A validation of the classical one-compartment model for perfusion

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Index Terms

Porous media flow, indicator, brain DCE-MRI

Abstract

I. MATHEMATICAL THEORY

A. A synthetic one-compartment single-phase flow model

A synthetic one-compartment (1C) model of the plasma flow within a porous organ can be developed from the principle of mass balance and Darcys law. Denote the porosity $\phi = \phi(x,t)$ with units m^3/m^3 as the relative quantity describing the fluid volume within a control volume Ω_i divided by the total volume of Ω_i . Within the classical theory of compartment modelling for human organs the porosity is better known as the cerebral blood volume (CBV). The fluid density has units (kg/m^3) and is denoted by $\rho = \rho(x,t)$. We apply the geomechanical understanding of flux as a surface flux q = q(x,t) with units $(m^3/s/m^2)$. The surface flux is a vector field describing the volume of fluid per unit time flowing across a sliced unit area of the sample. The continuity equation reflecting conservation of mass within the fluid states

$$\frac{\partial(\phi\rho)}{\partial t} + \nabla \cdot (\rho q) = \tilde{Q} \tag{1}$$

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where $\tilde{Q} = \tilde{Q}(x,t)$ is a source or sink of fluid mass with units $kg/s/m^3$. For the current task we want to model the perfusion of blood in the brain and we consider it a stationary process. This assumption is only partly true due to factors like external stress which can influence the flow pattern. However, it is still a reasonable and common approximation. By these simplifications the porosity, fluid density and flux are only functions of space, $\phi = \phi(x), \rho = \rho(x), q = q(x)$. Furthermore we assume that the fluid density ρ is constant in space, which is satisfied within all practical means for water within the human body. The continuity equation (1) then becomes

$$\nabla \cdot q = \frac{\tilde{Q}}{\rho}.\tag{2}$$

In order to scale away the density ρ we define another source term Q with units $m^3/s/m^3$ having the relation $\tilde{Q}=Q\rho$, thus transforming (2) into

$$\nabla \cdot q = Q. \tag{3}$$

The right hand side is only non-zero within the source or the sink. Elsewhere, (3) is concurrent with the incompressibility condition of divergence free flow.

Low velocity fluid flow in porous media is described by Darcys law

$$q = -\frac{K}{\mu} \Big(\nabla p + \rho g \nabla z \Big) \tag{4}$$

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where g is the gravitational acceleration, K is the permeability tensor in units m^2 , z is the spatial position along the gravitational field and $\mu = \mu(x)$ is the viscosity of the fluid with units $Pa \cdot s$. For the current project the flow is taking place perpendicular to the gravitational field and the gravitational term can thereby be discarded, such that a simplified version of Darcys law becomes

$$q = -\frac{K}{\mu} \nabla p. \tag{5}$$

Moreover, assuming a hermitian and positive definite permeability tensor K, and combining (3) and (5) results in the elliptical operator

$$\nabla \cdot \left(-\frac{K}{\mu} \nabla p \right) = Q \tag{6}$$

for the scalar pressure field p. Denote the boundary of Ω as $\partial\Omega$, with an outward normal vector n(x). We impose Neuman boundary conditions of no pressure gradient across the boundary, thus $\nabla p \cdot n = 0$ on $\partial\Omega$. This conditions is equivalent to no flow across the boundary. Equation (6) in combination with Neuman boundary conditions $\nabla p \cdot n = 0$ on $\partial\Omega$ is the flow model we apply to create the synthetic brain template. The flux field can be computed according to (4) from the obtained pressure map. An example of pressure and flux fields is shown in Fig. 5 (a) - (c).

B. The forward problem for indicator dilution

In this section we add a tracer to the flow entering via the source, where the aim is to describe how the tracer is carried along within the flow. The concentration map resulting from the simulation is later used as a model for observed concentrations as they could appear from a MR scan. We will apply this concentration for the inverse problem of determining the perfusion and porosity. By comparing the inversely reconstructed perfusion with the known perfusion we can validate the classical models for CBV and CBF.

Denote the space and time depending tracer concentration with respect to the fluid space as c(x,t), with units Mol/m^3 . Also, denote the tracer concentration with respect to a control volume as C(x,t), also with units Mol/m^3 . Thus, for c(x,t) the normalizing volume refers to the fluid volume and for C(x,t) the normalization refers to the control volume. The units Mol/m^3 are applied instead of the classical units mMol/L in order to be compatible with Q(x) having units $m^3/s/m^3$.

The porosity is from definition $\phi_i = v_i/V_i$ for a plasma volume v_i within a control region Ω_i having a total volume of V_i . Letting $|\Omega_i| \to 0$ leads to the continuos porosity field ϕ . Denote the number of tracer molecules within a control roi as n_i . From the definition of c and C, we obtain the relation $n_i = cv_i = CV_i$. Combining this with $\phi_i = v_i/V_i$ leads to $C_i = c_i\phi_i$, approaching the continuous formulation $C = c\phi$ as $|\Omega_i| \to 0$.

The rate of change of tracer molecules within a geometric control volume Ω_i can be phrased as

$$\frac{\partial}{\partial t} \int_{\Omega_i} C dx = \int_{\Omega_i} \frac{\partial}{\partial t} (\phi c) dx. \tag{7}$$

By discarding any dissipative diffusion process, the change in tracer mass within Ω_i occurs only from advective flow and source/sink terms as

$$-\int_{\Gamma_i} c(q \cdot \nu) ds + \int_{\Omega_i} c_{so} Q_{so} dx + \int_{\Omega_i} c Q_{si} dx \tag{8}$$

where $Q_{so}(x) \geq 0$ $[m^3/s/m^3]$ is the source term and $Q_{si}(x) \leq 0$ $[m^3/s/m^3]$ is the sink term which both are zero everywhere except from in the respective source/sink locations. Note that $\int_{\Omega} Q_{so} dx = -\int_{\Omega} Q_{si} dx$, in line with incompressible flow. The point wise relation between Q, Q_{so} and Q_{si} is $Q = Q_{so} + Q_{si}$. From preservation of the tracer mass, equations (7) and (8) must balance such that

$$\int_{\Omega_i} \frac{\partial}{\partial t} (\phi c) dx + \int_{\Gamma_i} c(q \cdot \nu) ds = \int_{\Omega_i} c_{so} Q_{so} dx + \int_{\Omega_i} c Q_{si} dx. \tag{9}$$

Assuming a stationary porosity ϕ , this can be rewritten as

$$\int_{\Omega_i} \phi \frac{\partial c}{\partial t} dx + \int_{\Gamma_i} c(q \cdot \nu) ds = \int_{\Omega_i} c_{so} Q_{so} dx + \int_{\Omega_i} c Q_{si} dx. \tag{10}$$

Upon application of the divergence theorem, Eq. (10) is consistent with the continuity equation on local form

$$\phi \frac{\partial c}{\partial t} + \nabla \cdot (cq) = c_{so}Q_{so} + cQ_{si}, \tag{11}$$

a quasi-linear transport equation in c(x,t).

