Optimal Supply Networks II: Blood, Water, and Truthicide

Complex Networks | @networksvox CSYS/MATH 303, Spring, 2016

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Metabolism and Truthicide

fractions

Measuring allometric exponents

River networks

Earlier theories

Geometric Real networks

Conclusion







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Outline

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Stories—The Fraction Assassin:



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Law and Order, Special Science Edition: Truthicide Department

"In the scientific integrity system known as peer review, the people are represented by two highly overlapping yet equally important groups: the independent scientists who review papers and the scientists who punish those who publish garbage. This is one of their stories."

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Fundamental biological and ecological constraint:

 $P = c M^{\alpha}$

P =basal metabolic rate M =organismal body mass







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Prefactor *c* depends on body plan and body temperature:

Birds	39−41 ° <i>C</i>
Eutherian Mammals	$36\text{-}38^{\circ}C$
Marsupials	34- 36 ° <i>C</i>
Monotremes	30 −31 ° <i>C</i>





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 $\alpha = 2/3$ because ...

Dimensional analysis suggests an energy balance surface law:

$$P \propto S \propto V^{2/3} \propto M^{2/3}$$

- Assumes isometric scaling (not quite the spherical cow).
- Lognormal fluctuations: Gaussian fluctuations in $\log P$ around $\log cM^{\alpha}$.
- Stefan-Boltzmann law for radiated energy:

$$\frac{\mathrm{d}E}{\mathrm{d}t} = \sigma \varepsilon S T^4 \propto S$$

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The prevailing belief of the Church of Quarterology:

$$\alpha = 3/4$$

 $P \propto M^{3/4}$

Huh?

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The prevailing belief of the Church of Quarterology:

Most obvious concern:

$$3/4 - 2/3 = 1/12$$

- An exponent higher than 2/3 points suggests a fundamental inefficiency in biology.
- Organisms must somehow be running 'hotter' than they need to balance heat loss.

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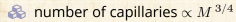


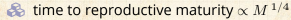




Related putative scalings:

Wait! There's more!:





 \Leftrightarrow heart rate $\propto M^{-1/4}$

 \ref{sphi} cross-sectional area of aorta $\propto M^{3/4}$

 \Leftrightarrow population density $\propto M^{-3/4}$

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The great 'law' of heartbeats:

Assuming:

 $\red {\mathbb A}$ Average lifespan $\propto M^{eta}$

 $\red {\mathbb A}$ Average heart rate $\propto M^{-\beta}$

 $\mbox{\&}$ Irrelevant but perhaps $\beta = 1/4$.

Then:

Average number of heart beats in a lifespan \simeq (Average lifespan) \times (Average heart rate) $\propto M^{\beta-\beta}$

 $\propto M^0$

Number of heartbeats per life time is independent of organism size!

& ≈ 1.5 billion....

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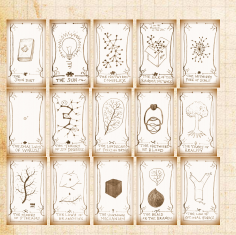
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A theory is born:

1840's: Sarrus and Rameaux [37] first suggested $\alpha = 2/3$.



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A theory grows:

1883: Rubner [35] found $\alpha \simeq 2/3$.



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Theory meets a different 'truth':

1930's: Brody, Benedict study mammals. [6] Found $\alpha \simeq 0.73$ (standard).



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Our hero faces a shadowy cabal:



- 3 1932: Kleiber analyzed 13 mammals. [22]
- \implies Found $\alpha = 0.76$ and suggested $\alpha = 3/4$.
- Scaling law of Metabolism became known as Kleiber's Law (2011 Wikipedia entry is embarrassing).
- 1961 book: "The Fire of Life. An Introduction to Animal Energetics". [23]

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When a cult becomes a religion:

1950/1960: Hemmingsen [19, 20] Extension to unicellular organisms. $\alpha = 3/4$ assumed true.



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Quarterology spreads throughout the land:

The Cabal assassinates 2/3-scaling:

1964: Troon, Scotland.

3rd Symposium on Energy Metabolism.

 $\alpha = 3/4$ made official ...

...29 to zip.



But the Cabal slipped up by publishing the conference proceedings ...

"Energy Metabolism; Proceedings of the 3rd symposium held at Troon, Scotland, May 1964," Ed. Sir Kenneth Blaxter [4] COcoNuTS

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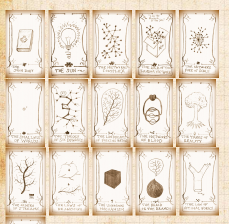
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So many questions ...

- Did the truth kill a theory? Or did a theory kill the truth?
- Or was the truth killed by just a lone, lowly hypothesis?
- Does this go all the way to the top? To the National Academies of Science?
- Is 2/3-scaling really dead?
- Could 2/3-scaling have faked its own death?
- What kind of people would vote on scientific facts?

