

Quantifying vasculature: new measures applied to arterial trees in the quail chorioallantoic membrane

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A wide variety of measures is currently in use in the morphometry of vascular systems. We introduce two additional classes of measures based on erosions and dilations of the image. Each measure has a clear biological interpretation in terms of the measured structures and their function. The measures are illustrated on images of the arterial tree of the quail chorioallantoic membrane (CAM). The new measures are correlated with widely-used measures, such as fractal dimension, but allow a clearer biological interpretation. To distinguish one CAM arterial tree from another, we propose reporting just three independent, uncorrelated numbers: (i) the fraction of tissue which is vascular (VF_0 , a pure ratio), (ii) a measure of the typical distance of the vascularized tissue to its vessels (CL, a length), and (iii) the flow capacity of the tissue (P, an area). An unusually large CL would indicate the presence of large avascular areas, a characteristic feature of tumor tissue. CL is inversely highly correlated with fractal dimension of the skeletonized image, but has a more direct biological interpretation.

Keywords: Vascular; Fractal; Chorioallantoic membrane; Angiogenesis; Cancer

1. Introduction

How does one count blood vessels? For small numbers, we begin with "one, two, three..." Over larger areas, we commonly think of some measure of vascular density, the number of vessels per unit area or volume (as in, e.g. [10]). There is a tremendous variety of measures in use (for an overview, see [9]), and many of them are not necessarily intuitive. For example, if we care primarily about vessel length, we should use a measure of *length density*. Length density is computed from a skeletonized image. If we prefer ease of measurement, we use *area density*, the fraction of a 2D image which is occupied by vessels and their lumens. Area density corresponds to volume fraction, the fraction of voxels in a 3D image which are occupied by vasculature.

Choosing the right measure of vascular density requires clarifying the purpose of the measurement. If our primary interest is in how much mass of vascular tissue is present, then volume fraction is the correct measure—but we must exclude the lumens. If we are instead interested in the total volume of the vasculature and its contents, we use volume fraction and include the lumens. However, often the feature that we are investigating is not anatomical but functional: in the case of vascularization our underlying concern is the flow. With the correctly derived measure, we can estimate flow capacity from an image.

Fractals have become a very popular metric for vascular systems e.g. [5], and software for computing the fractal dimension of an image has become fairly widespread. For many systems, the arterial tree is very well represented by a fractal. The fractal dimension is a unitless number, which can be tracked over time and compared across treatments. It is straightforward to compute, but it does come with some statistical liabilities [2].

We know what fractal dimension represents mathematically, but what it signifies biologically is less clear. We do know that fractal dimension of the chorioallantoic membrane (CAM) increases during development [4,6,11]. That mathematical fact, however, does not tell us which biological quantity is increasing over time, for which the fractal dimension is an indicator. Is it total flow? Is it flow homogeneity? We know that tumor vasculature has a higher fractal dimension than normal vasculature, and increases over time (for review, see [1]), yet a low fractal dimension can be associated with a high grade of tumor and poor patient outcome [8]. What is the biological interpretation of these observations? Figure 1 illustrates two CAM arterial trees with the same fractal dimension, yet there are obvious differences in the structures.

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Figure 1. Images from two CAM arterial trees. Scale is identical. Fractal dimension is the same (a) $D_f = 1.54 \pm 0.04$, (b) $D_f = 1.53 \pm 0.02$ yet there are major differences between the two trees.

The fractal dimension alone cannot explain the difference, which is obvious to the eye. The eye sees that there must also be significant differences in the function of these two trees.

The goal of this paper is to present a discussion of the issues of vascular quantification and to suggest additional measures of a vascular tree. We illustrate the measures using a set of images of the CAM arterial tree and discuss their biological significance.

2. Methods

Fertilized quail eggs were cultured and imaged at 72 pixels/in as in Parsons-Wingerter *et al.* [6]. Background and veins were erased by hand, arteries were filled by hand, and images were binarized, using NIH Image. Thresholds for binarization were determined by manually adjusting to retain the smallest arterial branches visible [6]. Skeletonization of binary images was done by NIH Image (figure 2). Image analysis was done on binary.tif files in Matlab.