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This is equivalent to

$$\phi \frac{\partial c}{\partial t} + c \nabla \cdot q + q \cdot \nabla c = c_{so} Q_{so} + c Q_{si}, \tag{12}$$

This is equivalent to

$$\phi \frac{\partial c}{\partial t} + cQ + q \cdot \nabla c = c_{so}Q_{so} + cQ_{si}$$
(13)

This is equivalent to

$$\phi \frac{\partial c}{\partial t} + cQ_{so} + cQ_{si} + q \cdot \nabla c = c_{so}Q_{so} + cQ_{si}$$
(14)

This is equivalent to

$$\phi \frac{\partial c}{\partial t} + cQ_{so} + q \cdot \nabla c = c_{so}Q_{so} \tag{15}$$

This is equivalent to

$$\phi \frac{\partial c}{\partial t} + q \cdot \nabla c = c_{so}Q_{so} - cQ_{so} \tag{16}$$

This is equivalent to

$$\phi \frac{\partial c}{\partial t} + q \cdot \nabla c = (c_{so} - c)Q_{so} \tag{17}$$

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The input tracer concentration c_{so} is assumed known at all time points as well as the source and sink terms Q_{so} and Q_{si} . Inserting (3), applying $Q = Q_{si} + Q_{so}$ and dividing by the non-zero porostiy ϕ leads to

$$\phi \frac{\partial c}{\partial t} + q \cdot \nabla c = (c_{so} - c)Q_{so}. \tag{18}$$

We refer to (10) and (18) as the *simulated flow model (SF model)*. Equation (10) is implemented under the assumption of (3), which the obtained flux q is fulfilling. Equation (10) is used for numerical simulations and (18) is only used for analytical considerations.

C. Conversion of flux into perfusion

The SF model uniquely determines the flux field q(x). However, it is not obvious how to transform a flux field into a scalar perfusion field P(x), a widely used parameter in the classical field of compartment modelling. There are at least two obvious differences between q and P. First, the flux is a vector field and the perfusion is a scalar field. Second, the flux relates to a surface area and the perfusion relates to a volume. Thus, these to quantities are strictly, mathematically different but still conceptually related. In the following we describe a method for converting flux into perfusion.

The classical understanding of perfusion is the amount of blood feeding a tissue volume per unit time. Thus, the perfusion P has the units $m^3/s/m^3$. It is common to scale this quantity to normalized perfusion $P_n(x)$ with units ml/min/100ml. One approach for converting flux into perfusion could be to estimate the perfusion as the total inflow (or outflow) of fluid (e.g. arterial blood) into a control region per unit time, and then normalizing with the control region volume. This is only a valid approach if every control region is separated from other control regions, and not feeding each other. Thus, this approach is valid for an entire organ being the control region. Such understanding is in line with the understanding of classical compartment models for perfusion where each voxel has its own source of feeding arterial blood, independent of the neighbour voxels. Clearly, this is a simplification since voxels will be fed by their neighbours. In our synthetic flow model SFM as well as in normal tissue this assumption is violated since the voxels are feeding their neighbour voxels with arterial blood. Simply

summing the total inflow into a voxel and dividing by the voxel volume will strongly over-estimate the perfusion since we would divide by the wrong normalization volume. The problematic issue is that the incoming blood is feeding more voxels than the current voxel. This phenomenon is demonstrated in Fig. 1 where the volume on the left has the true perfusion value of $P_1 = F_0/(2V)$ for an incoming flow F_0 [m^3/s] and distribution volume 2V [m^3]. However, for another discretization as shown in the middle, the perfusion within each of these sub-volumes becomes $P_2 = F_0/V = 2P_1$. Taking the average across both sub-volumes, it is clear that the perfusion is over-estimated with a factor of two. A discretization dependent perfusion value is not desirable, and the perfusion estimate of P_2 is clearly wrong.

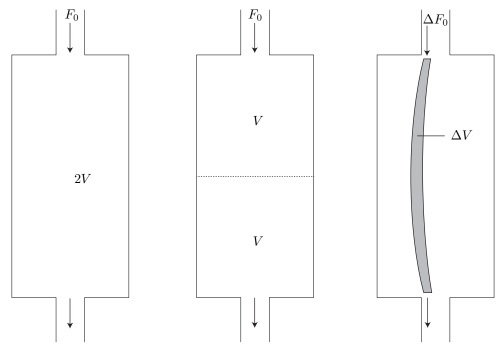


Fig. 1: Perfusion within a small volume. Left: A compartment with volume 2V is exposed to a flow F_0 $[m^3/s]$ of fluid. From definition, the overall perfusion within this object becomes $P_1 = F_0/(2V)$ $m^3/s/m^2$. Right: The volume is divided into two smaller compartment (e.g. voxels), and the perfusion for each of the compartments becomes $P_2 = F_0/V = 2P_1$. This discrepancy between the two discretisation regimes occurs because the flow is counted twice as it is feeded from one voxel to the other. Right: As a solution to the described problem we pick out a true distribution volume ΔV (area in this 2D sketch), which is a small area around a given streamline along the centre line of the grey area. This is the true distribution volume (area) which is feeded with arterial blood from the incoming fractional flow ΔF_0 . The correct perfusion within ΔV is therefore $\Delta F_0/dV$. The entire compartment can further be divided into infinitesimal distribution volumes, thus providing voxelwise perfusion values.

The reason for this discrepancy is that for P_2 the perfusion has been counted twice since we are dividing by the wrong distribution volume. Instead, we need to consider to the classical definition of perfusion. The concept of perfusion has a very precise meaning, as the amount of arterial blood per time unit delivered to a capillary bed in a biological tissue, and then scaled by the feeded tissue volume. Therefore, we must divide the incoming flow by the total distribution volume that is covered by the fluid streamlines. This formulation coincides with the classical understanding of perfusion, and the correct distribution volume will rather be the volume that the fluid particles within an infinitesimal cross-sectional area around the streamlines are covering. Assuming laminar flow, the streamlines are not crossing each other and we can with this formulation correctly estimate the true distribution volume that is fed by a given arterial blood flow.

