive); and carrier-mediated transport – entering cells via specific transporter proteins
embedded in the plasma membrane (passive or active).

The mathematical representation of microscale drug transport across a cell membrane can be
studied with a simple model considering the processes governing drug concentration
dynamics in two phases, inside and outside the cell, with a permeable barrier in-between.
Once inside the cell, the drug is removed via metabolism. We assume diffusion occurs at
different rates inside \(D_I\) and outside \(D_E\) of the cell, which we initially assume is spherical
of radius \(R\), but relax this assumption in section 2.3. The drug concentration \(C\) dynamics
inside the cell are given by the partial differential equation (PDE):

\[
\frac{\partial C}{\partial t} = D_I \nabla^2 C - \frac{V_{max} C}{C + K_m},
\]

\(1\)
where $V_{\text{max}}$ is the maximum metabolic rate and $K_m$ represents the drug concentration at which metabolism is half maximal. Since there is no flow within the in vitro system, and the dominant form of removal within the multiscale model is assumed to be due to intracellular metabolism, we assume that outside the cell drug transport is governed by diffusion processes only:

$$\frac{\partial C}{\partial t} = D_E \nabla^2 C. \tag{2}$$

For simplicity, we assume that the problem is radially symmetric and rescale the model with respect to cell radius and internal diffusion time (such that the cell boundary is now given by $r = 1$) to give

$$\frac{\partial C}{\partial t} = \frac{1}{r^2} \frac{\partial}{\partial r} \left( r^2 \frac{\partial C}{\partial r} \right) - \frac{V_{\text{max}} C}{C + K_m}, \quad r \leq 1, \tag{3}$$

$$\frac{\partial C}{\partial t} = \frac{D}{r^2} \frac{\partial}{\partial r} \left( r^2 \frac{\partial C}{\partial r} \right), \quad r > 1, \tag{4}$$

where $D = D_E / D_I$ due to rescaling (see supplementary material for details). We impose the following boundary conditions at the cell centre ($r = 0$), for radial symmetry, and a distance away from the cell ($r = r_{\text{max}}$):

$$\frac{\partial C}{\partial r} = 0, \quad r = 0, \tag{5}$$

$$C = C_{r_{\text{max}}}, \quad r = r_{\text{max}}, \tag{6}$$

where $C_{r_{\text{max}}}$ is a constant supply term. Assume that the flux at the cell boundary is equal such that mass is conserved, i.e.,
\[
D_l \frac{\partial C_l}{\partial r} = D_E \frac{\partial C_E}{\partial r}, \quad r = 1,
\]

where \( C_l \) and \( C_E \) are used to distinguish between interior and exterior drug concentrations at the cell membrane boundary. A further boundary condition must be specified at the cell membrane boundary in order to solve the coupled PDE system and investigate the effects of different means of drug transport.

2.1.1 Passive diffusion

The following boundary condition is imposed to describe the flux of drug into the cell due to passive diffusion:

\[
D_l \frac{\partial C_l}{\partial r} = D_E \frac{\partial C_E}{\partial r} = Q(C_E - C_l), \quad r = 1,
\]

where \( Q \) is the permeability coefficient. The mathematical model can be solved numerically in MATLAB R2017b. For methodological details regarding derivations, numerical solutions and simulations of microscale transport, see the supplementary material.

The impact of the permeability coefficient, \( Q \), on the steady-state distribution of drug concentration can be seen in Figure 1B (for temporal dynamics, see supplementary animations). For low permeability coefficients \( Q \ll 1 \), there is less drug penetration per unit time and so there is a low steady-state value inside the cell and a large discontinuity at the cell membrane. As \( Q \) increases, relatively more drug enters the cell per unit time and in the limit, as \( Q \to \infty \), the steady-state solutions converge such that the drug concentration profile is continuous \( (C_E = C_l) \) at the cell membrane boundary (which now provides no effective
barrier or resistance) and the steady-state profile represents the balance of supply via diffusion and removal via intracellular metabolism.

2.1.2 Carrier-mediated transport

For drugs whose physicochemical properties prohibit direct permeation across the cell membrane, specific transporter proteins are required that can mediate the transfer process. The reliance on transporter (or carrier) proteins dictates that the flux is now saturable with an explicit dependence on the surface area concentration, binding affinities, and activity of transporters in the cell membrane. In this scenario, the boundary condition representing membrane transport cannot be sufficiently represented by the passive diffusion condition in equation (8) and so we implement a simple carrier model as applied in other similar physiological membrane transport models, e.g., Keener & Sneyd (19) and Wood & Whitaker (20). This carrier model can be applied to define the flux boundary condition for the carrier-mediated transport model scenario:

$$D_L \frac{\partial C_i}{\partial r} = \frac{T_0 (C_E - \alpha_1 C_i)}{\alpha_2 + \alpha_3 C_E + \alpha_4 C_i + \alpha_5 C_E C_i}, \quad r = 1,$$

(9)

where $T_0$ represents transporter protein concentration on the cell membrane and $\alpha_1, \alpha_2, \alpha_3, \alpha_4, \alpha_5$ represent algebraic expressions dependent on kinetic rates in the carrier model such as binding rates (see supplementary material for more information).

