Hematocrit dispersion in asymmetrically bifurcating vascular networks

Krishna Sriram, Marcos Intagliaetta, and Daniel M. Tartakovsky

1Department of Mechanical and Aerospace Engineering, University of California, San Diego, La Jolla, California; and 2Department of Bioengineering, University of California, San Diego, La Jolla, California

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Sriram K, Intaglietta M, Tartakovsky DM. Hematocrit dispersion in asymmetrically bifurcating vascular networks. Am J Physiol Heart Circ Physiol 307: H1576–H1586, 2014. First published September 12, 2014; doi:10.1152/ajpheart.00283.2014—Quantitative modeling of physiological processes in vasculatures requires an accurate representation of network topology, including vessel branching. We propose a new approach for reconstruction of vascular network, which determines how vessel bifurcations distribute red blood cells (RBC) in the microcirculation, particularly in the smallest segments of the precapillary arteriolar network (24, 30, 42). This deviation from Murray’s law has been attributed to the non-Newtonian shear-thinning behavior of blood in the vessels of small radii (1, 45). Murray’s law fails to capture such a behavior, since it is derived by assuming blood to be a Newtonian fluid, whose flow within each vessel obeys the Poiseuille law (35). Alternative optimality criteria used to describe vascular bifurcations include the minimum energy hypothesis (24) and generalizations of Murray’s law that account for the role of muscle tone (53), alternative blood rheology (1, 45), and turbulent (54) or pulsatile (37) flow conditions. These and other similar optimality criteria aim to predict the radii of daughter vessels, relying on empirical closure assumptions to prescribe partitioning of suspended red blood cells (RBCs) between daughter vessels.

Both blood viscosity and its shear-thinning behavior vary with concentrations of dissolved chemicals, e.g., fibrinogen, and density of RBCs in the blood column, i.e., hematocrit. Of direct relevance to the present study are observations suggesting that blood viscosity and shear-thinning behavior increase with hematocrit (38, 41, 44, 47). This phenomenon was ignored by Revellin et al. (45), who modified Murray’s law by treating blood as an Ostwald de Waele fluid whose rheology and apparent viscosity are independent of either hematocrit or vessel radius. The latter assumptions contradict in vitro (41) and in vivo (44) observations that revealed the strong dependence of apparent viscosity on both hematocrit and vessel radius. Alarcon et al. (1) accounted for these effects by employing the Pries et al. (44) constitutive relation, according to which apparent blood viscosity varies with vessel radius and hematocrit. In applying this generalization of Murray’s law to modeling a network, they assumed that hematocrits in daughter vessels at bifurcations are given by the ratio of average velocities in each daughter vessel. This leads to predictions of hematocrit values in the terminal regions of the network, which are unrealistically low (1).

The question of how hematocrit is partitioned between the parent and daughter vessels remains open. In vivo and in vitro experimental data on hematocrit partitioning at bifurcations typically relate the flux fraction $F_{\text{RBC}}$, i.e., the fraction of RBCs flowing from the parent vessel into the larger daughter vessel, to the flow fraction $F_{\text{blood}}$, i.e., the fraction of total fluid flow from the parent vessel that enters the larger daughter vessel (see Ref. 4 and the references therein). Mathematical models (2, 3) of hematocrit partitioning at bifurcations are limited to two-dimensional channel flows. They suggest an approximately linear dependence of $F_{\text{RBC}}$ on $F_{\text{blood}}$ over a wide range of $F_{\text{blood}}$. While some experimental studies (12, 16, 26, 50) observed a linear relation between $F_{\text{RBC}}$ and $F_{\text{blood}}$, others (22, 40, 43) found this relationship to be nonlinear. Even when the linear behavior is observed, the corresponding slopes and
intercepts tend to be different. Furthermore, experimental data indicate that endothelial dysfunction significantly affects the distribution of hematocrit at arterial bifurcations with important implications for the adequacy of tissue perfusion (14, 15). Thus an analysis of baseline RBC partition at bifurcations solely dependent on network properties is warranted (17).

We propose a mathematical framework for construction of vascular networks, which possess both optimal daughter vessel radii and optimal partition of hematocrit between daughter vessels. These two goals are achieved by postulating that healthy vasculatures are constructed in a way that optimizes oxygen delivery to the surrounding tissue. Our model builds upon the analysis of Alarcon et al. (1) in the sense that it generalizes Murray’s law by accounting for both the non-Newtonian nature of blood flow in microcirculation and the dependence of blood rheology on hematocrit. Unlike Alarcon et al. (1), we impose no prior restrictions on the hematocrit partition between daughter vessels. Instead it is determined by solving an optimization problem and describes how vessel bifurcations distribute RBCs in the microcirculation.

The outcome of our model is a vascular network in which both the bifurcation asymmetry and WSS vary from one generation of the network to the next. We demonstrate that the resulting vascular networks satisfy a number of properties of healthy vasculatures constructed in a way that optimizes oxygen delivery to the tissue downstream of the bifurcation.