2.1 Definitions of measures

The two most commonly used measures in the vascular morphometry literature go by almost as many names as the number of papers in which they appear. We will call them the *vascular fraction* and the *fractal dimension of the skeletonized tree*.

2.2 Vascular fraction

This is the fraction of the tissue volume (in 3D) or area (of a 2D slice) which is occupied by the vessels. It is the



Figure 2. CAM image (a) is skeletonized by removing boundary pixels until the remaining object (b) is only 1 pixel thick.

most obvious and straightforward measure of the gross amount of vasculature in a tissue. In this study, vascular fraction VF counts only the arterial tree. It is unitless, and in our sample ranges from 15-20%. It is computed for a 2D structure by counting vascular pixels in the 2D binary image, as a fraction of the total pixel number. The vascular fraction of a 3D structure can be determined either by voxel vascular fraction in a 3D stack, or extrapolated from pixel vascular fraction of a 2D slice. There are issues relating to the proper thresholding in the image analysis, and these can be exacerbated by heterogeneity in marker uptake [3,10], but vascular fraction remains conceptually the most robust measure of vascular tissue.

2.3 Fractal dimension

The fractal dimension is a unitless quantity. It can be defined as the negative of the slope of the log-log plot of the number of pixels in the vascular portion against the size of those pixels. As in [4,6], we use the box-counting method: for each of several box sizes, the image is divided into a grid, and boxes that contain some vessel are counted. The slope of the log-log graph is computed by linear regression. We define D_f as the fractal dimension of the arterial tree, and D_{fsk} as the fractal dimension of the skeletonized tree. They are different measures. Although D_{fsk} is far more commonly used in the vascular morphometry literature than is D_f , it is most often referred to simply as the fractal dimension. In this paper, we make a clear distinction between D_f and D_{fsk} , because they are correlated with different biological quantities. Our method for calculating D_f was tested on several artificial fractal images (gif) whose D_f is known from mathematical



Figure 3. CAM image (a) modified by (b) erosion, (c) dilation, and (d) pruning processes. Pruning is accomplished by erosion followed by dilation. In each case, the gauge is a disk of radius 4 pixels. Hence pruning by a gauge of 4 pixels digitally removes all structures of radius smaller than 4 pixels, while preserving the size of all thicker structures.

principles (Koch snowflake and Sierpinski triangle, square, pentagon and hexagon). Our algorithm slightly underestimates D_f for the test images by a mean of 0.05. When this factor is applied to correct the measurements, the error in D_f as compared to the true (mathematically derived) D_f is within 0.02 for all test images. We do not include this correction factor in our reported D_f or D_{fsk} for the natural (CAM) images.

2.4 Erosion and dilation

We examined a set of dilations and prunings of the vascular tree. A *dilation* of the tree is the region in the image which is within a particular distance of the vessels (figure 3). The complement of the dilation is the fraction of the tissue which is at least a particular distance from the vessels. The complement farthest from the vessels is, for example, more hypoxic. Finding the vascular complement fractions CF(r) at all possible distances r from the vasculature (figure 4) allows us to compute measures of perfusion efficiency, such as the median distance to a vessel, or to locate regions in the tissue which are unusually far from a vessel (figure 5).

A *pruning* of the vascular tree is a subset of the vascular tree which has all its branches below a certain radius trimmed (figure 6). The set of vascular fractions VF(r) of all possible pruning radii r could allow us to make measurements of the flow efficiency, for example. Pruning is accomplished by first eroding the image by a disk of radius r, then dilating by the same amount (figure 3).

2.5 Regression

The measured functions VF(r) and CF(r) can be approximated by curves, using nonlinear regression, or linear regression on transformed data. We thus determine parameters tuning these curves, which distinguish between vascular trees with different anatomical features. In any regression, it is important not simply to have small residuals (R^2 large) but also to observe no trend in the residuals.

3. Results

3.1 Fractal measures

The CAM images that we analysed (displayed in figure 11) were all very well-described by fractals D_f and D_{fsk} .



Figure 5. Significance of complement fraction. Outlined in image is the portion of the tissue which is in the upper 5% of distance to a vessel. The 95th percentile distance is $325 \,\mu$ m; hence we say that 95% of the tissue is within 1/3 mm of a vessel, and $CF(325 \,\mu$ m) = 0.05. The spatial distribution of these poorly vascularized regions is fairly uniform in the CAM; however, in tumor tissue we would expect much more patchiness.