The barrier effect provided by the carrier-mediated transport of drugs across the cell membrane allows for a discontinuity in the steady-state profile of the drug concentration distribution when there is a constant external supply that diffuses towards a metabolically active cell (as before with the passive diffusion case with low permeability). Indeed the
carrier mediated transport condition can be reduced to the passive diffusion condition mathematically with appropriate parametrisation (e.g., $T_0 = Q, \alpha_{1,2} = 1, \alpha_{3,4,5} = 0$). Furthermore, the flexibility of the carrier-mediated condition facilitates the implementation of implicit active processes whereby the flux of drug can move uphill against its concentration gradient (e.g., see Figure 1C). This can be achieved with appropriate parameterisation of the simple carrier model such that $\alpha_1 < 1$, e.g. when binding affinity/dissociation in the interior is lower/higher than exterior binding/dissociation.

Figure 1: Modelling transmembrane transport in a single cell. (A): Drug transport schematics across the cell membrane. Two modes of transport are considered, passive diffusion (pink substrates/circles) and carrier-mediated transport (green substrates/circles). Drugs that permeate the cell via passive diffusion move down a concentration gradient directly through the membrane and are limited by their associated permeability coefficient. This coefficient is dependent on the
physicochemical properties of the drug and drugs which cross the membrane via this mechanism are typically small and lipophilic. Other drugs may require the action of specific membrane-bound transporter proteins to enter the cell (carrier-mediated transport). In this study, it is assumed that this mechanism is dependent on carrier proteins/receptors (depicted in cyan) which can reversibly bind to the substrate and undergo conformational changes to transport the substrate across the membrane. Species within the figure are annotated with symbols related to mathematical models described in the main text and supplementary material. (B): Variation in the permeability coefficient determines the steady-state concentration profile of drug concentration in a single cell for the passive diffusion transport mechanism. Low permeability results in a discontinuity at the cell membrane \( (D = 2, V_{\text{max}} = 10, K_m = 0.5, C_{r_{\text{max}}} = 1) \). (C): For specific parameter choices within the carrier-mediated transport model, a steady-state can be reached such that the drug is transported against its concentration gradient, implicitly simulating an active process \( (D = 2, V_{\text{max}} = 1, K_m = 0.5, C_{r_{\text{max}}} = 1, T_0 = 1, \alpha_1 = 0.5, \alpha_2 = 1, \alpha_3 = 0, \alpha_4 = 0, \alpha_5 = 0) \). Full spatiotemporal dynamics can be found for (B) and (C) in supplementary animations.

### 2.2 Parameterisation

For the full multiscale model, describing the transport and metabolism of drugs in a multicellular *in-vitro* environment, it is useful to include quantitative, dimensional parameter values based on experimental data to directly represent the laboratory scenario for drugs with a range of physicochemical properties. Therefore, it is important to identify relevant parameter ranges for the microscale model before upscaling the problem to the multicellular/tissue level by introducing hepatic spheroid geometry. There are currently three key processes that determine drug dynamics in our system and require parameterisation: diffusion, metabolism and permeation. For simplicity and more general applicability, we will focus on the passive diffusion case and not cover carrier mediated transport during analysis of the multicellular model.

#### 2.2.1 Diffusion of small molecule drugs

Most drugs, and nearly all drugs that cross the cell membrane via passive diffusion, are categorised as small molecule drugs. These are low molecular weight compounds and
comprise most drugs on the market today (21). For a sample database of 321 such drugs (22), we calculated diffusion coefficients based on physical measurements of weight and density (MW ~100-1,200Da; density ~0.6-2.6g/m³). Thus we propose the feasible diffusion coefficient range of approximately $5 \times 10^{-10}$ to $1 \times 10^{-9}$ m$^2$/s (further information in the supplementary material). This narrow range supports the assertion that the main determinants of drug disposition are the ability to translocate across hydrophobic diffusion barriers (permeability) and chemical transformation (metabolism), while variations in the aqueous diffusion rate have only minor effects on overall pharmacokinetics (13). A representative value of $7.5 \times 10^{-10}$ m$^2$/s for both $D_l$ and $D_E$ will be considered as default for further simulations.

2.2.2 Permeability as a function of lipophilicity

The permeability of a drug transported via passive diffusion is related to its lipophilicity, a measurable physicochemical property that can be used to define our permeability coefficient, $Q$. Menochet et al. (23, 24) discovered a log-linear relationship for hepatic uptake between passive diffusion clearance, $P_{diff}$, and lipophilicity,

$$\log P_{diff} = 0.6316 \times \log D_{7.4} - 0.3143, \quad (10)$$

where $P_{diff}$ has units of $\mu$L/min/10$^6$cells and Log$D_{7.4}$ is a partition coefficient measure of lipophilicity at a physiologically relevant pH (pH = 7.4). This relationship allowed us to derive, $Q$, as a function of $P_{diff}$, and the radius of the cell, $R$, by taking into account passive uptake across the whole cell membrane of surface area $4\pi R^2$: 
\[ Q = \frac{P_{\text{diff}}}{4\pi R^2} = \frac{1}{10^6} \frac{10^{(0.6316 \times \text{LogD}_{7.4} - 0.3143)}}{4\pi R^2}. \] (11)

For the full derivation see the supplementary material. LogD\(_{7.4}\) values between 1 and 5 are considered within this study to represent relatively lipophilic, small-molecule drugs (relevant for passive diffusion), with LogD\(_{7.4}\) = 3 as default.