STATE-OF-THE-ART IN VASCULATURE REPRESENTATION

Optimal vessel radius. The starting point of our analysis is the Murray cost function (34),

\[
Q = \frac{JR^4}{8\pi\mu(R, H)}.
\]

In the \(n\)th generation of the network, a parent vessel of radius \(R_n\) bifurcates into smaller daughter vessels with radii \(R_{n,1}\) and \(R_{n,2}\). The discharge hematocrit \(H_n\) in the parent vessel partitions into the discharge hematocrits \(H_{n,1}\) and \(H_{n,2}\) in the corresponding daughter vessels. If the oxygen flux in a parent vessel is \(Q_{O_2}\), then mass conservation requires the oxygen fluxes in its daughter vessels to be \(aQ_{O_2}\) and \((1 - a)Q_{O_2}\), where the flux-asymmetry parameter \(a\) is a number between 0 and 1. In the following sections, we compute the daughter vessel radii and hematocrits by postulating that daughter vessels bifurcate in a way that minimizes the total cost associated with oxygen delivery to the tissue downstream of the bifurcation.

Fig. 1. Dependence of the normalized apparent blood viscosity. \(\mu/\mu_p\) on the vessel radius \(R\) and hematocrit \(H\) predicted with the rheological law of Pries et al. (44).
\[ W = QJL + \alpha \pi R^2L \] (5)

which combines the mechanical work \((QJL)\) necessary to drive blood through a blood vessel of radius \(R\) and length \(L\) with the “metabolic cost” \((\alpha \pi R^2L)\). The latter is linearly proportional to the vessel’s volume \(\pi R^2L\) with the coefficient of proportionality \(\alpha\). According to Murray’s law, blood vessels have radii that minimize the cost function \(W\) for given flow rate \(Q\) and pressure gradient \(J\), i.e., they satisfy an equation \(dW/dR = 0\). Combined with Eq. 4, this defines the optimal vessel radius \(R^*\) as a solution of

\[ \frac{d}{dR} \left( \frac{8\mu Q^2}{\pi R^4} + \alpha \pi R^2 \right) = 0. \] (6)

If the blood viscosity \(\mu\) were independent of the vessel radius \(R\), this equation would yield Murray’s law, according to which \(Q\) is proportional to \(R^3\) (34). For small \(R\) (larger than 200 \(\mu\)m), the correction factor \(\Lambda\) is approximately constant (Fig. 2) and Murray’s law is recovered. For physiological values of hematocrit and vessel radii typically seen in the microcirculation (less than 100 \(\mu\)m), \(\Lambda\) has a strong dependence on \(R\) that is not captured by Murray’s law.

Substituting the optimal \(Q\) and \(R^*\) into Eq. 5 gives the optimal (minimum) cost of supplying a volume of blood to a block of tissue,

\[ W^* = \alpha \pi LR^2 \left[ \frac{2\mu (R^*, H)}{\Lambda (R^*, H)} + 1 \right]. \] (8)

This analysis enables one to determine the optimal radius of a single blood vessel, if the discharge hematocrit \(H\) and the flow rate \(Q\) are known. Determining the radii of daughter vessels (and hematocrits) from a bifurcating parent vessel requires additional assumptions.

Models of vessel bifurcation. The bifurcation of a parent vessel of radius \(R_n\) into daughter vessels with radii \(R_{n,1}\) and \(R_{n,2}\) is accompanied by the partitioning of the discharge hematocrit \(H_n\) in the parent vessel into the discharge hematocrits \(H_{n,1}\) and \(H_{n,2}\). The volumetric blood flow rates in the parent \((Q_n)\) and daughter \((Q_{n,1} \text{ and } Q_{n,2})\) vessels satisfy mass conservation,

\[ Q_n = Q_{n,1} + Q_{n,2}. \] (9)

Mass conservation of red blood cells imposes a constraint on the discharge hematocrits in the parent parent \((Q_n)\) and daughter \((Q_{n,1} \text{ and } Q_{n,2})\),

\[ Q_n H_n = Q_{n,1} H_{n,1} + Q_{n,2} H_{n,2}. \] (10)

The volumetric flow rates in each vessel are given by the modified Murray’s law (Eq. 7)

\[ Q_n = \frac{\pi \sqrt{\alpha R_n^3}}{2 \Lambda (R_n, H_n)}; \quad Q_{n,1} = \frac{\pi \sqrt{\alpha R_{n,1}^3}}{2 \Lambda (R_{n,1}, H_{n,1})}; \quad Q_{n,2} = \frac{\pi \sqrt{\alpha R_{n,2}^3}}{2 \Lambda (R_{n,2}, H_{n,2})}. \] (11)