The curves of pixel size against vessel count were highly linear for $D_f (R^2 > 0.998)$ and $D_{fsk} (R^2 > 0.97)$, though for D_{fsk} there was a slight trend in the residuals, a slight downward curvature at the coarser scales. Hence, we may reasonably assume that the CAM arterial tree is fractal rather than multifractal [13]. The range of fractal dimensions was quite small in our collection of CAM images. D_f ranged from 1.43 ± 0.02 to 1.54 ± 0.04 . D_{fsk} ranged from 1.02 ± 0.05 to 1.13 ± 0.10 . Note that both measures had an observed range in the CAM of 0.11, but D_{fsk} had a much larger standard error of each individual measurement.

3.2 Vascular fractions

The vascular fractions of the prunings, VF(r), were modelled by several curves, in particular, a linear, an exponential, a power law, a Weibull function, and a quadratic function. Surprisingly, all these function families had either poor fits or strong trends in the residuals, except for two functions.



Figure 4. Calculation of complement fraction CF(r). Original image (a) and regions of the tissue which are at least (b) 10, (c) 20, and (d) 30 pixels from any vessel. Complement fractions CF(r): (a) 78%, (b) 38%, (c) 15%, (d) 5%.



Figure 6. Prunings of a CAM arterial tree at different radii: (a) original arterial tree, (b-e) all vessels of radius below the gauges (b) 48 μ m, (c) 96 μ m, (d) 144 μ m, (e) 192 μ m, (f) 240 μ m have been digitally pruned (removed). Note that digitally-pruned vessels may not remain connected. Vascular fraction *VF*(*r*) can be determined from this process as a function of pruning radius *r*. *VF*(0 μ m) = 15.4%, *VF*(48 μ m) = 11.4%, *VF*(96 μ m) = 6.2%, *VF*(144 μ m) = 3.1%, *VF*(192 μ m) = 2.2%, *VF*(240 μ m) = 0.0%.

One was the quadratic function $VF_q(r) = VF_0[1 - (r/L_q)]^2$ where *r* and L_q are in μ m or mm and VF_0 is unitless. Regression was linear on transformed data, fixing VF_0 at the measured value, with $R^2 > 0.93$. Range of L_q was 233–581 μ m with SE 8–37 μ m.

The other function which performed well was a compound exponential, $VF_e(r) = \exp(\ln(VF_0)\exp(r/L_e))$, fitting VF_0 and L_e for each curve. The range of L_e was $158-421 \,\mu\text{m}$, with SE $8-28 \,\mu\text{m}$, and $R^2 > 0.86$. Interestingly, the fitted values of VF_0 for the compound exponential model differed from the observed (raw) VF_0 , with a correlation of only 0.72. The compound exponential fit least well at r = 0, where the data for VF(r) were unusually level (figure 7). This may be an artifact of the imaging method, or it may be an artifact of the vascular growth process itself. It is reasonable to assume that there is a minimum size of capillary below which we see no vessels at all; this would naturally tend to level off the curve of VF(r) near r = 0, as we observe.



Figure 7. Vascular fraction VF(r) after prunings to radius r. Shown are VF(r) for the images with L_e and L_q the smallest \bullet ($L_e = 158 \pm 9 \,\mu\text{m}$, $L_q = 233 \pm 8 \,\mu\text{m}$) and largest \blacksquare ($L_e = 421 \pm 16 \,\mu\text{m}$, $L_q = 581 \pm 17 \,\mu\text{m}$). Fitted curves are $VF_e(r)$ (dashed) and $VF_q(r)$ (solid).

The R^2 values are fairly low, but that is because the data for VF(r) have natural irregularity (figure 7).

3.3 Complement fractions

The complement fraction, CF(r) (see figure 8), was fitted to several test functions, including an exponential decay and several hyperbolas, but the only functional form examined which had an excellent fit to all images and had no trend in the residuals was CF(r) = $CF_0 \exp(1 - \exp(r/CL))$, where CL is a characteristic length (µm) and CF_0 is the highest value of CF (the same as $1 - VF_0$). For this function, all images had R^2 > 0.99.