2.2.3 Simplified drug metabolism in hepatocytes

Metabolism represents the principal sink/removal term in our model and the metabolic rate is likely to vary greatly depending on the chemical makeup of the drug of study, as well as the quantity and activity of metabolising enzymes present. Therefore this term is likely to have a significant impact on the overall disposition of drug concentration in a metabolically active in-vitro spheroid system. Metabolic rates are assumed to be independent of space in the model for simplicity, although zonal variation may exist. Brown et al. (25) reported kinetic parameters for a range of compounds to predict metabolic clearance by using cryopreserved human hepatocytes. This publication provided pharmacologically feasible \( V_{\text{max}} \) \((5 \times 10^{-6} \text{ to } 4.5 \times 10^{-1} \text{ mol/m}^3/\text{s})\) and \( K_m \) \((5 \times 10^{-4} \text{ to } 1.4 \times 10^{-1} \text{ mol/m}^3)\) ranges for drugs primarily metabolised in the liver and were thus used as conservative guidance for this model parameterisation, given that cells cultured in 3D often display improved drug metabolism functions. As default, we consider parameters values of \( V_{\text{max}} = 5 \times 10^{-3} \text{ mol/m}^3/\text{s} \) and \( K_m = 1 \times 10^{-2} \text{ mol/m}^3 \).

2.3 Macroscale – hepatocyte spheroid geometry

The impact of the hepatic spheroid environment on drug transport is considered by upscaling our microscale model to consider multiple discrete cells in a realistic spheroid geometry within an extracellular space (culture medium). This hepatocyte spheroid geometry was
generated based upon histological staining of hepatic spheroids to provide representative cell
sizes, number, and arrangement thereby replicating the in-vitro scenario within the multiscale
mathematical model.

2.3.1 Mathematical description of spheroid geometry

Histological staining of a hepatocyte spheroid revealed the spatial distribution of the cell
nuclei within a section (Figure 2A). This spatial information, as well as the spheroid
boundary, was quantified digitally with WebPlotDigitizer (26) and imported into MATLAB.
Due to the abundant expression of extracellular matrix in the hepatic spheroid histological
images, it was not possible to visualise and/or quantify the location of the hepatocyte
membranes. Therefore, we estimated the location of cell boundaries using Voronoi
tessellation (Figure 2B). Briefly, Voronoi tessellation involves assigning regions to each
nucleus such that any point in space within that region is closer to that nucleus than any other.
The boundaries of these regions can be determined by drawing perpendicular bisectors
between adjacent pairs of nuclei. This technique has been shown to provide viable estimates
for the qualitative morphology of cells in a tissue (27).

Cellular ultrastructure was visualised by transmission electron microscopy (TEM). TEM
revealed that the space between hepatocytes was narrow (~0.1-0.5µm, see Figure 2C). These
values are supported by the literature which states intercellular spaces from 100nm to the µm-
scale (28, 29). Furthermore, it should be noted that fixation methods can shrink such
morphological features (30) and therefore we consider both narrow and wide intercellular
space geometries. This was achieved by contracting the vertices of each model cell towards
the cell’s respective centre of mass by 1% (“narrow”, ~0.2µm) or 10% (“wide”, ~2µm) (Figure 2D).

2.3.2 Experimental methods

Primary rat hepatocyte spheroids with an initial seeding density of 5,000 cells were produced using the liquid-overlay technique as described by Kyffin et al. (31). After 11 days in culture the spheroids were washed in PBS, fixed in 4% paraformaldehyde and subjected to routine histological processing before staining with haematoxylin or processed for TEM analysis. For TEM imaging, spheroids were fixed in 3% glutaraldehyde and processed as previously described (31). Ultrathin (~70–90nm) sections were examined using a FEI Tecnai Transmission Electron Microscope at an accelerating voltage of 80Kv and images taken using a Gatan digital camera.

2.3.3 Numerical simulation

The finite-element simulation software, COMSOL Multiphysics® 5.3, was used to solve the multiscale model PDEs. The 2D spheroid slice geometry was imported into COMSOL and the PDEs were defined as before to calculate the dynamics of drug concentration, \( C \), for two separate phases (intracellular, \( C_I \), and extracellular, \( C_E \)):

\[
\frac{\partial C_I}{\partial t} = D_I \nabla^2 C_I - \frac{V_{\text{max}} C_I}{C_I + K_m}, \quad C = C_I, \quad (12)
\]

\[
\frac{\partial C_E}{\partial t} = D_E \nabla^2 C_E, \quad C = C_E, \quad (13)
\]