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Modern Quarterology, Post Truthicide

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3/4 is held by many to be the one true exponent.



In the Beat of a Heart: Life, Energy, and the Unity of Nature—by John Whitfield Metabolism and Truthicide

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But: much controversy ...



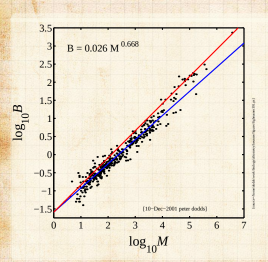
See 'Re-examination of the "3/4-law" of metabolism' by the Heretical Unbelievers Dodds, Rothman, and Weitz [13], and ensuing madness...







Some data on metabolic rates



Heusner's data $(1991)^{[21]}$



391 Mammals



Blue line: 2/3



Arr (B=P)

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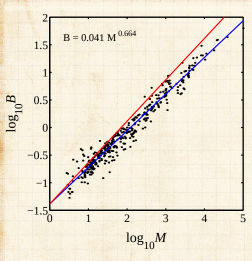






Some data on metabolic rates





Bennett and Harvey's data $(1987)^{[3]}$



Blue line: 2/3

red line: 3/4.

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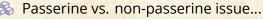
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Important:

- Ordinary Least Squares (OLS) Linear regression is only appropriate for analyzing a dataset $\{(x_i,y_i)\}$ when we know the x_i are measured without error.
- \Leftrightarrow Here we assume that measurements of mass M have less error than measurements of metabolic rate B.
- Linear regression assumes Gaussian errors.

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Measuring exponents

More on regression:

If (a) we don't know what the errors of either variable are,

or (b) no variable can be considered independent, then we need to use Standardized Major Axis Linear Regression. [36, 34] (aka Reduced Major Axis = RMA.)

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For Standardized Major Axis Linear Regression:

 $\mathsf{slope}_{\mathsf{SMA}} = \frac{\mathsf{standard} \ \mathsf{deviation} \ \mathsf{of} \ y \ \mathsf{data}}{\mathsf{standard} \ \mathsf{deviation} \ \mathsf{of} \ x \ \mathsf{data}}$

- Very simple!
- Minimization of sum of areas of triangles induced by vertical and horizontal residuals with best fit line.
- The only linear regression that is Scale invariant
 ∴.
- Attributed to Nobel Laureate economist Paul Samuelson , [36] but discovered independently by others.
- #somuchwin

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Relationship to ordinary least squares regression is simple:

$$\begin{aligned} \mathsf{slope}_{\mathsf{SMA}} &= r^{-1} \times \mathsf{slope}_{\mathsf{OLS}\,y\,\mathsf{on}\,x} \\ &= r \times \mathsf{slope}_{\mathsf{OLS}\,x\,\mathsf{on}\,y} \end{aligned}$$

where r = standard correlation coefficient:

$$r = \frac{\sum_{i=1}^{n}(x_i - \bar{x})(y_i - \bar{y})}{\sqrt{\sum_{i=1}^{n}(x_i - \bar{x})^2}\sqrt{\sum_{i=1}^{n}(y_i - \bar{y})^2}}$$

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Heusner's data, 1991 (391 Mammals)

range of M	N	\hat{lpha}
	467	
$\leq 0.1 \text{ kg}$	167	0.678 ± 0.038
$\leq 1~{\sf kg}$	276	0.662 ± 0.032
$\leq 10~\mathrm{kg}$	357	0.668 ± 0.019
$\leq 25~{ m kg}$	366	0.669 ± 0.018
$\leq 35~{ m kg}$	371	0.675 ± 0.018
$\leq 350 \text{ kg}$	389	0.706 ± 0.016
_ 330 N 8		0.00
$\leq 3670~\mathrm{kg}$	391	0.710 ± 0.021

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Bennett and Harvey, 1987 (398 birds)

$M_{\sf max}$	N	\hat{lpha}
≤ 0.032	162	0.636 ± 0.103
≤ 0.032	102	0.030 ± 0.103
≤ 0.1	236	0.602 ± 0.060
≤ 0.32	290	0.607 ± 0.039
≤ 1	334	0.652 ± 0.030
≤ 3.2	371	0.655 ± 0.023
≤ 10	391	0.664 ± 0.020
≤ 32	396	0.665 ± 0.019
≤ 100	398	0.664 ± 0.019

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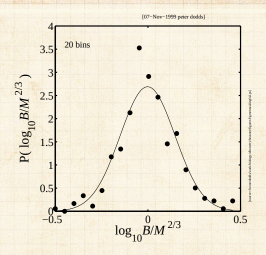
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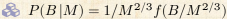
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Fluctuations—Things look normal...





Use a Kolmogorov-Smirnov test.