Formally, the direction of the streamline vector field is identical to the flux vector field mediating the flow. Pick an area S around a streamline, where S is formed as a disc with area $A = \pi r^2$ and infinitesimal area $dA = 2\pi r dr$. The flow in m^3/s passing the cross-sectional area S becomes

$$F_S = \int_S (q \cdot n) dA = \int_0^R (q \cdot n) 2\pi r dr \tag{19}$$

for a unit vector n pointing along with the flow, and where R is the radius of S. Since the flux is carrying the flow, n=q/|q| and the flow F_S across S becomes

$$F_S = 2\pi \int_0^R |q| r dr. \tag{20}$$

The total distribution volume that the flow is passing through can be estimated as

$$V_S = \int_0^L \int_S dA dl \tag{21}$$

which is equivalent to

$$V_S = 2\pi \int_0^L \int_0^R r dr dl, \tag{22}$$

where L is the upper line integration limit of a streamline from inlet to outlet. We can now formalize a precise perfusion measure in agreement with the physiological concept of perfusion as

$$P = \lim_{R \to 0} \frac{F_S}{V_S} = \lim_{R \to 0} \frac{\int_0^R |q| r dr}{\int_0^L \int_0^R r dr dl}.$$
 (23)

Using L'Hopitals rule we reach a locally well defined perfusion measure

$$P(x) = \frac{|q(x)|}{|\mathbb{C}|(x)|} \tag{24}$$

where $||\mathbb{C}(x)| = \int_0^L dl$ is the length of the streamline passing through the point x. This is an explicit formula for converting flux into perfusion and is later used for evaluation of the classical model for perfusion. With the model in (24) we realize that every point along a streamline will have the same value of perfusion. This is reasonable since these voxels are fed by the same flow.

D. Estimating the porosity

The principle of mass balance of fluid and tracer particles is described by Eq. (3) and (11). For locations where the source and sink is zero, $Q_{so}(x) = 0$ and $Q_{si}(x) = 0$, and (11) becomes

$$\phi \frac{\partial c}{\partial t} = -q \cdot \nabla c. \tag{25}$$

Integrating from t_0 to t_1 results in the model

$$\phi[c(x,t_1) - c(x,t_0)] = -\int_{t_0}^{t_1} q \cdot \nabla c dt.$$
 (26)

Inserting $t_0=-\infty$ to $t_1=\infty$ and boundary conditions $c(x,-\infty)=c(x,\infty)=0$ leads to

$$q \cdot \nabla \int_{-\infty}^{\infty} c(x, t)dt = 0 \tag{27}$$

since q is independent of time. Equation (27) states that $\int_{-\infty}^{\infty} c(x,t)dt$ is constant along the streamlines of the fluid flow. Thus, we can choose any voxel in the downstream from the arterial input such that

$$\int_{-\infty}^{\infty} c(a,t)dt = \int_{-\infty}^{\infty} c(x,t)dt$$
 (28)

where c(a,t) is the tracer concentration of the arterial input, and assuming the region of interest (ROI) for the arterial input is a infinitesimal small point in space for the spatial position x = a. Using $C = \phi c$ and assuming the porosity of the arterial inlet is known (and typically close to one), here denoted by ϕ_a , we have an analytical expression for the porosity at any position x,

$$\phi(x) = \phi_a \frac{\int_{-\infty}^{\infty} C(x, t)dt}{\int_{-\infty}^{\infty} C(a, t)dt}.$$
(29)

Note the boundary conditions that were used to arrive at this expression. Practially, $c(x, t \to \infty)$ will never go to zero and one possible solution to this problem is to fit a gamma curve to the data, approaching zero as $t \to \infty$. The expression reached in (29) coincides with the classical formula for CBV and is hereby analytically proven for a one-compartment model.

E. Classical model for perfusion

Denote as N the number of tracer molecules within a given compartment. Conservation of tracer molecules within the compartment leads to the differential equation

 $\frac{dN}{dt} = N_a - N_{out} \tag{30}$

where N_a and N_{out} are the number of tracer molecules entering or leaving the control region per unit time. Equation (30) can be written as

$$\frac{dN}{dt} = \tilde{P}_a c_a(t) - \tilde{P}_{out} c_{out}(t) \tag{31}$$

where \tilde{P}_a , \tilde{P}_{out} are the non-normalized perfusion $[mm^3/s]$ entering or leaving the system, respectively, and where $c_a(t)$ is the tracer concentration at the inlet. Due to the incompressibility condition, $\tilde{P}_a = \tilde{P}_{out} = \tilde{P}$, and assuming a well-mixed fluid compartment leads to

$$\frac{dN}{dt} = \tilde{P}(c_a(t) - c(t)) \tag{32}$$

where c(t) is the tracer concentration within the compartment. Dividing by the control volume V we get

$$\frac{dC}{dt} = P(c_a(t) - c(t)). \tag{33}$$

for C=N/V and $P=\tilde{P}/V$, noting that P is the fluid volume flow per unit time per unit tissue volume $[mm^3/s/mm^3]$, known as the perfusion or CBF when applied within the cortex of the brain. The subscript refers to the classical formulation of perfusion. This ODE can be rephrased as

$$\frac{dc}{dt} - \frac{P}{\phi}c(t) = \frac{P}{\phi}c_a(t). \tag{34}$$

For a stationary perfusion P this equation has the general solution

$$c(t) = \frac{P}{\phi} \int_{t_0}^{t} e^{-\frac{P}{\phi}(t-s)} c_a(s) ds + De^{-\frac{P}{\phi}(t)}$$
(35)

Assuming no tracer at $t = t_0$ and setting $t_0 = 0$ leads to the requirement D = 0, and the solution becomes

$$c(t) = \frac{P}{\phi} \int_0^t e^{-\frac{t-s}{T}} c_a(s) ds \tag{36}$$

for the mean transit time $T \equiv \phi/P$. Incorporating $C = c\phi$ leads to

$$C(t) = P \int_0^t e^{-\frac{t-s}{T}} c_a(s) ds.$$
 (37)

Deconvolution of (37) is the classical model for estimating the perfusion P with a known residue function $R(t) = e^{-t/T}$. Denote the impulse response function I(t) = PR(t). Equation (37) can then equivalently be written as a convolution

$$C(t) = I(t) * c_a(t) \tag{38}$$

provided $c_a(t)$ is zero for t < 0.

Applying the forward model in (38) for a known perfusion P leads to a simulated flow model based on convolution, in accordance with the classical theory. We refer to this model as the *simulated convolution model (SC model)*. We have now defined two competing models for indicator dilution, the SF model and the SC model. This was done in order to be able to discover any discrepancies from the true perfusion field and relate them to shortcomings in the model itself or to instability of the numerical deconvolution.

F. Relating the simulated flow (SF) model with the classical model for perfusion

Consider the solute transport equation (11), and rewrite it as

$$\frac{\partial c}{\partial t} + \frac{q}{\phi} \cdot \nabla c = \frac{(c_{so} - c)Q_{so}}{\phi}.$$
(39)

We follow the method of characteristics and first convert from Eulerican coordinates (x,t) to Lagrangian coordinates (x_0,t_0) , the beginning of a streamline. Here (x_0,t_0) is fixed the parameterization parameter for the streamline emerging at (x_0,t_0) will be denoted by s. Substituting $x=x(s),\,t=t(s)$ and denoting $c(s):=c(x(s),t(s)),\,Q_{\rm so}(s):=Q_{so}(x(s)),\,\phi(s):=\phi(x(s))$ and $c_a(s):=c_a(t(s))$ yields the following system of ODEs [cite Evans]:

$$c'(s) + \frac{Q_{\text{so}}(s)}{\phi(s)}C(s) = \frac{c_a(s)Q_{\text{so}}(s)}{\phi(s)}, \quad c(0) = 0$$
(40a)

$$x'(s) = \frac{q(s)}{\phi(s)},$$
 $x(0) = x_0$ (40b)

$$t'(s) = \phi(s),$$
 $t(0) = t_0,$ (40c)

Here x_0 denotes the starting-point (in space) of the streamline and t_0 denotes the starting-timepoint of the streamline. c(s) hence can be regarded the concentration curve of a particle which originated at (x_0, t_0) and (x(s), t(s)) is the wavefront at artificial timepoint s. For a visualization see Fig. 2.