with boundary conditions at the every cell membrane within the spheroid,

\[
(D_I \nabla C_I) \cdot \mathbf{n} = (D_E \nabla C_E) \cdot \mathbf{n} = Q(C_E - C_I), \quad (14)
\]
for the general inward fluxes, where \( \mathbf{n} \) is the unit normal vector pointing out of each cell. An illustrative example of the multiscale model steady-state with a constant supply of drug at the outer boundary of the media phase \( (C_{r_{\text{max}}} = 500 \mu\text{M}) \) can be seen in Figure 2E, simulated for a drug with physicochemical properties based on the default parameter set described above. Note that permeability \( Q \) is related to LogD\(_{7.4} \) according to equation (11). A 1D cross-section is plotted in Figure 2F for visualisation, highlighting the discontinuities in drug concentration between intra- and inter-cellular space and the heterogeneity in drug concentration between cells in different regions.
Figure 2: **The multiscale model including hepatocyte geometry.** (A): Histological staining of a hepatocyte spheroid slice indicating the location of cell nuclei (blue). (B): Voronoi diagram constructed to provide estimates of hepatocyte boundaries (red) based on location of hepatocyte nuclei (blue). (C): Representative TEM image of a hepatocyte spheroid showing the size of the space between adjacent cells. The intercellular space is indicated by the red arrows. (D): Intercellular space
was introduced into the model geometry by contracting the vertices of Voronoi cells (indicated by black arrows) towards the centre of each cell (yellow stars; nuclei in blue). (E): Steady-state distribution of an example drug ($\text{LogD}_{7.4} = 3$ with default parameters and wide intercellular space), formed with a constant supply of 500µM at the outer media boundary (disc of radius 750µm). The drug distribution is denoted by the colour-bar, demonstrating that there are lower drug concentrations in the central hepatocytes. (F): A 1D cross-section of the simulation (position indicated by white line in (E)) signifies the variation of drug concentration inside and outside of the cells within the spheroid structure, as well as the heterogeneity of intracellular drug concentration in different regions of the spheroid.

3 Results

3.1 Impact of drug permeability on spatiotemporal distribution throughout spheroid

The diffusion rate of a drug depends mainly on size, a property that has minimal variation in small molecule drugs (a detail supported by our analysis of over 300 compounds during parameterisation) and thus has relatively little impact upon drug distribution when compared with the ability to translocate across the hydrophobic diffusion barrier of the cell membrane (13). This translocation ability is determined by the lipophilicity of the drug during passive diffusional transfer across the membrane. Therefore we consider the impact that drug permeability (as determined by lipophilicity) has upon the overall dynamics within the representative in-vitro spheroid system. This analysis is illustrated by simulating the model, dosed for 3 example drugs with different permeability coefficients (corresponding to $\text{LogD}_{7.4} = 1, 3, 5$, within the otherwise default parameter set) via constant supply at the external boundary and comparing the steady-state spatial distribution of drug concentration (Figure 3). Spatiotemporal dynamics can be found in supplementary animations.

The results indicate that for highly lipophilic drugs ($\text{LogD}_{7.4} = 5$), the cell membrane does not represent a significant barrier to drug penetration and there is relatively little difference between drug concentrations in cells and the intercellular space. For relatively lowly
lipophilic drugs ($\text{LogD}_{7.4} = 1$), the membranes represent a significant barrier. Drug concentration is very low within the cells but relatively high in the intercellular space throughout the spheroid. However, in the intermediate case ($\text{LogD}_{7.4} = 3$), there is relatively little drug in the spheroid centre, both inside and outside of the hepatocytes. This is due to the balance between the overall processes of drug transport towards the spheroid centre (diffusion, permeability and metabolism), impacting penetration potential. Overall, it is clear that an increase in permeability results in higher intracellular drug concentration but there is a non-linear response in the intercellular space as permeability is increased, with a potential local minimum for drugs of intermediate lipophilicity. The same observations are made for narrow intercellular spaces and when varying transporter expression in the carrier-mediated transport model (data not shown). This result highlights the potential importance of not only permeability, but intercellular space on overall drug delivery.
Figure 3: **Impact of permeability on drug dynamics distribution.** Top row: Spatial distribution of drug concentration within a hepatic spheroid for 3 different permeability coefficients (permeability decrease from left to right, LogD$_{7.4}$ = 5, 3, 1). The plots represent steady-state values after constant supply of 500µM at the outer media boundary (disc of radius 750µm). Bottom row: Corresponding representative 1D plots are provided for each drug using the same-cross section position as Figure 2E.

3.2 **Impact of intercellular dimensions on spatiotemporal distribution throughout spheroid**

Many mathematical models of cellular spheroids consider geometrical simplifications such as radial symmetry and a homogenous continuum of cells. The consideration of a spheroid with individual hepatocytes modelled as discrete regions in space, and accompanying intercellular space, has a visible impact upon the radial drug concentration profile. This can be seen most clearly in the case of low permeability with large fluctuations in the drug concentration between intra- and inter-cellular space (Figure 3). There is a considerable range of intercellular gap sizes within spheroids, a feature which can be magnified by fixation issues.
and cell type, with tumour spheroids notoriously exhibiting higher porosities (32). Therefore it is prudent to also consider the impact of porosity (gap size) on drug delivery by simulating our model for both narrow and wide intercellular space geometries, as well as a model without intercellular space altogether for comparison. Steady-state spatial distributions in Figure 4 suggest that intercellular space has a considerable impact upon drug penetration, with increased porosity resulting in higher drug concentration for the spheroid interior.