Substituting Eq. 11 into Eqs. 9 and 10 yields

\[ \frac{R_n^3}{\Lambda (R_n, H_n)} = \frac{R_{n,1}^3}{\Lambda (R_{n,1}, H_{n,1})} + \frac{R_{n,2}^3}{\Lambda (R_{n,2}, H_{n,2})} \] (12a)

and

\[ \frac{H_n R_n^3}{\Lambda (R_n, H_n)} = \frac{H_{n,1} R_{n,1}^3}{\Lambda (R_{n,1}, H_{n,1})} + \frac{H_{n,2} R_{n,2}^3}{\Lambda (R_{n,2}, H_{n,2})}. \] (12b)

Since these two equations contain four unknowns \((R_{n,1}, R_{n,2}, H_{n,1}, \text{ and } H_{n,2})\), the determination of the optimal bifurcation radii requires additional assumptions. For example, one can postulate that the daughter vessels have identical radii, \(R_{n,1} = R_{n,2} = R_{n,d}\) (i.e., \(Q_{n,1} = Q_{n,2} = Q_{n,d}\)), and assume that the discharge hematocrit in all the vessels is the same, \(H_n = H_{n,1} = H_{n,2} = H\). This model implies that the discharge hematocrit remains constant throughout the vascular network and relates the daughter-vessel radius to the radius of its parent by an implicit relation \(2R_{n,d}^3 / \Lambda (R_{n,d}, H) = R_n^3 / \Lambda (R_n, H)\). Unfortunately symmetrically bifurcating networks are not representative of typical vasculatures.

The construction of asymmetrically bifurcating vascular networks relies on Eqs. 12a and 12b, supplemented with the following two assumptions. First, one assumes the ratio of the two daughter-vessel radii, \(R_{n,1}/R_{n,2}\), to be known \((1, 24, 36)\). Second, one postulates a constitutive relation that governs the partition of discharge hematocrit between the two daughter vessels \((1)\). The shortcomings of the latter assumption are discussed in the introduction.

**BIFURCATIONS OPTIMIZED FOR OXYGEN DELIVERY**

We posit that biological vascular networks are structured in a way that maximizes their ability to deliver oxygen. Specifically, we postulate that 1) asymmetric bifurcations occur because the volumes of tissue downstream of each daughter vessel have different oxygen needs; 2) these needs are quantified by a known constant \(a\) \((0 < a < 1)\), which serves to partition the oxygen flux \(Q_{O_2}\) in any given parent vessel into the oxygen fluxes \(aQ_{O_2}\) and \((1 - a)Q_{O_2}\) in its two daughter vessels.
vessels; 3) an optimal bifurcation is one in which the oxygen demands of each downstream tissue volume are supplied at a “minimal total cost”; and 4) the amount of oxygen transported through a blood vessel is proportional to the number of RBCs flowing through that vessel (51), i.e., the oxygen flux is given by \( Q_{O_2} = \rho Q_n H_n \), where the constant \( \rho \) denotes the amount of oxygen transported by a unit volume of RBCs. The value of \( \rho \) depends on the physiology of the downstream volumes of tissue supplied by each daughter vessel (symmetric bifurcations imply that these volumes are identical, so that \( \rho = 0.5 \)).

If the oxygen flux in the \( n \)th vessel is \( Q_{O_2} \), then the oxygen fluxes in the two daughter vessels are \( a Q_{O_2} = \rho Q_n H_n,1,1 \) and \( (1-a)Q_{O_2} = \rho Q_n H_n,2,2 \). It follows from Eq. 11 that the oxygen flux in the \( n \)th vessel,

\[
Q_{O_2} = \frac{\pi \sqrt{\alpha}}{2} \frac{H_n R_n^3}{\rho} \frac{H_n R_n^3}{\rho} \lambda (R_n, H_n),
\]

(partitions into the oxygen fluxes in its two daughter vessels, \( a Q_{O_2} \) and \( (1-a) Q_{O_2} \), according to

\[
a Q_{O_2} = \frac{\pi \sqrt{\alpha}}{2} \frac{H_n R_n^3}{\rho} \lambda (R_n, H_n,1,1) \quad \text{and} \quad (1-a) Q_{O_2} = \frac{\pi \sqrt{\alpha}}{2} \frac{H_n R_n^3}{\rho} \lambda (R_n, H_n,2,2).
\]

A constraint on the hematocrit partitioning between the two daughter vessels is obtained by substituting \( Q_n = a Q_{O_2} / (\rho H_n,1) \) and \( Q_{n,2} = (1-a) Q_{O_2} / (\rho H_n,2) \) into Eq. 9, which yields

\[
1 = \frac{a}{H_n,1} + \frac{1-a}{H_n,2}.
\]