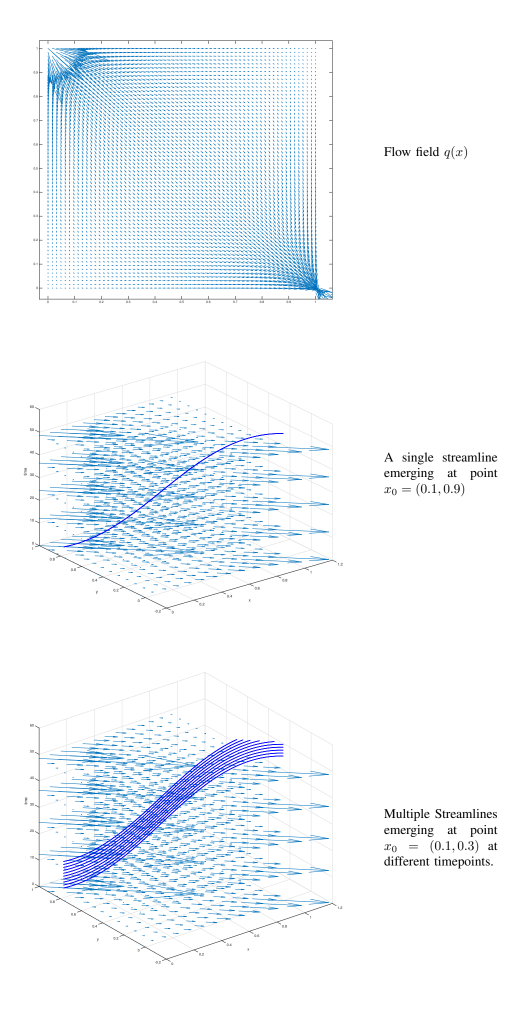


Fig. 2: Flow-Field and Streamlines

Let the source term be non-zero within the ball B_{ϵ} and zero elsewhere. Since contrast-agent is only created in the source, we need to study the problem on streamlines passing through this ball. In order to do this, we place a small ball with radius δ with $\delta << \epsilon$ in the middle of the source. We can now study the solution along streamlines starting at ∂B_{δ} , the boundary of B_{δ} (cf. Fig. 3). Letting $\delta \to 0$, we see that the streamlines connecting to ∂B_{δ} fill whole Ω .

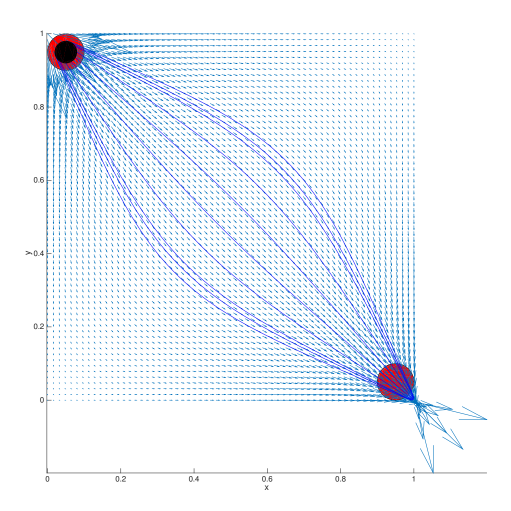


Fig. 3: Streamlines passing through the source. Source- and sink areas are shown in red. Streamlines are supposed to begin on ∂B_{δ} (black)

Hence we conclude that if we know the solution on ∂B_{δ} we know the solution everywhere in Ω . ODE (40a), which governs the behavior along the streamline, has the general solution

$$c(s) = \int_0^s e^{-(R(s) - R(v))} c_a(v) \frac{Q_{so}(v)}{\phi(v)} dv.$$
(41)

Here $R(v) = \int_0^v Q_{\rm so}(x(u))/\phi(x(u))du$. We can interpret R(v) as the amount of contrast agent which is "accumulated" while the particle moves through the source field. Since in our setting $Q_{\rm so}(x) = q_{\rm in}$ for $x \in B_{\epsilon}$ and $Q_{\rm so}(x) = 0$ for $x \notin B_{\epsilon}$, we find that R(v) is constant if v is bigger then some (streamline-dependent) threshold. Backpropagating $C = c\phi$ and inserting

the expression for F results in

$$C(x_0, t) = Q_{so}(x_0) \int_0^t e^{-(t-s)/T} c_{so}(x_0, s) ds$$
(42)

for the mean transit time $T := \phi(x_0)/Q_{so}(x_0)$. A comparison with (37) shows that these expressions are equal provided $P \to Q_{so}$ and $c_{so} \to c_a$. Thus, we have shown that the classical model is equivalent to a field model for tracer transport for specific requirements of the perfusion P and the arterial input concentration c_a . (i) The first observation is that the computed perfusion P is the actual flow at the inlet of the entire domain Ω , and not the inlet of a further spatial sub-division. (ii) The second observation is the fact that the arterial input concentration c_a is identical to the concentration at the source of Ω . This is unproblematic since it is in agreement with the understanding of c_a . However, the first observation regarding the perfusion implies that the computed perfusion along the streamlines will be constant for various t due to stationarity, and hence, computing the voxel wise perfusion for all voxels lying on the streamlines will over-estimate the perfusion with a factor varying with the streamline length. From this argument, we can predict that applying the classical model voxel wise will over-estimate the total perfusion within an arbitrary domain Ω .