Table 1 presents the measurements for a sample of 8 images. Standard errors are reported for all quantities except VF_0 and CF_0 , which are dependent solely on the pixel size, which was small enough that we assume negligible error.



Figure 8. Complement fraction CF(r) of material a radius *r* from any vessel. Shown are CF(r) for the images with the smallest and largest CL ($\blacklozenge CL = 168 \pm 4 \,\mu\text{m}$; $\blacklozenge CL = 311 \pm 4 \,\mu\text{m}$), with fitted curves. *CL* is the distance (vertical reference line) marking the 18th percentile (horizontal reference line) of non-vascular tissue's proximity to a vessel.

Table 1. Eight measures for eight images.

Image	VF_0	D_f	D_{fsk}	CL (µm)	$CF_0 = 1 - VF_0$	$L_q \ (\mu \mathrm{m})$	L_e (µm)	VF_0 (fitted)
CAM0	0.17	1.48 ± 0.03	1.07 ± 0.07	226 ± 8	0.83	233 ± 8	158 ± 9	0.21 ± 0.02
CAM1	0.16	1.47 ± 0.03	1.05 ± 0.07	237 ± 8	0.83	256 ± 25	218 ± 28	0.19 ± 0.03
CAM2	0.15	1.43 ± 0.02	1.02 ± 0.05	239 ± 5	0.85	285 ± 17	245 ± 12	0.16 ± 0.01
CAM3	0.20	1.52 ± 0.02	1.04 ± 0.06	268 ± 4	0.80	405 ± 8	284 ± 8	0.22 ± 0.02
CAM4	0.23	1.54 ± 0.04	1.13 ± 0.10	168 ± 4	0.76	416 ± 37	255 ± 15	0.21 ± 0.01
CAM5	0.19	1.49 ± 0.03	1.09 ± 0.08	204 ± 3	0.80	264 ± 13	197 ± 8	0.21 ± 0.02
CAM6	0.17	1.47 ± 0.02	1.05 ± 0.07	259 ± 2	0.82	263 ± 15	199 ± 21	0.20 ± 0.01
CAM7	0.20	1.53 ± 0.02	1.03 ± 0.06	311 ± 4	0.80	581 ± 17	421 ± 16	0.22 ± 0.02

 VF_0 , vascular fraction, and CF_0 , complement fraction, assumed to have no error. VF_0 (fitted) is usually larger than VF_0 from the raw image, because of a lower limit of capillary size. D_f , fractal dimension, and D_{fsk} , fractal dimension of the skeletonized image. CL, characteristic length, derived from shape of CF(r). L_q (quadratic) and L_e (exponential), are lengths derived from shape of VF(r). For definitions, see Appendix.

3.4 Derived quantities

Once we know the gauge-dependent vascular fraction VF(r), the measure can be used to calculate derived measures for area density, length density and volume density at the different gauges. In the case of planar vasculature (such as in the CAM), the area, length and volume of vessels in a particular range of gauges r_1 to r_2 per unit area of tissue are given in table 2.

Hence, for example, the area density of vessels at all gauges is $-\int_0^{\infty} (d/dr)(VF(r))dr = VF(0) - VF(\infty) = VF_0$, the vascular fraction. Also, we see that for planar vasculature, area density is equivalent to vascular fraction, since $AD(r, \infty) = -\int_r^{\infty} (d/dr)(VF(r))dr = VF(r)$.

Note that it is not possible to calculate the length density of vessels at the smallest gauges from the formula, because the formula for length density blows up at a radius $r_I = 0$. The most widespread method of measuring length density, the grid intersection method, avoids this difficulty by missing many of the vessels smaller than the grid size. Real vessels, however, do have a minimum size, so the blowup of the length density at small radii presents little practical difficulty.

The volume density is well-behaved, and can be used as another measure of a vascular tree, if we are particularly concerned, for instance, with volumes of planar vascular trees, or with quantities closely related to volumes. Figure 9 illustrates the volume density distribution (integrand of volume density) for a CAM image whose VF(r) was fitted by the quadratic and exponential functions $VF_q(r)$ and $VF_e(r)$.

Similarly, we can use VF(r) measured from an image to derive any quantity which may depend on the radius of a vessel. For example, mean flow rate depends on the 4th power of the vessel radius, if Poiseuille's law applies.

Table 2.