Figure 4: **Impact of intercellular space on drug dynamics distribution.** Spatial distribution of drug concentration within a hepatic spheroid for 3 different intercellular space geometries (no spaces (zero porosity, A); narrow spaces (~0.2µm, B); wide spaces (~2µm, C)). The figures represent steady-state values after constant supply of 500µM at the outer media boundary (disc of radius 750µm) with default parameters.

3.3 **Translating the multiscale model to a simple continuum model**

From Figure 4 it is clear that, when using quantitative, measurable, microscale parameters, the assumption of a homogenous continuum of hepatocytes in the spheroid will be insufficient for simulating spatial drug distributions, particularly for wider intercellular spaces. Therefore, we consider if there are any parameter modifications that can be made such that the continuum model can be said to sufficiently replicate the simulations provided by the more spatially complex discretised model. Such a model would be highly beneficial for the quantification of drug dynamics with greater computational efficiency. For this
investigation we compare the average behaviour of the full discrete, multiscale, dimensional
model ("cell-based model") with the idealised radially symmetric, homogenised sphere
model ("continuum model") in 2D (cylindrical coordinates) given by:

\[
\frac{\partial C_S}{\partial t} = \frac{D_{l}^{\text{Eff}}}{r} \frac{\partial}{\partial r} \left( r \frac{\partial C_S}{\partial r} \right) - \frac{V_{\max} C_S}{C_S + K_m}, \quad r \leq R_s, \quad (15)
\]

\[
\frac{\partial C_0}{\partial t} = \frac{D_E}{r} \frac{\partial}{\partial r} \left( r \frac{\partial C_0}{\partial r} \right), \quad r > R_s, \quad (16)
\]

where \( C_S \) and \( C_0 \) represent spheroid and outer drug concentrations respectively, and \( R_s = 135 \)
\( \mu \text{m} \) (the average radius of the hepatocyte sphere slice in Figure 2), with boundary conditions

\[
D_{l}^{\text{Eff}} \frac{\partial C_S}{\partial r} = 0, \quad r = 0, \quad (17)
\]

\[
D_{l}^{\text{Eff}} \frac{\partial C_S}{\partial r} = D_E \frac{\partial C_0}{\partial r} = Q^{\text{Eff}} (C_0 - C_S), \quad r = R_s, \quad (18)
\]

for effective parameters \( D_{l}^{\text{Eff}} \) and \( Q^{\text{Eff}} \) which represent the parameters to be modified. These
parameters are logical targets for translation since they determine interior transport via
internal diffusion and translocation across cell membranes in the cell-based model.
Homogenisation here can be thought of as an extreme modification of the spheroid structure
such that we reduce the system to a very large single cell with a single permeable membrane.
The effective parameter values of the continuum model were optimised to fit the average
behaviour of the cell-based models for both intercellular space geometries and a
physicochemically relevant range of permeability coefficients (corresponding to LogD_{7.4} = 1,
2, 3, 4, 5). For information regarding parameter optimisation, see the supplementary
material.
The required modifications of effective parameters, both collectively and individually as functions of drug lipophilicity and intercellular space, are summarised in Figure 5, as well as corresponding error metrics. A combined parameter change metric in Figure 5E is introduced to quantify the relative amount of modification required for each scenario (intercellular width and lipophilicity) and defined as

$$\Delta P = \sqrt{\left(\frac{D_{l}^{\text{Eff}} - D_{l}}{D_{l}}\right)^2 + \left(\frac{Q^{\text{Eff}} - Q}{Q}\right)^2}.$$  \hfill (19)

From Figure 5 it is clear that $\Delta P$ is dominated by relative changes in the effective permeability coefficient, $Q^{\text{Eff}}$ (compare Figure 5E with Figure 5A-B). Permeability must be increased to account for the intercellular space in the cell-based models (all lipophilicities), i.e., $Q^{\text{Eff}}/Q \geq 1$ for all LogD$_{7.4}$ (Figure 5B). This effectively makes the spheroid boundary in the continuum model more porous (virtually simulating gaps between cells) and the discontinuity at the spheroid boundary is reduced. It should be noted that in the dimensional cell-based models, while $D_{l}$ remains constant throughout all simulations, $Q$ will change dependent on LogD$_{7.4}$ (recall equations (10)-(11)). This is seen in Figure 5C-D with absolute changes in $Q^{\text{Eff}}$ and $Q$. Permeability must be increased by a greater amount for wider intercellular spaces to be effectively simulated by the continuum model (e.g., Figure 5D) for all LogD$_{7.4}$. This is expected due to the increased porosity provided by wider gaps. Finally, effective permeability must be increased by a greater amount for low lipophilicities. This can be seen in Figure 5B where the effective permeability $Q^{\text{Eff}}$ decreases towards the dimensional value $Q$ with increasing lipophilicity, for both gap sizes, in a monotonic fashion.
This reflects the increased discrepancy between transport through cells and transport between cells found for drugs that are poorly lipid soluble.

In order for the continuum model to effectively simulate intercellular space, intracellular diffusion must be decreased for all lipophilicities, i.e., \( D_{i}^{\text{Eff}} < D_{i} \) for all \( \log D_{7.4} \) (Figure 5A).