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Test to see if α' is consistent with our data $\{(M_i, B_i)\}$:

$$H_0: \alpha = \alpha'$$
 and $H_1: \alpha \neq \alpha'$.

- Assume each \mathbf{B}_i (now a random variable) is normally distributed about $\alpha'\log_{10}M_i+\log_{10}c$.
- Follows that the measured α for one realization obeys a t distribution with N-2 degrees of freedom.
- \Leftrightarrow Calculate a p-value: probability that the measured α is as least as different to our hypothesized α' as we observe.
- See, for example, DeGroot and Scherish, "Probability and Statistics." [10]

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Revisiting the past—mammals

Full mass range:

	N	\hat{lpha}	$p_{2/3}$	$p_{3/4}$	
Kleiber	13	0.738	$< 10^{-6}$	0.11	
Brody	35	0.718	$< 10^{-4}$	$< 10^{-2}$	
Heusner	391	0.710	$< 10^{-6}$	$< 10^{-5}$	
Bennett and Harvey	398	0.664	0.69	$< 10^{-15}$	

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Revisiting the past—mammals

 $M \leq 10 \text{ kg}$:

	N	\hat{lpha}	$p_{2/3}$	$p_{3/4}$	
Kleiber	5	0.667	0.99	0.088	
Brody	26	0.709	$< 10^{-3}$	$< 10^{-3}$	
Heusner	357	0.668	0.91	$< 10^{-15}$	

 $M \ge 10$ kg:

	N	\hat{lpha}	$p_{2/3}$	$p_{3/4}$	
Kleiber	8	0.754	$< 10^{-4}$	0.66	
			0		
Brody	9	0.760	$< 10^{-3}$	0.56	
			10		
Heusner	34	0.877	$< 10^{-12}$	$< 10^{-7}$	

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- 1. Presume an exponent of your choice: 2/3 or 3/4.
- 2. Fit the prefactor ($\log_{10} c$) and then examine the residuals:

$$r_i = \log_{10} B_i - (\alpha' \log_{10} M_i - \log_{10} c).$$

- 3. H_0 : residuals are uncorrelated H_1 : residuals are correlated.
- 4. Measure the correlations in the residuals and compute a *p*-value.

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We use the spiffing Spearman Rank-Order Correlation Coefficient

Basic idea:

 Given $\{(x_i,y_i)\}$, rank the $\{x_i\}$ and $\{y_i\}$ separately from smallest to largest. Call these ranks R_i and S_i .

 $\red{\$}$ Now calculate correlation coefficient for ranks, r_s :

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$$r_s = \frac{\sum_{i=1}^n (R_i - \bar{R})(S_i - \bar{S})}{\sqrt{\sum_{i=1}^n (R_i - \bar{R})^2} \sqrt{\sum_{i=1}^n (S_i - \bar{S})^2}}$$

 $\ref{eq:special}$ Perfect correlation: x_i 's and y_i 's both increase monotonically.

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We assume all rank orderings are equally likely:

- R_s is distributed according to a Student's t-distribution $\mathcal L$ with N-2 degrees of freedom.
- Excellent feature: Non-parametric—real distribution of x's and y's doesn't matter.
- Bonus: works for non-linear monotonic relationships as well.
- See Numerical Recipes in C/Fortran which contains many good things. [32]

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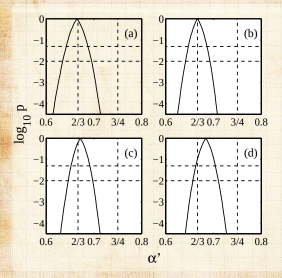
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Analysis of residuals—mammals



- (a) M < 3.2 kg,
- (b) M < 10 kg,
- (c) M < 32 kg,
- (d) all mammals.

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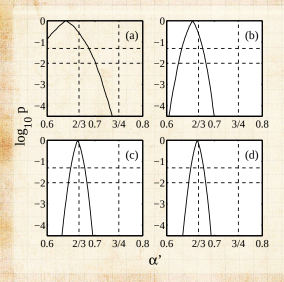
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Analysis of residuals—birds



- (a) M < 0.1 kg
- (b) $M < 1 \, \text{kg}$
- (c) M < 10 kg
- (d) all birds.

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Clauset, Shalizi, Newman: "Power-law distributions in empirical data" [9] SIAM Review, 2009.

See Clauset's page on measuring power law exponents (code, other goodies).

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- So: The exponent $\alpha = 2/3$ works for all birds and mammals up to 10–30 kg
- For mammals > 10-30 kg, maybe we have a new scaling regime
- Possible connection?: Economos (1983)—limb length break in scaling around 20 kg [14]
- But see later: non-isometric growth leads to lower metabolic scaling. Oops.