Three Eqs. 14 and 15 contain four unknowns \( (R_n,1, R_n,2, H_n,1, H_n,2) \). The fourth equation needed to close this system is obtained by assuming that vascular networks are formed in a way that minimizes the work necessary to distribute oxygen throughout the vasculature. The cumulative work of forcing the blood through the two daughter vessels of the \( n \)th parent vessel is computed from Eq. 5 as

\[
W = \left( Q_{n,1} L_n,1 + \alpha \pi R_n^2 L_n,1 \right) + \left( Q_{n,2} L_n,2 + \alpha \pi R_n^2 L_n,2 \right)
\]

where \( R_n,1, R_n,2, L_n,1, L_n,2 \) are the (yet unknown) lengths of the daughter vessels. By analogy with Eq. 8, for any given partitioning of the hematocrit the minimum Murray’s work has the form

\[
W^* = \alpha \pi L_n,1 R_n^2,1 \left[ \frac{2 \mu (R_n,1, H_n,1)}{\Lambda^2 (R_n,1, H_n,1)} + 1 \right] + \alpha \pi L_n,2 R_n^2,2 \left[ \frac{2 \mu (R_n,2, H_n,2)}{\Lambda^2 (R_n,2, H_n,2)} + 1 \right].
\]

The optimal hematocrit partitioning minimizes the total work in Eq. 16, giving rise to a fourth equation,

\[
\frac{dW^*}{dH_n,1} = 0,
\]

where Eq. 15 is used to express \( H_n,2 \) in terms of \( H_n,1 \).

The system of Eqs. 14, 15, and 17 remains unclosed due to the presence of two additional unknowns: the daughter vessel lengths \( L_n,1 \) and \( L_n,2 \). These are often related to the corresponding vessel radii \( R_n,1 \) and \( R_n,2, \) e.g., by assuming the radius-to-length ratio \( R_n,1/L_n,1 = R_n,2/L_n,2 \) to be constant throughout the vascular network (i.e., for any \( n \)) (10, 11, 24, 36). This assumption is supported by in vivo morphological studies of arterial trees (19, 32). Rather than forcing the radius-to-length ratio to be constant, we supplement the four postulates listed above with the following hypothesis: 5) a blood vessel’s volume, \( \pi R^2 L \), is linearly proportional to the volume of tissue it oxygenates.

We show in the Appendix that this assumption leads to a radius-to-length relationship,

\[
L = \frac{\kappa HR}{\lambda (R, H)} \quad (18)
\]

where \( \kappa \) is a constant model parameter. Setting \( \kappa = 650 \sqrt{\mu} \) in Eq. 18 results in the \( LR \) ratios between 50 and 100 depending on the value of discharge hematocrit \( H \) (Fig. 3), which falls within the range of the reported length-to-radius ratios (10, 19, 24, 36). Figure 3 reveals that in blood vessels with \( R > 150 \mu m \), the length-radius ratios do become constant, with their value decreasing with hematocrit \( H \). This leads us to conclude that the length-to-radius ratio \( (LR) \) may be assumed constant over the bulk of a vascular tree, with deviations from its constant value occurring in small, precapillary arterioles.

Given values of the discharge hematocrit \( H_n \) and the oxygen flux \( Q_{O_2} \) in the parent vessel \( n \) and its radius \( R_n \), the radii \( (R_n,1, R_n,2) \) and lengths \( (L_n,1, L_n,2) \) of, and the hematocrits \( (H_n,1, H_n,2) \), in the bifurcating daughter vessels are uniquely determined by the system of nonlinear Eqs. 14, 15, 17, and 18.

The radii of the daughter vessels, \( R_n,1, R_n,2, \) predicted with our model are reported in Fig. 4 in terms of their ratio \( R_n,1/R_n,2 \). The symmetric bifurcation \( (R_n,1 = R_n,2) \) occurs when the bifurcation parameter \( a = 0.5 \). The values \( 0.5 < a < 1 \) result in \( R_n,1 < R_n,2 \), while values \( 0 < a < 0.5 \) (not shown in Fig. 4) yield \( R_n,1 < R_n,2 \). The bifurcation asymmetry increases with the parent vessel’s radius \( R_n \), as long as \( R_n \leq 80 \mu m \). After that threshold the ratio \( R_n,1/R_n,2 \) is independent of \( R_n \), so that the curves \( R_n,1/R_n,2 vs. a \) overlap with that for \( R_n = 80 \mu m \).

Figure 5 shows the partitioning of hematocrit \( H_n \) = 0.45 in the parent vessel into hematocrits \( H_n,1, H_n,2 \) in the daughter vessels for several values of the parent vessel radius \( R_n \) and the

![Fig. 3. Variation of length-to-radius ratio LR with vessel radius R, for several values of discharge hematocrit H. The length-to-radius ratio LR may be assumed constant over the bulk of a vascular tree, with deviations from its constant value occurring in small, precapillary arterioles.](image-url)
bifurcation parameter $a$. While hematocrit in the larger daughter vessel ($H_{n,1}$) is nearly the same as hematocrit in the parent vessel ($H_n$), hematocrit in the smaller daughter vessel ($H_{n,2}$) is significantly different from $H_n$. This is consistent with the observations reported in Refs. 12, 16, and 22 and supports the idea that discharge hematocrit at the bifurcation partitions in a way that minimizes the work necessary to induce blood flow in microcirculation.