Integrating over Ω on both sides of (42), and defining the average concentration $\overline{C}(t) := \frac{1}{|\Omega|} \int_{\Omega} C(x_0, t) dx$,

$$\overline{C}(t) = \frac{1}{|\Omega|} \int_{\Omega} Q_{so}(x_0) \int_0^t e^{-(t-s)/T} c_{so}(x_0, s) ds dx$$
(43)

Since $Q_{so}(x_0)$ is only nonzero across B_{ϵ} we can write

$$\overline{C}(t) = \frac{1}{|\Omega|} \int_{B_{\epsilon}} Q_{so}(x_0) \int_0^t e^{-(t-s)/T} c_{so}(x_0, s) ds dx$$
(44)

For a constant source flow \tilde{Q}_{so} within B_{ϵ} , $\tilde{Q}_{so}=Q(x_0), x_0 \in B_{\epsilon}$, this is equivalent to

$$\overline{C}(t) = \frac{\tilde{Q}_{so}}{|\Omega|} \int_{B_{\epsilon}} \int_{0}^{t} e^{-(t-s)/T} c_{so}(x_0, s) ds dx \tag{45}$$

Further, for a content arterial input concentration $\tilde{c}_{so}(t)$ across B_{ϵ} , $\tilde{c}_{so}(t) := c(x_0, t), x_0 \in B_{\epsilon}$ we can write

$$\overline{C}(t) = \frac{\tilde{Q}_{so}|B_{\epsilon}|}{|\Omega|} \int_0^t e^{-(t-s)/T} \overline{c}_{so}(s) ds$$
(46)

This is recognized as the total perfusion $P = \overline{Q}_{so}|B_{\epsilon}|/|\Omega|$, which is the total source flow divided by the total distributing domain $|\Omega|$. Hence, we can state that the averaging PDE model is consistent with

$$\overline{C}(t) = P \int_0^t e^{-(t-s)/T} \overline{c}_{so}(s) ds, \tag{47}$$

This expression is identical to the classical model in (37) for Ω as the averaging domain and by taking $c_{so}(t) = c_a(t)$.

Thus, we have in this section formally proven that the classical model for perfusion is correct when applying Ω as the averaging domain. On the other hand, applying the same model voxel wise will lead to an over-estimation of the perfusion, with a scaling factor growing along with the streamline length.

II. NUMERICAL IMPLEMENTATION

A. Discretization of the single phase flow model

As a discretizastion scheme for the SF model we apply the two-point fixed approximation (TPFA) finite volume method. This scheme is based on conservation of physical quantities across cell volumes. Denote a grid cell (voxel) as Ω_i and apply to (6) an integral over Ω_i

$$\int_{\Omega_{\delta}} \nabla \cdot \left(-\frac{K}{\mu} \nabla p \right) dx = \int_{\Omega_{\delta}} Q dx. \tag{48}$$

This can in terms of the divergence theorem be rewritten as

$$\int_{\partial\Omega_i} -(\lambda \nabla p) \cdot \nu ds = \int_{\Omega_i} Q dx = F_i. \tag{49}$$

for the conductivities $\lambda = K/\mu$. The integrated source term is per definition named as F_i and has units of absolute flow, m^3/s . Let $\gamma_{ij} = \partial \Omega_i \cap \partial \Omega_j$ be the adjacent face between voxel i and voxel j. Only the component of the pressure gradient perpendicular to γ_{ij} will drive the flow between voxel i and j. The component of ∇p pointing along the normal vector of γ_{ij} can therefore be replaced in terms of the cell centred pressure values p_i and p_j

$$\delta p_{ij} = \frac{2(p_j - p_i)}{\Delta x_i + \Delta x_j} \tag{50}$$

where Δx_i and Δx_j are the cell dimensions in the respective grid direction for voxel i and j, respectively. The total flux v_{ij} across the face γ_{ij} , the left hand side of (49), is thereby approximated by

$$v_{ij} = \delta p_{ij} \int_{\gamma_{ij}} \lambda ds. \tag{51}$$

The conductivies λ are defined at the cell centers, and must be approximated on the faces where the flux is measured. This can be achieved by a distance-weighted harmonic averaging. Let $\lambda_{i,ij} = n_{ij} \cdot \lambda_i n_{ij}$ and $\lambda_{j,ij} = n_{ji} \cdot \lambda_j n_{ij}$ and the directional permeability along n_{ij} is computed as

$$\lambda_{ij} = (\Delta x_i + \Delta x_j) \left(\frac{\Delta x_i}{\lambda_{i,ij}} + \frac{\Delta x_j}{\lambda_{j,ij}} \right)^{-1}.$$
 (52)

Thus, for an orthogonal grid along the main coordinate axes we can approximate the flux across γ_{ij} as

$$v_{ij} = -|\gamma_{ij}|\lambda_{ij}\delta p_{ij} = 2|\gamma_{ij}| \left(\frac{\Delta x_i}{\lambda_{i,ij}} + \frac{\Delta x_j}{\lambda_{j,ij}}\right)^{-1} (p_i - p_j)$$
(53)

where $|\gamma_{ij}|$ is notation for the surface area of γ_{ij} . The terms not depending on the pressure p can be collected into the transmissibilities t_{ij}

$$t_{ij} = 2|\gamma_{ij}| \left(\frac{\Delta x_i}{\lambda_{i,ij}} + \frac{\Delta x_j}{\lambda_{j,ij}}\right)^{-1}.$$
 (54)

From equation (49), summing over all faces γ_{ij} surrounding voxel $i, j \in \mathbb{N}_i$ for the set of neighbour indices \mathbb{N}_i at voxel i, results in a linear equation in the unknowns p_i and p_j

$$\sum_{i \in \mathbb{N}} t_{ij}(p_i - p_j) = F_i \quad \forall \ \Omega_i \subset \Omega$$
 (55)

for each voxel. Denote by $n = n_x n_y n_z$ the total number of voxels within Ω . The left hand side term in (55) yields a band diagonal matrix A for the transmissibilities. Solving the linear system in (55) provides the cell centered voxel wise values of the pressure p_i . Note that the flux is discretized on a staggered grid, corresponding to the voxel faces, whereas the pressure, porosity and permeability are all discretized on a cell-centered grid, as shown in Fig. 4. Thus, the obtained flux field from applying (5) is descretized on a staggered grid, along with the transmissibilities.

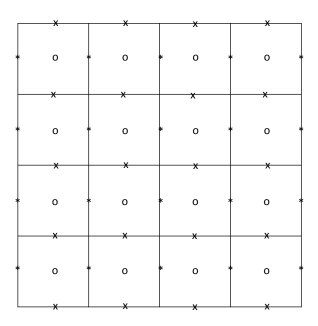


Fig. 4: Grid for discretisation of the flow model in (5) and (6). The circles represent cell centers where the pressure, porosity and permeability are defined. The "x" and the "*" are staggered grid representations for discretisation of the flux in the vertical and horizontal direction, respectively.

B. Discretization of the indicator dilution flow model

Let c_i be the center voxel tracer concentration and c_j the neighbour voxel concentration which the common face γ_{ij} . Consider Eq. (10) for averaged quantities across cell volumes and faces,

$$\phi_i V_i \frac{c_i^{t+1} - c_i^t}{\Delta t} = -\sum_{j=1}^{nf} c_{E_j}^t q_{\gamma_{ij}} |\gamma_{ij}| + c_{so,i}^t F_{so,i} + c_i^t F_{si,i}$$
(56)

where the understanding of $q_{\gamma_{ij}}$ is the normal flux across face γ_{ij} , $c_{\gamma_{ij}}^t$ is the tracer concentration at face γ_{ij} and $c_{so,i}^t$ is the tracer concentration of the source fluid entering the system (the AIF). The terms $F_{so,i} = Q_{so,i}V_i$ and $F_{si,i} = Q_{si,i}V_i$. However, the tracer concentrations are not discretized at the faces but rather at the cell centres, and we must apply an upwinding Godunov scheme for a proper representation. Let

$$c_{E_j}^t = \left\{ \begin{array}{l} c_i^t & \text{if } q_{\gamma_{ij}} > 0, \text{ outward flux} \\ c_j^t & \text{if } q_{\gamma_{ij}} < 0, \text{ inward flux}. \end{array} \right.$$

From (56), this leads to the explicit forward scheme

$$c_i^{t+1} = c_i^t + \frac{\Delta t}{\phi_i V_i} \left[(c_{so,i}^t F_{so,i} + c_i^t F_{si,i}) - \sum_{j=1}^{nf} c_{E_j}^t q_{\gamma_{ij}} |\gamma_{ij}| \right]$$
(57)

for each voxel i. A conversion of c_i into C_i is performed via the relation $C_i = c_i \phi_i$. The overall concentration map C_i is later used for the inverse problem of restoring CBV and CBF.