The primary effect of decreasing this parameter in the model is to increase the gradient of concentrations within the spheroid. For high lipophilicity and narrow intercellular spaces, the continuum model can provide a representative simulation of the cell-based model by reducing \( D_{i}^{\text{Eff}} \) alone. This property is observed by comparing the negligible changes in \( Q^{\text{Eff}} \) relative to \( D_{i}^{\text{Eff}} \) at high lipophilicity and narrow intercellular spaces. For example, when \( \log D_{7.4} = 4 \) and 5, \( D_{i} \) is decreased by 94% and 86%, while \( Q \) is unchanged (Figure 5A-B). Theoretically, given a high enough value of \( \log D_{7.4} \), this behaviour is expected for wide spaces too, but this is beyond relevant parameter space.

Regardless of lipophilicity, the optimised continuum model compares better with the cell-based model of narrow intercellular gaps (Figure 5F, solid lines). This is likely due to the relatively lower amount of fluctuations in the mean 1D profiles as there is less extracellular space in general within the spheroid. These fluctuations represent the local drug concentration variation at the cellular scale due to discrepancies between intra- and extracellular phases, which can be very high for drugs that are poorly lipid soluble (e.g., see Figure 3 1D profiles). Prior to any optimisation and re-scaling of dimensional parameters to their effective counterparts \( (D_{i} \rightarrow D_{i}^{\text{Eff}}, Q \rightarrow Q^{\text{Eff}}) \), there was a clear pattern in the fit quality between the simple continuum model approximation and the cell-based models of different sized intercellular spaces (Figure 5F, dashed lines). Generally, the continuum model fits the
narrow spaces better for low membrane permeability and wider spaces better for high permeability. This feature appears to be correlated to the overall higher intracellular drug concentrations found in spheroids with wider spaces (since there is proportionally less transport across metabolising cells). The (pre-optimised) continuum model exhibits very low drug concentration within the spheroid at low permeability and so fits the narrow spaced model better. At higher permeabilities the continuum model has relatively high interior concentration and so fits the wide spaced cell-based model better (Figure S1 in the supplementary material). This switch in behaviour is likely due to the continuum model only providing a single barrier to permeation (spheroid boundary), which, once penetrated, facilitates drug penetration via diffusion solely.

Interestingly, Figure 5E indicates that the cell-based model with wider intercellular spaces requires more parameter modification for all drug lipophilicities. Despite the intra-spheroidal gradients being vastly different between the (pre-optimised) continuum and narrow cell-based model at high permeabilities (Figure S1), the boundary intracellular drug concentrations are similar. Therefore the continuum model can be optimised via sufficient reduction in $D_{l}^{Eff}$ whilst maintaining the original permeability coefficient ($Q^{Eff} = Q$). However, in order to simulate the wide cell-based model, and account for different concentrations in boundary cells, a relatively greater change in $Q^{Eff}$ was required (compare relative changes in effective parameters at LogD$_{7.4}$ = 4 and 5 for $D_{l}^{Eff}$ and $Q^{Eff}$ for both models, Figure 5A-B).
Figure 5: **Emulating the cell-based models with a simple, symmetric continuum model.** Effective parameters for intracellular diffusion and permeability ($D_{i}^{Eff}$ and $Q^{Eff}$) were optimised in the continuum model to fit the output for the cell-based models for a range of drug lipophilicities. (A): The optimised $D_{i}^{Eff}$ value provides the required relative changes in intracellular diffusion for the continuum model to match the cell-based models with narrow (blue) and wide (red) intercellular space geometries. (B) Required relative changes in permeability identified by optimising $Q^{Eff}$. Absolute values of $Q^{Eff}$ optimised for each cell-based model and drug lipophilicity (LogD$_{7,4}$) are...
plotted in (C). (D): Optimised $Q^{\text{Eff}}$ on a log scale (y-axis). (E): A combined parameter change metric ($\Delta P$) indicates the summarised amount of model modification required for the continuum model to effectively simulate the cell-based models. (F): Comparisons between model outputs are calculated by using the R$^2$ error metric to determine relative quality of fits (see supplementary material for definition). Comparisons are made both prior to optimisation (“pre”, direct comparison using dimensional parameters, i.e., $D_I^{\text{Eff}} = D_I, Q^{\text{Eff}} = Q$) and post-optimisation (“post”).

3.4 Investigating the impact of permeability on the dynamic process of drug delivery in different regions of the spheroid for a bolus dose