Dependence of the hematocrit ratio $H_{n,1}/H_{n,2}$ on the parent vessel radius $R_n$ is elucidated further in Fig. 6. Bifurcations of parent vessels with $R_n < 40 \mu m$ result in the hematocrit ratios $H_{n,1}/H_{n,2} > 1$, i.e., hematocrit in the larger daughter vessel exceeds hematocrit in the smaller daughter vessel. The situation is reversed in parent vessels with $R_n > 40 \mu m$, which yield $H_{n,1}/H_{n,2} < 1$. As $R_n$ increases, the hematocrit ratio $H_{n,1}/H_{n,2}$ asymptotically tends to 1. The inflection point of the $H_{n,1}/H_{n,2}$ vs. $R_n$ curves corresponds to the inflection point in the relationship between the apparent viscosity and vessel radius in Fig. 1.

Experimental data on flow behavior at vessel bifurcations are typically reported in terms of the RBC flux fraction $F_{RBC}$ and the blood flow fraction $F_{blood}$ (12, 16, 40, 50). These are defined as

$$F_{RBC} = \frac{Q_n}{H_n} \frac{H_{n,1}}{Q_{n,1}}$$

$$F_{blood} = \frac{Q_n}{H_n} \frac{Q_{n,1}}{Q_{n,2}}$$

Our model predicts the RBC flux fraction $F_{RBC}$ to vary linearly with the flow fraction $F_{blood}$ (Fig. 7), in agreement with the results reported in Refs. 12, 16, 50 but disputed by others (22, 40, 43). Our model also indicates that the relationship between $F_{RBC}$ and $F_{blood}$ is relatively insensitive to $R_n$ and $H_n$. Moreover, Fig. 7 suggests that $F_{RBC} \approx F_{blood}$ for all cases which, combined with Eq. 19, implies that $H_{n,1}$ is within a few percent of $H_n$ (see also Figs. 5 and 6). However, the analysis in the section below demonstrates that these small differences in hematocrit cannot be neglected since they accumulate from one generation of vessels to the next, resulting in large intravessel variability of hematocrit in terminal sections of the vascular network.

RESULTS

Comparison with Murray’s law. The radii of the daughter vessels are but one metric by which to compare the vascular networks predicted with our model and that given by Murray’s law; while the former requires one to solve a system of nonlinear Eqs. 14, 15, 17, and 18 in order to obtain these radii,
the latter is given by a closed-form relation \( R_{n}^{3} = R_{n,1}^{3} + R_{n,2}^{3} \).

Other metrics include the distributions of pressure \( P \) and WSS \( \tau \) throughout the vascular network.

According to our model, the WSS \( \tau_{n} = J_{n}R_{n}/2 \) in an \( n \)th generation vessel is computed from Eqs. 4 and 7 as

\[
\tau_{n} = 2\pi^{2}\sqrt{\frac{\mu(R_{n},H_{n})}{\Lambda(R_{n},H_{n})}}.
\]

(20)

Since \( J = \Delta P/L \), it follows from Eqs. 4, 7, and 18 that the pressure drop \( \Delta P \) across a vessel of length \( L \) is given by \( \Delta P = 4\kappa\pi^{2}\sqrt{\alpha H\mu/L^{2}} \). Let us suppose that a vascular network consists of \( N \) generations of vessels, and ends in the capillary bed where the blood pressure is \( P_{\text{cap}} \). Then intraluminal pressure \( P_{n} \) at the start of an \( n \)th generation vessel is

\[
P_{n} = P_{\text{cap}} + \sum_{i=0}^{n-1} \Delta P_{N-i}, \quad \Delta P_{m} = 4\kappa\pi^{2}\sqrt{\alpha \frac{H_{m}\mu(R_{m},H_{m})}{\Lambda(R_{m},H_{m})}}.
\]

(21)

A vascular network that obeys Murray’s law has the constant length-to-radius ratio \( \bar{e} = L/R \), and the bifurcation relation \( R_{n}^{3} = R_{n,1}^{3} + R_{n,2}^{3} \) for all \( n \leq N \). Murray (34, 35) and subsequent studies (49) treat the viscosity of blood, \( \mu^{M} \), flowing through such a network as constant. Under these assumptions, Eqs. 4 and 7 predict the WSS \( \tau^{M} \) that is constant throughout the vasculature,

\[
\tau^{M} = \pi^{2}\sqrt{\frac{\alpha \mu^{M}}{\Lambda^{M}(R_{m},H_{m})}}.
\]

(22)

and the intraluminal pressure \( P_{n}^{M} \) in an \( n \)th generation vessel that is given by

\[
P_{n}^{M} = P_{\text{cap}} + 2n\pi^{2}\sqrt{\frac{\mu^{M}}{\Lambda^{M}(R_{m},H_{m})}}(N - n + 1).
\]

(23)