C. Estimating the streamline length

The streamline length $|\mathbb{C}|$ was computed according to

$$r_{i+1} = r_i + \epsilon q(r_i), \quad i = 0, 1, \dots$$

for a small step length $\epsilon=0.3$ and the parameterised streamline r. Initiating the integration from the source or sink was not possible since the streamlines then are diverging. Instead, we computed a continuous line of equidistant voxels from the source and the sink and initiated the integration in both directions from these positions. Thus, we would instead follow the vector field as it is joining instead of diverging. The length of the streamlines was assigned as the total integration length in both directions towards the source and the sink from the line of equidistant voxels as initial positions r_0 . Due to discretization errors, increasingly multiple assignments of stream line length will take place as approaching the source and sink. For these voxels we assign the minimum length for every voxel.

III. NUMERICAL EXPERIMENTS

A. Simulations on the single phase flow field

A table of parameter values used for the numeral simulations are shown in Table I. We aimed at creating a transparent synthetic test case and kept all optional parameters as simple as possible. Therefore, the permeability and porosity were constant valued in space.

Description	Symbol	Value(s)	Unit
Average input perfusion	\overline{P}	50	ml/min/100ml
Flow source, derived from \overline{P}	F_{so}	0.08333	mm^3/s
Flow sink	F_{si}	$-F_{so}$	mm^3/s
Dynamic viscosity blood [?]	μ_b	$5 \cdot 10^{-6}$	$kPa \cdot s$
Fluid density blood [?]	ρ	1	mg/mm^3
Permeability	k	$5 \cdot 10^{-6}$	mm^2
Porosity	ϕ	0.05	mm^3/mm^3
Spatial resolution	R	(64, 64, 1)	-
Physical dimension	D	(10, 10, 1)	mm
Voxel size	h	(0.156, 0.156, 1)	mm

TABLE I: Parameters used in the numerical experiments, optimized for a slab of the capillary system in the human brain. For the viscosity of non-Newtonian blood we used a suitable setting of dynamic viscosity. The scalar permeability k was added to the diagonal terms of K, such that $K_{ii} = k$ for $i = \{1, 2\}$. The porosity had a fixed value everywhere (cite paper Constantin).

Results from the discretized flow model described in (55) are shown in Fig. 5 for the pressure field p, the flux vector field q and the perfusion P. The pressure field can only be estimated up to a constant and was therefore scaled such that the minimum value is always zero. We used average input perfusion values of $\overline{P} = 50ml/min/100ml$, targeted towards the human brain. For the specified physical dimension D we computed the flow source from the average input perfusion by multiplying with the physical volume D and converting to mm and s as units for space and time. The computed perfusion field P had an average value of 50.5ml/min/100ml, which is close to the average input perfusion \overline{P} used to compute the source flow F_{so} .

The source term was chosen to the upper left voxel and the sink term was chosen in the lower right voxel. The inflow in the source was set negatively to the outflow in the sink, $F_{so} = -F_{si}$. Note that the source term is only nonzero within the domain of the source, and similar for the sink

$$F_i(x) = \begin{cases} F_{so} & \text{if } x_i \in \Omega_{so}, \\ F_{si} & \text{if } x_i \in \Omega_{si}, \\ 0 & \text{elsewhere} \end{cases}$$

for the source domain Ω_{so} and sink domain Ω_{si} .

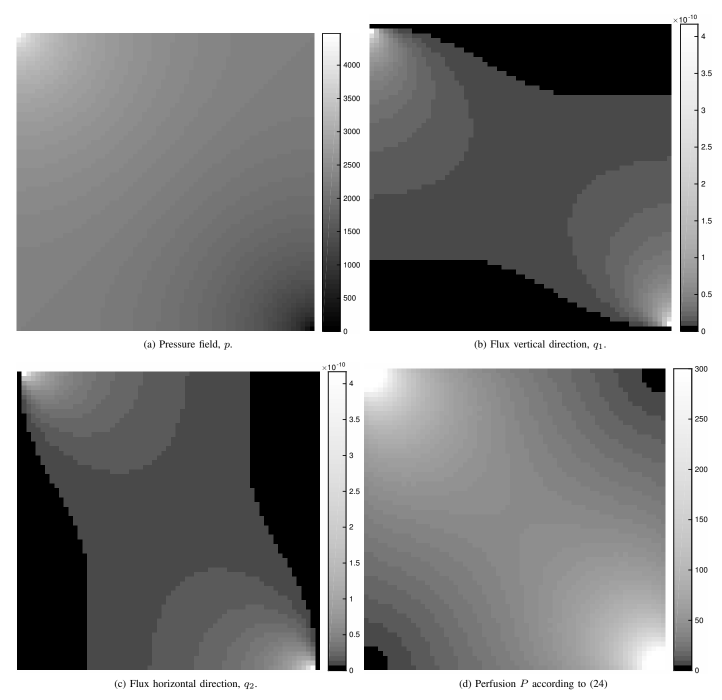


Fig. 5: Synthetic flow model with a source in the upper left corner and a sink in the lower right corner. (a) Pressure field from solving the linear system in (55), (b-c) flux multiplied by voxel face area $q_1h_yh_z$, $q_2h_xh_z$ $[mm^3/s]$ in vertical and horizontal direction with positive directions downward and right, respectively. (c) Voxelwise estimated perfusion [ml/min/100ml] according to (24).

The voxelwise streamline length was computed from the flux field q as described in section II-C. The streamline length for the flux field in Fig. 5 is shown in Fig. 6.

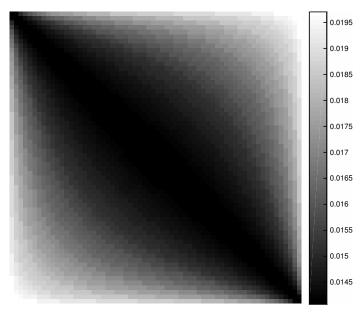


Fig. 6: Length of streamlines $|\mathbb{C}|$ (mm) of the flux field shown in Fig. 5.