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Now we're really confused (empirically):

- Nhite and Seymour, 2005: unhappy with large herbivore measurements [47]. Pro 2/3: Find $\alpha \simeq 0.686 \pm 0.014$.
- Glazier, BioScience (2006) [17]: "The 3/4-Power Law Is Not Universal: Evolution of Isometric, Ontogenetic Metabolic Scaling in Pelagic Animals."
- S Glazier, Biol. Rev. (2005) [16]: "Beyond the 3/4-power law: variation in the intra- and interspecific scaling of metabolic rate in animals."
- Savage et al., PLoS Biology (2008) [38] "Sizing up allometric scaling theory" Pro 3/4: problems claimed to be finite-size scaling.

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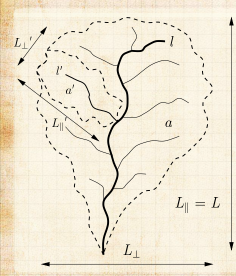
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Somehow, optimal river networks are connected:





🚵 a = drainage basin area



♣ length of longest (main) stream



& $L=L_{\parallel}$ = longitudinal length of basin COcoNuTS

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Mysterious allometric scaling in river networks

3 1957: J. T. Hack [18] "Studies of Longitudinal Stream Profiles in Virginia and Maryland"

 $\ell \sim a^h$

 $h \sim 0.6$

Anomalous scaling: we would expect h = 1/2...

Subsequent studies: $0.5 \lesssim h \lesssim 0.6$

Another quest to find universality/god...

A catch: studies done on small scales.

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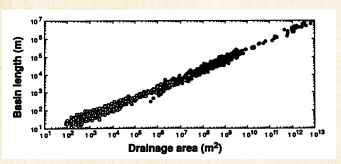


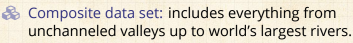


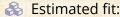


Large-scale networks:

(1992) Montgomery and Dietrich [30]:







 $L \simeq 1.78a^{0.49}$

Mixture of basin and main stream lengths.

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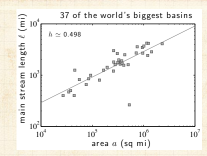








World's largest rivers only:



Data from Leopold (1994) [26, 12]



 \Leftrightarrow Estimate of Hack exponent: $h = 0.50 \pm 0.06$

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Building on the surface area idea:

- & McMahon (70's, 80's): Elastic Similarity [27, 29]
- Idea is that organismal shapes scale allometrically with 1/4 powers (like trees...)
- Disastrously, cites Hemmingsen [20] for surface area data.
- Appears to be true for ungulate legs ... [28]
- Metabolism and shape never properly connected.

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"Size and shape in biology"

T. McMahon, Science, **179**, 1201–1204, 1973. [27]

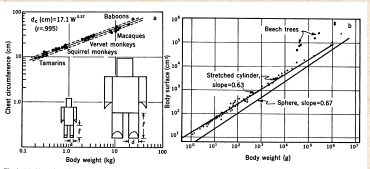


Fig. 3. (a) Chest circumference, d., plotted against body weight, W, for five species of primates. The broken lines represent the standard error in this least-squares fit [adapted from (21)]. The model proposed here, whereby each length, I, increases as the ½ power of diameter, d., is illustrated for two weights differing by a factor of 16. (b) Body surface area plotted against weight for vertebrates. The animal data are reasonably well fitted by the stretched cylinder model [adapted from (3)].

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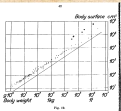
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The relation of body surface to body weight in vertebrates. The points surrounded by a circle represent beech trees. The nuthorities of the data are in approximate order of hody sizes of organisms: Fishes (Yinca, Esax, Salaro, Pleuronectes Resus, Annuilla, Creniinbrus, Lu-Srue: 9.84 p-2 kg), Jan Bokrius (unpublished). Frogs (3.5-32 g), lisards (3-13 g), Fay, 1914, p. 191. Ross excalente (25 and 50 g), Krosss, 1904, p. 404. Lizards (Locerta muralis and piridis, August fragilis: 5-26 g) and Ringed Snake (47-100 g), Isano, 1911, pp. 7-8. Teuch (Times: 211 g), frog (44 g), rabbit (3.6 kg), Very, 1930, no. 239, 244, 245, Dogs (7 and 30 kg), pigs, (3 and 100 kg), horses (175 and 900 kg), monkeys (2.5 and 5.5 kg), man (6 and 65 kg), Bacer, Convers and Marrietws, 1928, pp. 8, 30, 33 and 51. Snakes (rattle-snake, small and large python, box: 8.5-32 kg), Busineers, 1932, p. 146. Rate (20 and 250 g), cattle (20 and 460 kg), Becov, 1945, pp. 360, 361. Giant shark (2.75 I), rhinoseres (1 I), Hassensusen, 1950, pp. 30 and 43. Beech trees without leaves and roots (30 kg-1.3 t), Maller, Nittlsess and Mileage, 1954, tables 2-4 on pp. 277-281.