In the simulations reported in Fig. 8, we consider a symmetrically bifurcating network \( (a = 0.5 \text{ or } R_{n,1} = R_{n,2} \text{ for all } n \leq N) \) that consists of \( N = 22 \) generations, terminating with capillaries of radius \( R_{N} = 3.0 \text{ \( \mu \text{m} \)} \). The capillary pressure is set to \( P_{\text{cap}} = 30.0 \text{ \text{mmHg} (9).} \) To facilitate the comparison between the two models, we chose the value of the constant blood viscosity \( \mu^{M} \) in the Murray model to coincide with the asymptotic value of \( \mu(R,H) \), which corresponds to large vessel radii. Setting \( H = 0.45 \) and \( R = 1,000.0 \text{ \text{m}} \), this gives \( \mu^{M} = 3.198 \text{ cP.} \)

The vessel radii of the vascular networks reconstructed with the two models are shown in Fig. 8B. The difference between the two predictions exceeds 20% for the larger vessels (early generations of the network). The predicted distributions of the WSS \( \tau \) (Fig. 8A) highlight the physiological differences between the two models. While Murray’s law implies a constant WSS \( \tau^{M} \) across the entire network (37), our model captures the experimentally observed variability in the WSS \( \tau \) between the vessels of different generations. Specifically, it predicts the amplification of the WSS in the microcirculation (vessel radii \( R < 25 \text{ \( \mu \text{m} \)} \)), wherein the WSS appreciably increases as the vessel radii \( R \) become smaller (the vessel generation \( n \) becomes larger), reaching its maximum in the terminal, precapillary arterioles. This behavior is in agreement with the observed amplification of WSS in the microcirculation (24, 30, 31, 42). In larger vessels \( (R > 25 \text{ \( \mu \text{m} \)} \), the WSS increases by a small amount with \( R \), reaching a constant value in large arterioles. This is in general agreement with the observations reported in Refs. 42 and 52.

Both models predict that the intraluminal blood pressure \( P \) increases with vessel radius \( R \), with the bulk of the pressure drop occurring in the smaller vessels (Fig. 8A). As \( R \) approaches the values typical of large arterioles and small arterioles.
ies, blood pressure in these vessels becomes almost equal to systemic arterial pressure. This indicates that the smaller arterioles contribute most to vascular resistance. These small vessels are often referred to as “resistance vessels.” The predicted dependence of intraluminal pressure $P$ vessel radius $R$ is in qualitative agreement with the observations (9, 30, 42).

Impact of branching asymmetry. The results presented in Fig. 8 are obtained for a symmetric vascular network (the bifurcation parameter $a = 0.5$). Figure 9 demonstrates the effect of network asymmetry on the distributions of vessel radii and hematocrit throughout the networks with $a = 0.7$ and $a = 0.9$. The largest vessel in these simulations has radius $R_1 = 500.0 \, \mu m$ and hematocrit $H_1 = 0.45$. Each branch of the network is assumed to terminate once daughter vessel radius reaches $R = 3.0 \, \mu m$, corresponding to termination of the arteriolar network, which feeds the capillary bed. The network asymmetry ($a \neq 0.5$) causes the number of vessels in each branch to vary. This implies that asymmetric vascular networks defy idealized fractal descriptions, which is in line with several in vivo studies (20, 24, 26, 36, 39). Only the first seven generations of the vessels (i.e., before any branch reaches the $R = 3.0 \, \mu m$ threshold) are shown in Fig. 9, with each circle representing a blood vessel with the radius $R$ or hematocrit $H$ at a given vessel generation $n$. Note that the number of bifurcating vessels in each generation increases as $2^{n-1}$, which might not be apparent in Fig. 9 since many of the data points overlap.

Our model predicts a large variability of hematocrit values across the vessels of the same generation (Fig. 9), including at the terminal regions of the network and in the capillary beds supplied by these terminal branches. This finding is supported by the in vivo measurements in capillary beds (20, 26, 39) that show large variations of observed hematocrit values across capillary beds, even among vessels of similar diameters. Our model accounts for this effect by allowing for the asymmetric partitioning of hematocrit at every bifurcating vessel.

A measure of the deviation of vascular network from Murray’s law is provided by a network branching exponent $\xi$ defined such that

$$\sum_{i=1}^{2^{n-1}} R_i^\xi \approx \sum_{i=1}^{2^{n-1}} (R_{i,1}^\xi + R_{i,2}^\xi).$$

(Murray’s law states that the sum of powers of all the vessel radii in the $n$th generation must equal to the sum of the powers of all the vessel radii in the $(n + 1)$-th generation, and sets $\xi = 3$. Reported values of $\xi$ range from 2.7 to 3 (10, 27, 28, 33, 36, 46), which suggests small but meaningful deviations from Murray’s law. For the networks presented in Fig. 9, Eq. 24 holds across all generations with $\xi = 2.96$ for $a = 0.7$ and $\xi = 2.84$.)

Fig. 9. Variation of vessel radii $R$ and discharge hematocrit $H$ within the first $n = 7$ generations of asymmetric vascular networks with bifurcation parameters $a = 0.7$ and $a = 0.9$. 