The perfusion map in Fig. 5 was computed from Eq. (24) using the flux field shown in Fig. 5 and the length of streamlines shown in Fig. 6.

B. The AIF input curve

For the incoming arterial input function we applied two well established variants of the AIF, First, we model the AIF as an analytical gamma function according to [Wu, Ostergaard]. Second, we use the empirically averaged Parker AIF [Parker 2006]. These two AIF functions are shown in Fig. 7.

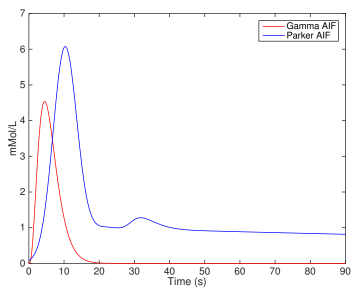


Fig. 7: The two AIF curves used for simulations. The Gamma AIF is an analytical function while the Parker AIF is an averaged response curve observed in real experiments.

C. Simulations on the simulated flow model

The SF model was implemented according to (57) and the previously obtained flux field. The results are shown in Figs. 8 and 9 for the gamma and Parker AIF, respectively.

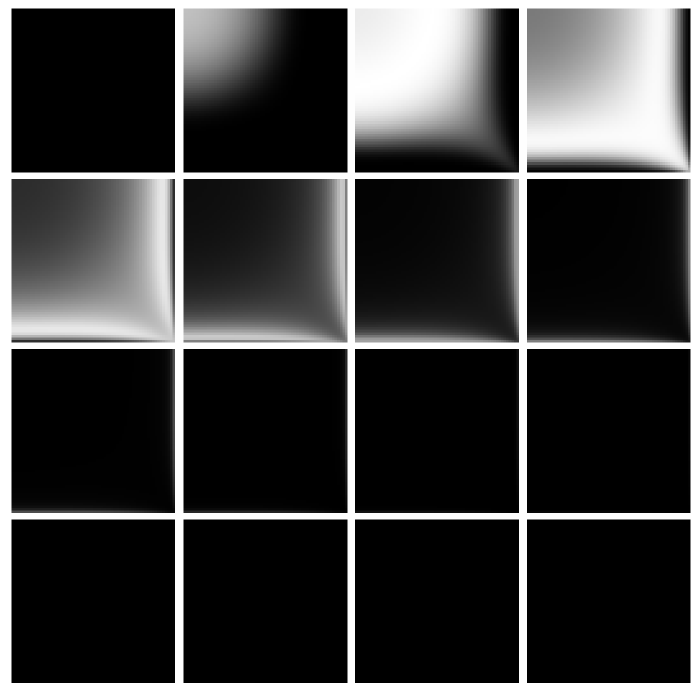


Fig. 8: Development of tracer C(x,t) as a function of time using the gamma AIF. The plots show the tracer concentration for equidistant time points, from top to bottom and left to right of the panels. The entire simulation was run for 90s but only the first half of the simulation is plotted.

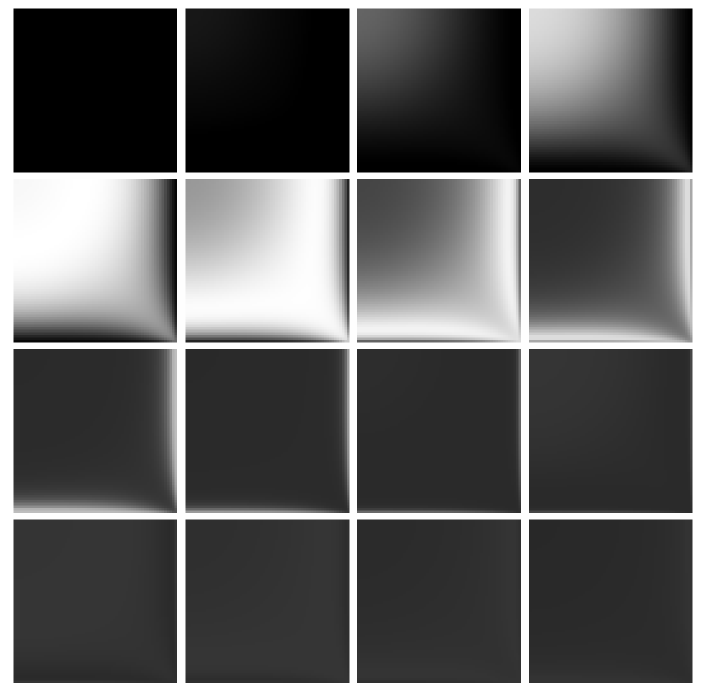


Fig. 9: Development of tracer C(x,t) as a function of time using the Parker AIF. The plots show the tracer concentration for equidistant time points, from top to bottom and left to right of the panels. The entire simulation was run for 90s but only the first half of the simulation is plotted.

D. Experiments on the simulated convolution model

In this section we present experimental data of the SC model Section II-A. We use realistic parameter values of the included parameters, as described in Table I, optimized for the human brain capillary system. We seek to model a small volume of the capillary system with physical dimension of $1cm \times 1cm \times 0.1mm$. This corresponds to a 2D slab with thickness equal to 0.1mm, and has a physically realistic extension in x and y. All tissue parameters are constant in space.

Given the previously estimated voxelwise flow field from the PDE model we want simulate the flow field using (38). However, the classical theory does not take into account the timing of the arrival of the tracer, and the feeding of the indicator starts everywhere at the same time. Therefore we estimate a voxelwise delay field according to the maximum tracer concentrations

within the synthetic field model. Denote this delay field as $\mu(x)$, and we get a slightly modified classical model

$$C(x,t) = P(x) \int_0^t e^{-\frac{t-s}{T}} c_A(s) ds$$
 (58)

From now we also assume that the concentrations are now spatially dependent as C = C(x, t), and similarly for the perfusion P = P(x). The delay field is essentially scaleable with a time-to-peak map and will not change the results of the forthcoming deconvolution but it generates a concentration map which is physically plausible.

Simulation results using the PDE derived flow field as well as the classical model (58) is shown in Figs. ?? and ??.

E. Comparing the SF and SC model in terms of concentration maps

The percentage difference between the SF and the SC model was computed as

$$d_i = 100 \frac{\frac{1}{|\Omega|} \sum_{x_i \in \Omega} \left(C_{SF}(x_i, t_i) - C_{SC}(x_i, t_i) \right)}{\overline{C}_{SF}(x_i, t_i)}$$

$$(59)$$

where $\overline{C}_{SF}(x_i)$ is the voxelwise average value of C_{SF} across time. The average deviation \overline{d} was 88.5% and 31.2% for the gamma and Parker AIF, respectively. These average tracer concentration is plotted over time in Fig. 10 and we see that the convolution partly fails to describe the flow field from the SF model.