assuming a specific gravity of 1.0. Naturally, the inclination of this line corresponds to a proportionality power of 0.87. Of the unicellular organisms represented in fig. 1 not a few are subtrical in shape (the bacterium Sorcella, Saccharomyces, marine ergs); and most of the others have surfaces exceeding those of subserve of courst volume by rarely more than what corresponds to 0.1 decade in the log, ordinate system (Photobacterium abouthorsecray: 12 %, i.e. 0.05 decade, Escherichia coli: 34 %, i. c. 0.13 decade, the ciliates Colpidium and Paramaccium: 19-

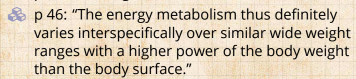
22 %, i.e. about 0.08-0.09 decade; calculated on the basis of data of Pürren, 1924, table 7 on p. 108, and Hanvey, 1928, table 1). Similar figures probably hold for other ciliates. Only the fluorilates represented (Trapanosomidae, Astesia Mebril) and certain amorboe are likely to deviate by higher figures. The surface values of the unicellular organisms represented in fig. 1 will, therefore, fall either on, or in most other cases less than 0.1 decade above, a line representing the relation between surface

It will be seen from fig. 10 that the points representing the hody surfaces of the metazoic minute in question are grouped parallel to the sphere line; that is, also corresponding to a proportionality power of 0.67. An average line through the points would fall about 0.30 logarithmic decade above the sphere line. meaning that on the average the body surface is roughly 2 (antilog, 0,30) times higher in the animals under study than in soberry of must weight or volume. In organisms of extreme shapes as the python (1045 g) and the beech trees (especially marked in fig. 3) the surface is about 3 and 10 times, respectively, greater than in a solvere of equal weight and volume. These facts agree well with the values 9-11.8 for the constant k in the

body surface in $em^2 = k \cdot body$ weight^{0,6}

as tabularized by Busumer (1938, p. 175) for various birds and mammals weighing 8 g-14 kg; because this is about double the value of k for sphere surface (4.83). The value of k (13.95) found by Kabses (1940) for Azcariz is 2.9 times 4.83, and this corresponds well with the above mentioned figure 3 for the much larger python of similar shape.

Hemmingsen's "fit" is for a 2/3 power, notes possible 10 kg transition. [?]



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Earlier theories (1977):

Building on the surface area idea...

Blum (1977)^[5] speculates on four-dimensional biology:

$$P \propto M^{(d-1)/d}$$

$$d = 3$$
 gives $\alpha = 2/3$

$$\Leftrightarrow d=4 \text{ gives } \alpha=3/4$$

So we need another dimension...

Obviously, a bit silly... [39]

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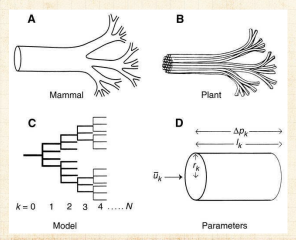




Nutrient delivering networks:

1960's: Rashevsky considers blood networks and finds a 2/3 scaling.

3/4 scaling. 46 use a network story to find



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Nutrient delivering networks:

West et al.'s assumptions:

- 1. hierarchical network
- 2. capillaries (delivery units) invariant
- 3. network impedance is minimized via evolution

Claims:



 $P \propto M^{3/4}$

networks are fractal

quarter powers everywhere

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Impedance measures:

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Poiseuille flow (outer branches):

$$Z = \frac{8\mu}{\pi} \sum_{k=0}^{N} \frac{\ell_k}{r_k^4 N_k}$$

Pulsatile flow (main branches):

$$Z \propto \sum_{k=0}^N \frac{h_k^{1/2}}{r_k^{5/2} N_k}$$

- Wheel out Lagrange multipliers ...
- Arr Poiseuille gives $P \propto M^1$ with a logarithmic correction.
- Pulsatile calculation explodes into flames.

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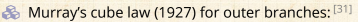




Actually, model shows:

- $Rrightarrow P \propto M^{3/4}$ does not follow for pulsatile flow
- networks are not necessarily fractal.

Do find:



$$r_0^3 = r_1^3 + r_2^3$$

- Impedance is distributed evenly.
- Can still assume networks are fractal.