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2.97 for \( a = 0.9 \). These values are within 1% of the values reported in Refs. 27 and 33.

Model validation. The vascular networks constructed with using our model exhibit the following characteristics observed in a number of in vivo and in vitro studies. 1) RBC flux fraction \( F_{\text{RBC}} \) at bifurcations varies approximately linearly with the blood flow fraction \( F_{\text{blood}} \) over a broad range of hematocrits and vessel radii. This is in agreement with the data reported in Refs. 3, 12, 16, and 50. 2) The asymmetry in discharge hematocrit at asymmetric bifurcations increases with the degree of asymmetry. This is in agreement with the data reported in Refs. 12, 16, and 22. 3) WSS is nonuniform in the microcirculation, significantly increasing in the terminal sections of the network. This is in agreement with the data reported in Refs. 24, 30, 31, 37, and 42. 4) Pressure gradients increase sharply in the terminal sections of the network. This is in agreement with the data reported in Refs. 9 and 30. 5) Both the hematocrit partitioning at vessel bifurcations and hematocrit across the capillary bed in asymmetric networks is nonuniform. This is in agreement with the data reported in Refs. 20, 26, and 39. 6) Microvascular discharge hematocrit is comparable (within 10–20%) to the macrocirculatory systemic hematocrit (see Fig. 9 for \( a = 0.7 \)). This is in qualitative agreement with the direct measurements of discharge hematocrit (13) that observed analogous results, albeit with a significant amount of scatter. 7) Predicted values of the branching exponent \( \xi \) fall within the range of their measured counterparts (27, 28, 33, 46).

Inverse modeling of vascular networks. The modeling framework described above treated the bifurcation parameter \( a \) as an input in order to construct a vascular network, e.g., to identify the ratio of the radii of daughter vessels \( R_{n,1}/R_{n,2} \). A problem with such “forward modeling” is that the model parameter \( a \), which determines the hematocrit partitioning at bifurcations, is harder to measure than the model output \( R_{n,1}/R_{n,2} \), which is more readily measured in morphometric studies (10, 21, 24, 32, 33, 36). The goal of “inverse modeling” is to infer the bifurcation parameter \( a \) from measurements of the parent vessel radius \( R_n \), the ratio of the daughter vessel radii \( R_{n,1}/R_{n,2} \) and the hematocrit \( H_n \) in the \( n \)th parent vessel.

This goal is facilitated by the one-to-one relationship between the bifurcation parameter \( a \) and the daughter vessel radii \( R_{n,1}/R_{n,2} \), for any given value of \( R_n \) and \( H_n \) (Fig. 10). This figure is constructed by running our forward model for multiple values of \( a \), while keeping the values of \( R_n \) and \( H_n \) fixed. The four curves in Fig. 10 correspond to \( H_n = 0.45 \) and four values of the parent vessel radius \( R_n \). We found these curves to be essentially independent of \( H_n \) over a physiologically relevant range of its values. The dependence of \( a \) on the radii ratio \( R_{n,1}/R_{n,2} \) in Fig. 10 is fitted with a second-degree polynomial

\[
a = c_{a_0} \left( \frac{R_{n,1}}{R_{n,2}} \right)^2 + c_{a_1} \frac{R_{n,1}}{R_{n,2}} + c_{a_2}
\]

The fitting coefficients \( c_{a_0}, c_{a_1}, \) and \( c_{a_2} \) vary with the parent vessel radius \( R_n \), such that

\[
c_{a_0} = -0.58, \ c_{a_1} = 1.43, \ c_{a_2} = -0.36, \ \text{for} \ R_n > 50 \mu m
\]

and

\[
c_{a_0} = -3.1 \times 10^{-4} R_n^2 + 0.03 R_n - 1.31, \ c_{a_1} = 4.9 \times 10^{-4} R_n^2 - 0.05 R_n + 2.55, \ c_{a_2} = -1.7 \times 10^{-4} R_n^2 + 0.02 R_n - 0.74, \ \text{for} \ R_n \leq 50 \mu m.
\]

Figure 10 enables one to infer a value of the bifurcation parameter \( a \) from a measurement (or the average of multiple measurements) of the ratio of daughter vessel radii \( R_{n,1}/R_{n,2} \). Then one can use our model to reconstruct the whole vascular network. Examples of such reconstructions are shown in Fig. 11. Large degrees of asymmetry imply large spreads of discharge hematocrits and vessel radii across a given generation of vessels.

DISCUSSION AND CONCLUSIONS

We proposed a new approach for simulation of vascular networks. Our method follows the foundation premise of Murray’s law (34) in postulating the existence of functional optimality of such networks. The optimality criterion adopted in our approach is the physiological cost of supplying oxygen to the tissue surrounding a blood vessel. Bifurcation asymmetry is expressed in terms of the amount of oxygen consumption associated with the respective tissue volumes being supplied by each daughter vessel. Similar to Ref. 1, our approach accounts for the non-Newtonian behavior of blood by allowing the apparent blood viscosity to vary with discharge hematocrit and vessel radius in accordance with Ref. 44.