Intercellular space has a discernible impact on the spatiotemporal drug dynamics in the in-vitro spheroid environment and moreover, a non-linear effect was revealed for local concentrations within intercellular space as permeability is increased (Figure 3). Since this phenomenon (i.e., a monotonic decrease in intracellular drug concentration with decreasing permeability, but a non-monotonic response in the intercellular regions) cannot be described by the simple continuum model, it is worth considering the potential impact of this feature on drug penetration. Here we choose to examine drug delivery and subsequent effects by calculating the total uptake/metabolism of the drug in different regions of the spheroid. To investigate drug delivery via metabolism, we introduce the following “metabolism” variable, $M$, with dynamics:

$$\frac{\partial M}{\partial t} = \frac{V_{\text{max}} C}{C + K_m}, \quad C = C_I,$$  

which corresponds to accumulated drug metabolised and is only relevant inside model cells. Corresponding model simulations are conducted with a finite bolus dose initially supplied in the outer medium, uniformly distributed in the extracellular space outside the spheroid, and zero-flux boundary conditions are imposed on the outer boundary of the media phase.
Two separate regions are defined, “outer” and “inner”, corresponding to cells of comparable size in the outer boundary layer of the spheroid, \((x, y) = (-10\mu m, 110\mu m)\), and the spheroid centre, \((x, y) = (0\mu m, 0\mu m)\). Simulations are run to the drug-free steady-state whereby all of the initial dose has been removed from the system and accumulated in the effective sink variable, \(M\). For highly lipophilic drugs, the concentration dynamics are relatively similar between inner and outer regions as the drug is able to be transported throughout the spheroid quickly, unrestricted by permeability. However, the outer cells are exposed to slightly higher concentrations and consequently more drug is metabolised in this region, demonstrated by similar rates of metabolism (Figure 6A). Simulations of lowly lipophilic drugs require much longer timespans in order to reach equilibrium due to the reduced uptake rate at the cell membranes. However, due to the intercellular transport via diffusion, even centrally located cells receive relatively high local drug exposure and metabolise at a similar rate to outer cells (Figure 6C). It is the in-silico drugs of intermediate lipophilicity in this model scenario that exhibit the most striking discrepancies between inner and outer cells (Figure 6B). The impact of these varying rates of metabolism between drug lipophilicities and regions of the spheroid can be evaluated by comparing the total drug metabolised (Figure 6D). The greatest discrepancy in drug uptake between outer and inner hepatocytes is revealed for drugs of intermediate permeability (1,250% increase from inner to outer cells for \(\text{LogD}_{7.4} = 4\) compared to just +13% for \(\text{LogD}_{7.4} = 1\) and +219% for \(\text{LogD}_{7.4} = 6\)). Furthermore, outer cells in this case receive the most drug out of all three cases studies and the inner cells receive the least (Figure 6D). This effect can potentially be exacerbated when carrier-mediated transport kinetics are modelled at the cell membrane, due to the saturating effects of this uptake mechanism (arbitrary transporter parameterisation, data not shown). This feature has
the potential to significantly impact experimental design considerations and *in-vitro* drug efficacy and toxicity evaluation.

![Figure 6: Impact of drug lipophilicity on uptake and metabolism in different regions of the spheroid.](image)

Figure 6: **Impact of drug lipophilicity on uptake and metabolism in different regions of the spheroid.** Metabolism rates are plotted against time as a result of model simulations following a bolus dose of 100µM, initially uniformly distributed in the medium for varying drug lipophilicities (LogD₇.₄ = 6 (A), 4 (B), 2 (C)). (D): Total metabolised drug (after complete clearance) is compared for inner and outer hepatocytes within the spheroid for a range of drug lipophilicities (LogD₇.₄ = 1-6).

### 4 Discussion

The enhanced sophistication of current cell culture methodologies due to increasing advancements in scientific understanding and technological developments has allowed for *in-vitro* studies to become more physiologically relevant. There are a range of different *in-vitro* models that span varying levels of complexity, reproducibility, high-throughput potential, and
cost. Spheroids represent an intermediate experimental model that allows for increased physiological relevance over 2D monolayers due to the 3D environment, as well as more appropriate cell morphology and functionality while remaining cost-effective, consistent, and easy to use (1). The subsequent prevalence of liver spheroid cultures for studying hepatocyte behaviour *in-vitro* is evident and represents a key component of drug development such that drug candidates can be tested for efficacy and toxic potential in a 3D environment with physiological gradients (31, 33-35). However, such sophisticated adaptations offered by technological advancements are often prone to rapid widespread adoption, misuse and consequent distrust due to a lack of deep mechanistic understanding of the new tool. This is related to what is sometimes known as the hype cycle for emerging technologies (36). Data-driven multiscale mathematical models provide an ideal platform from which to try and enhance mechanistic understanding of new biotechnologies by simulating the underlying physical processes. Additionally, the development of spatiotemporal data generated by 3D cell imaging offers tremendous opportunities for developing, parameterising and testing multiscale mathematical models and in response, mathematical modelling can be successfully used to optimise these developing technologies.

In this study we developed a mathematical model of drug transport and metabolism in a multiscale spheroid framework, accounting for microscale processes such as membrane transport kinetics and how they relate to the physicochemical properties of a drug, and macroscale features such as the geometry of a hepatocyte spheroid, informed by imaging data. Cellular uptake of drugs was modelled by the two major processes of transport across the cell membrane, passive diffusion and carrier-mediated transport (17). The carrier-mediated transport microscale model was innately more complex, depending on quantities
such as transporter protein expression, binding kinetics and rates of conformational change and this complexity allowed for a wider array of dynamic mechanisms such as enzymatic saturation and active processes. The extensive parameterisation required to quantify the carrier-mediated transport model depends on more compound-specific information, and so the passive diffusion case became the main focus of investigations within the generalised multiscale framework, more relevant for relatively lipophilic compounds.