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Connecting network structure to α

1. Ratios of network parameters:

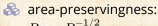
$$R_n = \frac{n_{k+1}}{n_k}, \ R_\ell = \frac{\ell_{k+1}}{\ell_k}, \ R_r = \frac{r_{k+1}}{r_k}$$

2. Number of capillaries $\propto P \propto M^{\alpha}$.

$$\Rightarrow \boxed{\alpha = -\frac{\ln R_n}{\ln R_r^2 R_\ell}}$$

(also problematic due to prefactor issues)

Obliviously soldiering on, we could assert:



$$R_n = R_n^{-1/2}$$

 $\Rightarrow \alpha = 3/4$

 \Re space-fillingness: $R_{\ell} = R_n^{-1/3}$

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Data from real networks:

Network	R_n	R_r	R_ℓ	$-rac{\ln R_r}{\ln R_n}$	$-rac{\ln R_\ell}{\ln R_n}$	α
West <i>et al.</i>	-	-	-	1/2	1/3	3/4
rat (PAT)	2.76	1.58	1.60	0.45	0.46	0.73
cat (PAT) (Turcotte <i>et al.</i> ^[43])	3.67	1.71	1.78	0.41	0.44	0.79
dog (PAT)	3.69	1.67	1.52	0.39	0.32	0.90
pig (LCX) pig (RCA) pig (LAD)	3.57 3.50 3.51	1.89 1.81 1.84	2.20 2.12 2.02	0.50 0.47 0.49	0.62 0.60 0.56	0.62 0.65 0.65
human (PAT) human (PAT)	3.03	1.60	1.49 1.49	0.42	0.36 0.33	0.83 0.94

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Some people understand it's truly a disaster:



"Power, Sex, Suicide: Mitochondria and the Meaning of Life" by Nick Lane (2005). [25]

"As so often happens in science, the apparently solid foundations of a field turned to rubble on closer inspection."

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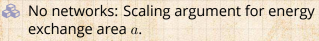


Let's never talk about this again:



"The fourth dimension of life: Fractal geometry and allometric scaling of organisms" (3

West, Brown, and Emquist, Science Magazine, , , 1999. [45]



Distinguish between biological and physical length scales (distance between mitochondria versus cell radius).

 $\red{8}$ Buckingham π action. [8]

 \red Arrive at $a \propto M^{D/D+1}$ and $\ell \propto M^{1/D}$.

New disaster: after going on about fractality of a, then state $v \propto a\ell$ in general.

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Really, quite confused:

Whole 2004 issue of Functional Ecology addresses the problem:

- 🚵 J. Kozlowski, M. Konrzewski. "Is West, Brown and Enquist's model of allometric scaling mathematically correct and biologically relevant?" Functional Ecology 18: 283-9, 2004. [24]
- 🚵 J. H. Brown, G. B. West, and B. J. Enquist. "Yes, West, Brown and Enquist's model of allometric scaling is both mathematically correct and biologically relevant." Functional Ecology 19: 735-738, 2005. [7]
- 🚵 J. Kozlowski, M. Konarzewski. "West, Brown and Enquist's model of allometric scaling again: the same questions remain." Functional Ecology 19: 739-743, 2005.

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Simple supply networks:

Banavar et al., Nature, (1999)[1].

- Flow rate argument.
- Ignore impedance.
 - Very general attempt to find most efficient transportation networks.

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Simple supply networks

Banavar et al. find 'most efficient' networks with

$$P \propto M^{d/(d+1)}$$

... but also find

$$V_{
m network} \propto M^{\,(d+1)/d}$$

$$d = 3$$
:

$$V_{\rm blood} \propto M^{4/3}$$

 \Leftrightarrow Consider a 3 g shrew with $V_{\text{blood}} = 0.1 V_{\text{body}}$

$$\Leftrightarrow$$
 3000 kg elephant with $V_{\rm blood}$ = $10V_{\rm body}$

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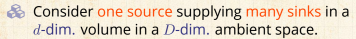




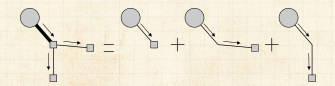


"Optimal Form of Branching Supply and Collection Networks"

Peter Sheridan Dodds, Phys. Rev. Lett., **104**, 048702, 2010. [11]



- Assume sinks are invariant.
- Assume sink density $\rho = \rho(V)$.
- Assume some cap on flow speed of material.
- See network as a bundle of virtual vessels:



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ightharpoonup Q: how does the number of sustainable sinks $N_{
m sinks}$ scale with volume V for the most efficient network design?

 \mathfrak{S} Or: what is the highest α for $N_{\mathsf{sinks}} \propto V^{\alpha}$?

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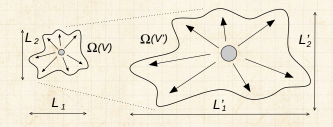








Allometrically growing regions:





Have d length scales which scale as

$$L_i \propto V^{\gamma_i}$$
 where $\gamma_1 + \gamma_2 + ... + \gamma_d = 1$.



 \Leftrightarrow For isometric growth, $\gamma_i = 1/d$.