Our approach to network reconstruction offers significant advantages over Murray’s law. Chief among them is its ability to capture the observed variability of WSS in the microcirculation. Our model predicts the sharp amplification of WSS in the smallest vessels of a network (\( R < 50.0 \mu m \)), which is consistent with in vivo observations (25, 30, 38, 42). WSS in intermediate vessels gradually increases with vessel radius, before reaching a constant value at large vessel radii (\( R > 200.0 \mu m \)). This WSS variability is absent in networks reconstructed with Murray’s law, which exhibit constant WSS throughout the vasculature.

The proposed approach captures both the asymmetric partitioning of hematocrit at vessel bifurcations and its effects on hematocrit variability in the terminal vessels of vascular networks. It provides theoretical support for the experimentally observed linear relationship between the RBC flux fraction and the blood flow fraction (3, 12, 16, 50), and for the in vivo...
observations of pronounced variability of hematocrit in capillary beds (20, 26, 39). The high-precision measurements reported in Ref. 48 suggest that this linear relationship is a result of averaging over multiple observations. To account for the scatter in the observed RBC distributions at various vessel bifurcations, one can treat the bifurcation parameter \( a \) as a random (e.g., Gaussian) variable whose realizations are assigned either to every bifurcation in the network or to all bifurcations of the same level.

Microvascular dispersion of hematocrit due to the asymmetric branching has important physiological implications. The variability of discharge hematocrit at bifurcation is cumulative, leading to the presence of capillaries with hematocrits higher than systemic, while the average capillary hematocrit is generally half of the systemic value (29). The literature refers to these high-hematocrit vessels as shunts or thoroughfare channels; they were proposed to arise from the distribution of RBCs at capillary bifurcations where they concentrate in the branch with the greater velocity (18, 23). Our analysis provides an alternative explanation: the location of these red blood cell shunts is determined by the nature of the preceding bifurcations. Therefore the microcirculation appears to be designed so that the asymmetry of bifurcations generates capillary vessels that act as RBC shunts between the arterial and venous circulatory compartments.

If the presence of shunts is indeed a consequence of properties of the capillary network, then their functional properties are difficult to control extrinsically since the mechanisms that control capillary flow within the capillary network are limited. Conversely, if the function of these shunts is regulated already at the arteriolar level, it can be controlled by the many interventions that influence arteriolar function. These findings may help in designing treatment strategies in clinical conditions of organ hypoxia and capillary ischemia wherein local blood flow is normal or even increased related to the malfunction of microvascular regulation (7, 8, 14, 15).

**APPENDIX**

The premise of Murray’s law is that a blood vessel’s volume is determined by a compromise between minimizing the vessel’s resistance to blood flow and minimizing the total volume of blood needed to serve the body’s metabolic needs (5, 49, 56). The latter condition is equivalent to positing that a blood vessel supplying a given oxygen flux \( Q_{O2} / H \) to the surrounding tissue has an optimal volume at which this compromise between minimizing resistance to flow and minimizing vessel volume is met. This implies an “optimal” vessel volume (and since the vessel is filled with blood, an optimal volume of blood) for a given tissue volume. (The existence of an optimal blood volume has been suggested in Refs. 6 and 51.) In other words, the volume of the tissue oxygenated by a vessel is related to both the vessel volume and the oxygen flux \( QH \) supplied by the vessel. We assume the linear relationships, \( V \propto \pi R^2 L \) and \( V \propto QH \), which gives

\[
\pi R^2 L \propto QH
\]  

Substituting the flow rate \( Q \) predicted by the modified Murray’s law in Eq. 7, we obtain \( R^2 L \propto R^3 H/\Lambda \). This is equivalent to \( L \Lambda/(HR) = \kappa \), where \( \kappa \) is a constant of proportionality, which leads to Eq. 18.

As an aside, we note that \( V \propto \pi R^2 L \) implies that the ratio of the vessel volume and the volume of the surrounding tissue it oxygenates
are constant throughout a vascular network. Since the oxygen flux $Q_{O_2} = QH$ is conserved from one vessel generation to the next (due to conservation of mass), the assumption that $V \propto \pi R^2L \propto QH$ for each generation of vessels implies that the vascular network is volume preserving (i.e., each successive generation of blood vessels has the same total volume as the preceding generation). This implies that any given tissue volume at any length scale (larger than the length of a capillary) will have a fixed volume fraction that is occupied by blood vessels. The assumption of a volume preserving network has previously been used in studies on scaling in biological systems (55).

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DISCLOSURES

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AUTHOR CONTRIBUTIONS

Author contributions: K.S. and D.M.T. conception and design of research; K.S. prepared figures; K.S. drafted manuscript; K.S., M.I., and D.M.T. edited manuscript; D.M.T. analyzed data.

REFERENCES

34. Murray CD. The physiological principle of minimum work applied to the angle of branching of arteries. J Gen Physiol 9: 835–841, 1926.