The explicit representation of individual hepatocytes based on imaging data allowed for an investigation into the effects of including a distinct cell-based geometry in the model. The model suggests that steady-state intracellular drug concentrations increase monotonically with increasing drug lipophilicity. However, a non-monotonic relationship was revealed between drug lipophilicity and intercellular drug concentration (Figure 3), while the width of the intercellular space further impacted spatial drug distribution (Figure 4). Intercellular space geometry, or spheroid porosity, is therefore a key physiological feature of the multicellular structure but is both difficult to accurately quantify and known to vary widely between cell types. This is particularly relevant in the case of tumour-derived spheroids, whose morphology tends to be more porous (32), and organoids that are increasingly being used in efficacy testing for tumour cells (37). We therefore studied two different average intercellular widths informed by TEM data and the literature which suggested a range of $10^{2}$-$10^{3}$nm-scale, with results varying due to cell type, tumour phenotype and experimental artefacts such as fixation (29, 30, 38).

While it is important to account for intercellular space within spheroids to correctly model drug delivery, the consequent increase in complexity by modelling this feature explicitly renders detailed analytic work intractable and deriving numerical solutions is costly with
respect to time and computational power requirements. Therefore it is appropriate to consider
the application of simplified models that consider averaged or homogenised system
behaviour and under what conditions they can provide valid approximations (39). We have
shown how to approximate the cell-based models using a simple, symmetric, continuum
model by reparameterising dimensional parameters to re-scaled effective counterparts. For
relatively narrow intercellular gaps, these approximations are more accurate and the required
parameter changes are reduced. The differences between the models, due to the explicit
representation of intercellular space (porosity) within the cell-based model, are largely
accounted for by increasing the effective permeability parameter. This parameter determines
the translocation of drug at the boundary between intra- and extracellular phases and, as a
result, the degree of discontinuity in drug concentration across the cell membrane. This is
particularly important at lower lipophilicities when permeability limitations are maximised.
For higher lipophilicities and narrow intercellular space, the spatially averaged dynamics of
the cell-based model can be effectively simulated with the symmetric continuum model by
appropriate reductions in the effective internal diffusion parameter only. Further work is
required to determine the impact of spatially varying quantities that might exist within a
spheroid such as intercellular space or transporters that vary zonally (40), and how these
might compare between continuum models and cell-based models. Metabolic rates are also
known to vary in space throughout multicellular structures due to gradients in environmental
factors such as oxygen and glucose (41). Alternative model simplifications that might
expedite analysis can be made by careful consideration of potentially redundant model
complexities such as intracellular diffusion, which may be neglected in some scenarios. The
model currently neglects any intracellular binding of the drug for simplicity, focusing on the
dominant mechanisms of transport and removal (metabolism) that drive the spatiotemporal
dynamics. However, for specific future applications of the model, intracellular binding could
be considered by ascertaining the relevant fraction unbound for a particular drug, as this will
lower the rate of metabolism for those drugs which bind strongly to intracellular proteins and
nuclear structures.

The discovery of an apparent local minimum in drug penetration, whereby intercellular
concentrations are lower for intermediate membrane permeation, motivated an investigation
into corresponding effects on drug delivery, uptake and metabolism in spheroid centres for a
bolus dose (Figure 6). The results of this investigation indicated that indeed, it is feasible to
observe minimal drug uptake at the spheroid centre for drugs of intermediate lipophilic
properties (with the majority of drug being metabolised at the outer regions). These
mechanistic insights and modelling results have potential impact for the dosing of spheroid
systems in-vitro as well as relevance for analogous in-vivo systems such as avascular
tumours. It is not necessarily sufficient to assume that increasing a chemical’s lipid solubility
will enhance its metabolism at the spheroid centre. Lowly lipid soluble drugs may require a
much longer time in culture but ultimately metabolise the drug more uniformly throughout
the spheroid. Accounting for reduced penetration due to the intermediate lipophilic property
may be alleviated somewhat by increasing the dose, but this could have potentially toxic
consequences from overdosing cells at the spheroid boundary. Other experimental design
options include manipulating permeability (by chemical modification or intervention, but this
could potentially further increase the divergent amounts of drug being metabolised in
different regions of the spheroid) or using smaller spheroids. These investigations could be
conducted within the in-silico framework, in the first instance, to guide strategy. The
implications of drug delivery characteristics based on permeability parameters could potentially be translated to targeting delivery in tissues of multiple cell types expressed zonally. For example, targeting the central zone of a spheroid that contains cells of a different phenotype (e.g. cancerous/hypoxic) may be aided by manipulating these properties regarding permeability, i.e., making certain that the permeability is either relatively high or relatively low to ensure delivery to the spheroid centre. The combination of mathematical modelling with experimental imaging provides a convenient in-silico testing toolkit to optimise the use of 3D cell culture systems in the laboratory and maximise the potential of spheroid models aiding drug discovery, toxicity testing and dose optimisation.

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Author Contributions

JAL wrote the manuscript; JAK, ALH, HEC, CM, & PS performed the experiments; JAL, SDW & RNB contributed to the mathematical modelling; JAL, DPW, SDW & RNB designed the research. All authors read and approved the final manuscript.

Declaration of Interests

The authors declare no competing interests.
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