Measure	units	formula
Length density $LD(r_1, r_2)$	length/area	$-\int_{r_1}^{r_2} \frac{1}{2r} \frac{\mathrm{d}}{\mathrm{d}r} (VF(r)) \mathrm{d}r$
Area density $AD(r_1, r_2)$	area/area	$-\int_{r_1}^{r_2} \frac{d}{dr} (VF(r)) dr = VF(r_1) - VF(r_2)$
Volume density $VD(r_1, r_2)$	volume/area	$-\int_{r_1}^{r_2} \frac{\pi r}{2} \frac{\mathrm{d}}{\mathrm{d}r} (VF(r)) \mathrm{d}r$

We can estimate the permeability coefficient *P* as $P = -\int_0^{\infty} r^2 (d/dr)(VF(r))dr$ such that the total flow rate through a vascular bed will be proportional to the product of *P* and the ratio of pressure drop to viscosity [4]. There will be additional factors determining the actual flow rate, such as tortuosity and elasticity of the vessels, but our permeability coefficient *P* gives a fair quantitative comparison between the flows expected in geometrically similar structures. *P* has area units (length⁴ per unit area).

In figure 10 we see how the permeability is shared among vessels of different radii, and we see that the estimation of the permeability's dependence on radius also depends on which function is chosen to approximate VF(r). The functions $VF_q(r)$ and $VF_e(r)$ are not fundamental to any measures, but are simply convenient ways of smoothing the VF(r) data, which naturally have some roughness. Because of the simple form of $VF_q(r)$, we can estimate P as the simple expression $(VF_0/6)L_q^2$. The numerical estimate of P by use of $VF_e(r)$ correlates very well (r = 0.98) with the estimate using $VF_q(r)$. Table 3 is a comparison of the permeability coefficients. Actual flow rate through a vascular bed would depend on vessel tortuosity and elasticity, the viscosity of the blood, and the pressure drop.



Figure 9. Planar volume density distribution (unitless) at a given gauge r (µm), shown calculated from the fitted functions $VF_q(r)$ (solid curve) and $VF_e(r)$ (dashed curve) for the same image. Note that the greatest contribution to volume is at an intermediate gauge, and this gauge is somewhat different for the two functions. The total volume per unit area for this arterial tree is estimated differently by the exponential (22.5 µm³/µm²) and quadratic(27.8 µm³/µm²) functions.



Figure 10. Distribution of flow capacity (arbitrary units) among vessels of gauge r (µm), as estimated by two different functions VF(r) for a particular CAM image. The quadratic function $VF_q(r)$ (solid curve) estimates a higher overall flow rate than the exponential function $VF_e(r)$ (dashed curve). Flow distribution units are arbitrary because actual flow rate depends on viscosity and pressure drop.

3.5 Correlations

Some of the measures we have examined for the CAM arterial trees are highly correlated, and others are uncorrelated (table 4). Interestingly, the two fractal dimensions D_f and D_{fsk} are only weakly correlated with each other. Fractal dimension D_f is strongly correlated with VF_0 , the fraction of the image which is vascular, and in fact the measured VF_0 correlates better with D_f than it does with the fitted VF_0 . On the other hand, the skeletonized fractal dimension D_{fsk} is strongly negatively correlated with CL. In turn, CL is uncorrelated with D_f and with VF_0 . Thus we could use VF_0 as a surrogate measure for D_{f} , and CL as a surrogate measure for D_{fsk} . L_e and L_q are highly correlated, as is expected for different measures of the same feature, and they in turn are strongly correlated with the permeability coefficients P_e and P_q . A principal component analysis (PCA) was performed, but the results do not provide new insight.

To distinguish one CAM arterial tree from another, in general it should suffice to report just three numbers. The measures should not be correlated with each other, or they will be presenting redundant information. A useful trio of independent, mostly uncorrelated measures could be VF_0 , CL, and P. They represent, respectively, the fraction of tissue, which is vascular (VF_0 , a pure ratio), a measure of the distance of the vascularized tissue to its vessels (CL, a

Table 3. Estimates of permeability coefficient $P(\mu m^2)$.