For allometric growth, we must have at least two of the $\{\gamma_i\}$ being different

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Spherical cows and pancake cows:

Assume an isometrically Scaling family of cows:



Extremes of allometry: The pancake cows-



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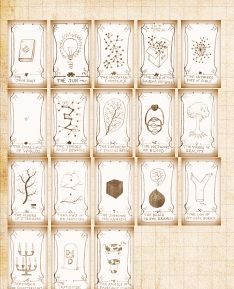
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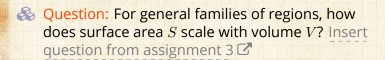






Spherical cows and pancake cows:

Question: How does the surface area S_{cow} of our two types of cows scale with cow volume V_{cow} ? Insert question from assignment 3 \square



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Metabolism and

Best and worst configurations (Banavar et al.)

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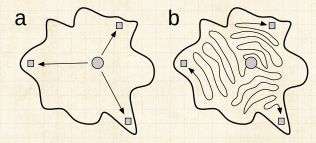
References











Rather obviously:

min $V_{\text{net}} \propto \sum$ distances from source to sinks.

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Minimal network volume:

Real supply networks are close to optimal:

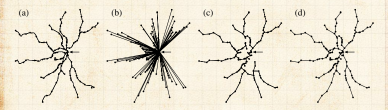


Figure 1. (a) Commuter rail network in the Boston area. The arrow marks the assumed root of the network. (b) Star graph. (c) Minimum spanning tree. (d) The model of equation (3) applied to the same set of stations.

Gastner and Newman (2006): "Shape and efficiency in spatial distribution networks" [15]

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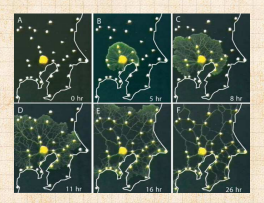








"Rules for Biologically Inspired Adaptive Network Design" Tero et al., Science, **327**, 439-442, 2010. [42]



Urban deslime in action:

https://www.youtube.com/watch?v=GwKuFREOgmo@

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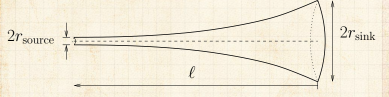
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We add one more element:



- Vessel cross-sectional area may vary with distance from the source.
- Flow rate increases as cross-sectional area decreases.
- e.g., a collection network may have vessels tapering as they approach the central sink.
- \implies Find that vessel volume v must scale with vessel length ℓ to affect overall system scalings.

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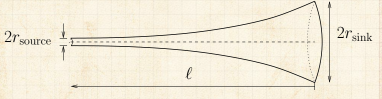
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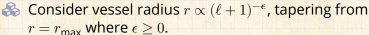


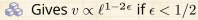




Effecting scaling:







$$\Leftrightarrow$$
 Gives $v \propto 1 - \ell^{-(2\epsilon - 1)} \to 1$ for large ℓ if $\epsilon > 1/2$

 \red Previously, we looked at $\epsilon = 0$ only.

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Minimal network volume:

For $0 \le \epsilon < 1/2$, approximate network volume by integral over region:

$$\min V_{\mathsf{net}} \propto \int_{\Omega_{d,D}(V)}
ho \, ||ec{x}||^{1-2\epsilon} \, \mathsf{d}ec{x}$$

Insert question , assignment 3 🗗 <2->

$$\propto
ho V^{1+\gamma_{\max}(1-2\epsilon)}$$
 where $\gamma_{\max} = \max_i \gamma_i$.

For $\epsilon > 1/2$, find simply that

$$\min V_{\rm net} \propto \rho V$$

So if supply lines can taper fast enough and without limit, minimum network volume can be made negligible.

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For $0 \le \epsilon < 1/2$:



$$\min V_{\rm net} \propto \rho V^{1+\gamma_{\rm max}(1-2\epsilon)}$$

 \mathfrak{R} If scaling is isometric, we have $\gamma_{\mathsf{max}} = 1/d$:

$$\min V_{\mathsf{net/iso}} \propto \rho V^{1+(1-2\epsilon)/d}$$



\$ If scaling is allometric, we have $\gamma_{\text{max}} = \gamma_{\text{allo}} > 1/d$: and

$$\min V_{\rm net/allo} \propto \rho V^{1+(1-2\epsilon)\gamma_{\rm allo}}$$



Isometrically growing volumes require less network volume than allometrically growing volumes:

$$\frac{\min V_{\text{net/iso}}}{\min V_{\text{net/allo}}} \to 0 \text{ as } V \to \infty$$

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For $\epsilon > 1/2$:



$$\min V_{\mathsf{net}} \propto
ho V$$

Network volume scaling is now independent of overall shape scaling.

Limits to scaling

- & Can argue that ϵ must effectively be 0 for real networks over large enough scales.
- Limit to how fast material can move, and how small material packages can be.
- & e.g., blood velocity and blood cell size.