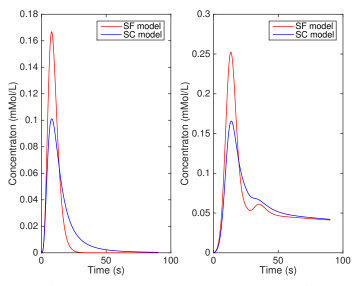


Fig. 10: Average tracer concentration using the SF model and the SC model. Left: Gamma AIF. Right: Parker AIF.

F. The maximum-slope model

The maximum-slope model is the simplest model to recover perfusion. We follow [?] in its the derivation. Given a tissue-curve C(t) and an arterial input function $c_a(t)$ as well as a venous output function $c_v(t)$, it is assumed that for any given timepoint the following relationship holds is fulfilled:

$$C(t + \Delta t) - C(t) \approx P \cdot \Delta t (c_a(t) - c_v(t + \Delta t)).$$

Here $\Delta t(c_a(t) - c_v(t + \Delta t))$ is the total amount contrast-agent introduces in time Δt and $C(t + \Delta t) - C(t)$ is the amount of contrast agent which has been measured to pass through the system. The Maximum-Slope model assumes that these two values are proportional with proportionality constant P, the perfusion. Rearranging and letting $\Delta t \to 0$ yields:

$$P = \frac{C'(t)}{c_a(t) - c_v(t)}.$$

We now assume that we are in an early uptake-phase, where no contrast agent is leaving the system yet. Hence it holds approximately

 $P = \frac{C'(t)}{c_a(t)} \iff Pc_a(t) = C'(t).$

Differentiating this equation with respect to t yields:

$$Pc'_{a}(t) = C''(t).$$

Since this equation is fulfilled for every timepoint t, it is fulfilled especially for timepoint $t_{max} := \operatorname{argmax}_t c_a(t)$. Since at this point it holds that $c'_a(t_{max}) = 0$ and hence also $C''(t_{max}) = 0$, we derive

$$P = \frac{\max_t C'(t)}{\max_t c_a(t)}.$$
(60)

IV. RESTORING THE POROSITY AND FLOW FIELD USING DIFFERENT MODELS

We tested the convolution based classical model (38) as well as maximum-slope model (MS, (60)) for their capability to recover the perfusion values. The deconvolution was performed using the block circulant SVD (bSVD) approach described in [?], which comes with a time-invariance with respect to arrival times. The success of the restoration was measured in terms of the relative error of the recovered perfusion with respect to the true perfusion

$$RE := \frac{P_{rec} - P_{true}}{P_{true}} \cdot 100\%.$$

The input data was downsampled to a final time-resolution of 1 s. In order to simulate different spatial resolutions, the data was spatially averaged over different block-sizes ranging from $b_1 = (1,1)$ to $b_5 = (64,64)$. Results are displayed in Figure 12,11 as well as in Table II. It can be observed that for the complete domain the perfusion can be restored by both methods accurately with an error of $\approx 2\%$. The impuls-response function reconstructed from the perfusion method is displayed in Figure 13. However, within the capillary field the model assumptions of one inlet and one outlet are violated, causing large errors in the recovered flow field. Typical impuls response functions of the capillary field are also visualized in Figure 13. These recovered gaussian-like functions seem to be describing dispersion effects, yielding a smoothing and broadening of the arterial-input.

TABLE II: Results of the perfusion restoration with median relative errors in percent.

	Model	Block Size				
		(1,1)	(5,5)	(10,10)	entire domain	
SFD: Perfusion	MS	170.09	165.03	158.57	2.65	
	bSVD	859.06	768.58	664.84	-1.25	
SCD: Perfusion	MS	-23.20	-24.26	-25.75	37.96	
	bSVD	2.43	-4.27	-8.80	20.72	
SFD: CBV		$4.37 \cdot 10^{-5}$	$4.37 \cdot 10^{-5}$	$4.37 \cdot 10^{-5}$	$4.37 \cdot 10^{-5}$	
SCD: CBV		$1.94 \cdot 10^{-2}$	$1.93 \cdot 10^{-2}$	$2.12 \cdot 10^{-2}$	1.13	

Fig. 11: Results of the restoration for the PDE Model (SFD)

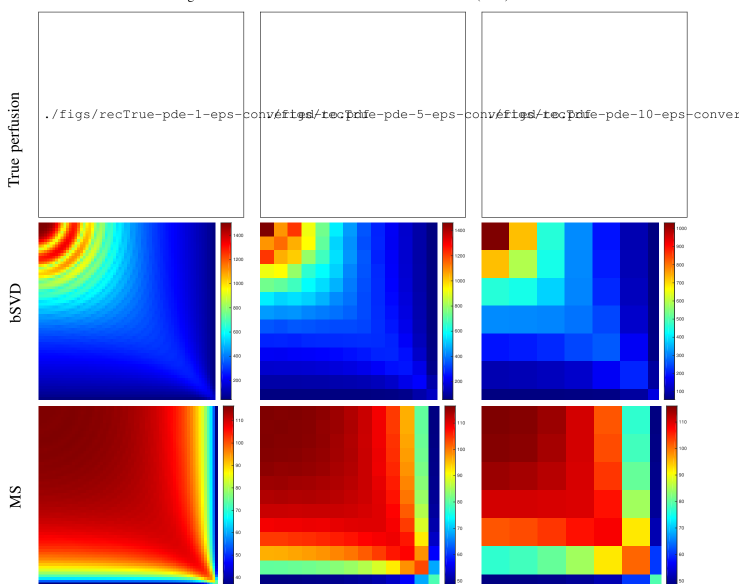


Fig. 12: Results of the restoration for the Convolution Model (SCD)

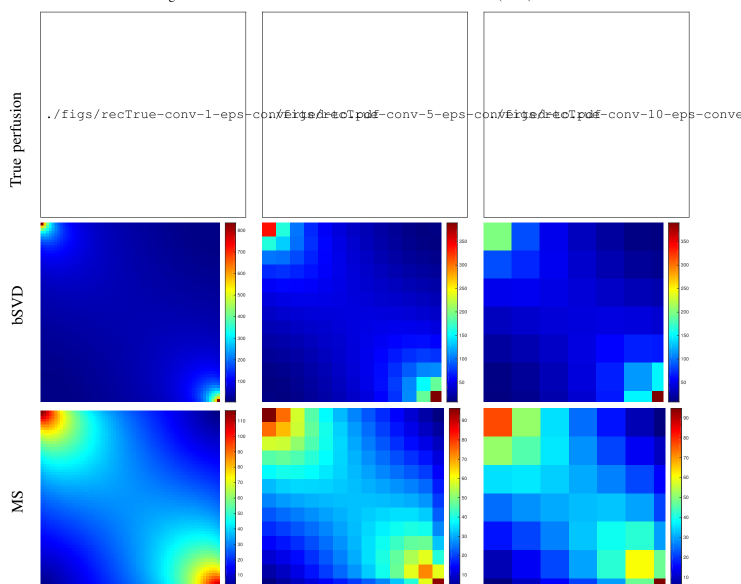


Fig. 13: Deconvolution results