Image	P_q	P_e
CAM0	1900	1562
CAM1	2075	2471
CAM2	2166	2291
CAM3	6014	5509
CAM4	6057	4069
CAM5	2439	2428
CAM6	2306	2263
CAM7	12377	12106

P is derived from vascular fraction VF(r) of vessels of different radii, $P = -\int_0^\infty r^2 (d/dr)(VF(r))dr$. P_q calculated from $VF_q(r) = VF_0(1 - r/L_q)^2$, P_e calculated from $VF_e(r) = \exp(\ln (VF_0)\exp(r/L_e))$. True flow rate depends on several other factors, such as pressure drop and tortuosity.

length), and the flow capacity of the tissue (P, an area). The three measures, along with fractal dimension D_{f} , are shown for several images in figure 11.

4. Discussion

The goal of this paper is to present a comparison of different common and uncommon measures of a vascular tree, with an eye to increasing the amount of biological insight gained from the use of these measures.

In particular, we find that the common fractal dimension D_f has some common problems. First, the fractal dimension which is most widely reported as a vascular measure is actually of the skeletonized image, not the raw image, yet typically this distinction is not recognized. Second, it is not clear whether any groups reporting fractal dimensions of natural images have calibrated their algorithms on mathematical images of known D_{f} . When we did so with the most commonly used algorithm (gridbased box counting), we found a consistent underestimate of D_{f} , possibly due to prior compression of the fractal images by the authors who had generated them. We strongly recommend that researchers reporting fractal dimension of their natural images (a) indicate whether the images have been skeletonized, and (b) calibrate their algorithms on mathematical fractals.

We find that the abstract fractal dimension D_f is really a surrogate measure for the vascular fraction VF_0 . Since VF_0 is much more straightforward to measure, it would seem

Table 4. Correlations among measures reported in tables 1 and 2.

	VF_0 meas	VF_0 fit	CL	L_q	L_e	D_{f}	D_{fsk}	P_q	P_{e}
VF_0 meas	1.00								
VF_0 fit	0.72	1.00							
CL	-0.24	0.14	1.00						
L_a	0.65	0.49	0.46	1.00					
L_{e}^{q}	0.42	0.29	0.62	0.95	1.00				
$\tilde{D_f}$	0.94	0.86	0.00	0.74	0.53	1.00			
D_{fsk}	0.64	0.36	-0.84	-0.09	- 0.34	0.48	1.00		
P_a^{jsk}	0.62	0.54	0.52	0.99	0.95	0.74	-0.12	1.00	
$P_e^{''}$	0.47	0.49	0.65	0.95	0.97	0.63	-0.27	0.98	1.00

Quantities which use different methods to measure the same physical feature (such as L_q and L_e) should be highly correlated. Quantities which measure very different physical features should have low correlations, unless there is some biological reason for a high correlation.



Figure 11. Eight images of CAM arterial trees. Trees on the left are in an intermediate range for the four measures shown. Trees on the right are at either a maximum or a minimum of one or more of the displayed measures (boldface).

superfluous to use D_f , which has no intrinsic biological meaning. More commonly used than D_f is D_{fsk} , the fractal dimension of the skeletonized image. We have seen that (at least for the CAM) D_{fsk} is a surrogate measure for CL, a

length which is characteristic of the complement fraction *CF*, the space between the vessels. What is the *CL*? To be precise, it is the negative reciprocal of the initial slope of the curve of CF(r); $\exp(1 - \exp(1)) = 18\%$ of the nonvascular tissue, or 13-15% of the whole tissue, is farther from a vessel than that distance (as illustrated in figure 8). Therefore, the CL is a measure of how close most of the interstitium is to the vessels, and it is a surrogate measure for D_{fsk} (conversely, D_{fsk} is a surrogate measure for how close most of the interstitium is to the vessels). The higher the fractal dimension D_{fsk} , the lower the characteristic length, and the closer most of the tissue is to the vessels. When we observe that the fractal dimension D_{fsk} is increasing over time during development of the CAM, we can interpret it as the average piece of tissue getting closer to a vessel over time, the vasculature increasing its geometric efficiency.

A large number of derived measures based on the complement fraction CF(r) are possible, and are straightforward to implement. For example, it is well known that tumor tissue has large avascular areas; this characteristic would appear quantitatively either as a large CL or small D_{fsk} , or visually as the easily computed regions which are more than a particular distance from a vessel (figure 5).

Other authors [6,11] have found high correlations between D_{fsk} and what is called either ρ_{ν} , vascular density, or S_i , vessel length. Note that ρ_{ν} is not the same as VF_0 ; it is the vascular fraction of the skeletonized image, hence is more correctly a measure of length density, or equivalent to S_i per unit area. We may conclude that for CAM arterial trees, D_{fsk} is a measure of both length density and of interstitial proximity, but is uncorrelated with any of the other measures we have examined.

The vascular measures based on digital "pruning" of the vascular tree allow us to estimate the effectiveness of functioning of the vascular tree with various derived measures. For example, we were able to use the measured functions VF(r) to estimate the permeability coefficient P of the vascular beds. If our primary biological interest is in studying the efficiency of a vascular tree and its dependence on our experimental conditions, the most important measurement of the tree may be P. If our interest is in determining whether a particular growth factor is affecting linear versus radial growth of the vasculature, the length density $LD(r_1, r_2)$ gives us a measure of the linear density of vessels in that particular range of gauges. LD may be easier to implement than measures of branch generations [7,12].

We proposed a trio of independent, uncorrelated measures: VF_{o} , CL, and P. They represent, respectively, the fraction of tissue which is vascular (VF_o , a pure ratio), the distance of the vascularized tissue to its vessels (CL, a length), and the flow capacity of the tissue (P, an area). These units may be more intuitive than the fractal dimension, and two of the measures are highly correlated with fractal dimensions. Our proposed trio can replace fractal dimension as a measure of a vascular tree, and each

Appendix: Measures discussed in this paper

Symbol	Name	Units	Definition
D_f	fractal dimension	_	-slope of log-log plot of box counts at each gauge
D_{fsk}	fractal dimension of skeletonized image	-	-slope of log-log plot of box counts at each gauge
r	gauge	length	pruning radius or dilation radius
VF(r)	volume fraction	-	fraction of image consisting of vessels larger than a given gauge <i>r</i>
CF(r)	complement fraction	-	fraction of image which is farther from any vessel than distance r
VF_0	volume fraction	_	fraction of image which is vascular tissue
CF_0	complement fraction	_	fraction of image which is not vascular tissue
CL	characteristic length of complement fraction	length	best-fitting parameter in $CF(r) = CF_0 \exp\left(1 - \exp\left(\frac{r}{CT}\right)\right)$
$VF_q(r)$	quadratic fit of volume fraction	-	$VF_q(r) = VF_0 \left(1 - \frac{r}{L_q}\right)^2$
$VF_e(r)$	compound exponential fit of volume fraction	-	$VF_e(r) = \exp\left(\ln\left(VF_0\right)\exp\left(\frac{r}{L_e}\right)\right)$
La	characteristic length of vascular fraction	length	see definition for $VF_{a}(r)$; best fit to data
L_e^q	characteristic length of vascular fraction	length	see definition for $VF_e(r)$; best fit to data
$LD(r_1,r_2)$	length density	length/area	$-\int_{r_1}^{r_2} \frac{1}{2r} \frac{\mathrm{d}}{\mathrm{d}r} (VF(r)) \mathrm{d}r$
$AD(r_1,r_2)$	area density	area/area	$-\int_{r_1}^{r_2} \frac{d}{dr} (VF(r)) dr = VF(r_1) - VF(r_2)$
$VD(r_1, r_2)$	volume density	volume/area	$-\int_{r_1}^{r_2} \frac{\pi r}{2} \frac{\mathrm{d}}{\mathrm{d}r} (VF(r)) \mathrm{d}r$
Р	permeability coefficient	area	$P = -\int_0^\infty r^2 \frac{\mathrm{d}}{\mathrm{d}r} (VF(r)) \mathrm{d}r$
P_q	permeability coefficient (quadratic method)	area	$P_q = -\int_0^\infty r^2 \frac{\mathrm{d}}{\mathrm{d}r} (VF_q(r)) \mathrm{d}r = \frac{VF_0}{6} L_q^2$
P_e	permeability coefficient (compound exponential method)	area	$P_e = -\int_0^\infty r^2 \frac{\mathrm{d}}{\mathrm{d}r} (VF_e(r)) \mathrm{d}r$

of our three measures has a straightforward biological interpretation.

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