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- Velocity at capillaries and aorta approximately constant across body size [44]: $\epsilon = 0$.
- Material costly ⇒ expect lower optimal bound of $V_{\mathsf{net}} \propto \rho V^{(d+1)/d}$ to be followed closely.
- A For cardiovascular networks, d = D = 3.
- Blood volume scales linearly with body volume [40], $V_{\mathsf{net}} \propto V$.
- Sink density must : decrease as volume increases:

$$\rho \propto V^{-1/d}$$
.

Density of suppliable sinks decreases with organism size.

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Blood networks

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 \aleph Then P, the rate of overall energy use in Ω, can at most scale with volume as

$$P \propto \rho V \propto \rho \, M \propto M^{\,(d-1)/d}$$

A For d=3 dimensional organisms, we have

$$P \propto M^{2/3}$$

Including other constraints may raise scaling exponent to a higher, less efficient value. Metabolism and Truthicide

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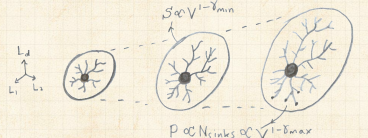
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Exciting bonus: Scaling obtained by the supply network story and the surface-area law only match for isometrically growing shapes. Insert question from assignment 3 2

The surface area-supply network mismatch for allometrically growing shapes:



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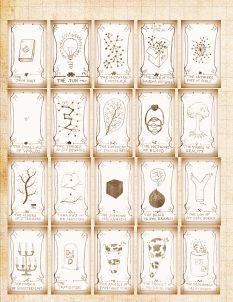
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- mammals up to 10-30 kg
- For mammals > 10-30 kg, maybe we have a new scaling regime
- 🙈 Economos: limb length break in scaling around 20 kg
- White and Seymour, 2005: unhappy with large herbivore measurements. Find $\alpha \simeq 0.686 + 0.014$

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Stefan-Boltzmann law:



$$\frac{\mathrm{d}E}{\mathrm{d}t} = \sigma S T^4$$

where *S* is surface and *T* is temperature.

Very rough estimate of prefactor based on scaling of normal mammalian body temperature and surface area S:

$$B \simeq 10^5 M^{2/3} {\rm erg/sec.}$$

 \clubsuit Measured for $M \leq 10$ kg:

$$B = 2.57 \times 10^5 M^{2/3}$$
erg/sec.

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Many sources and one sink.

& ε?

& Assume ρ is constant over time and $\epsilon = 0$:

$$V_{
m net} \propto
ho V^{(d+1)/d} = {
m constant} imes V^{3/2}$$

Network volume grows faster than basin 'volume' (really area).

It's all okay:

Landscapes are d=2 surfaces living in D=3 dimensions.

Streams can grow not just in width but in depth...

 \iff If $\epsilon > 0$, V_{net} will grow more slowly but 3/2 appears to be confirmed from real data.

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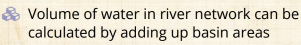
onclusion







Hack's law



Flows sum in such a way that

$$V_{\mathsf{net}} = \sum_{\mathsf{all \ pixels}} a_{\mathsf{pixel} \ i}$$

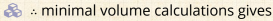
A Hack's law again:

$$\ell \sim a^h$$

🙈 Can argue

$$V_{\mathrm{net}} \propto V_{\mathrm{basin}}^{1+h} = a_{\mathrm{basin}}^{1+h}$$

where h is Hack's exponent.



$$h = 1/2$$

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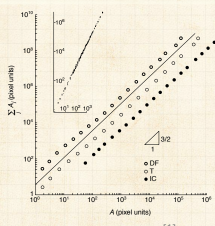
Conclusion







- Banavar et al.'s approach [1] is okay because ρ really is constant.
- The irony: shows optimal basins are isometric
- Optimal Hack's law: $\ell \sim a^h$ with h = 1/2
- (Zzzzz)



From Banavar et al. (1999)^[1]

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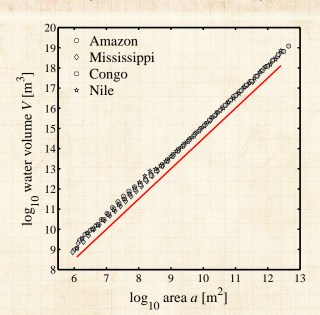








Even better—prefactors match up:



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Banavar et al., 2010, PNAS: "A general basis for quarter-power scaling in animals." [2]

#It has been known for decades that the metabolic rate of animals scales with body mass with an exponent that is almost always < 1, > 2/3, and often very close to 3/4."

Sough, cough, cough, hack, wheeze, cough.

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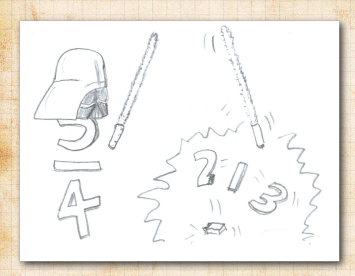
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Stories—Darth Quarter:



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Some people understand it's truly a disaster:





Peter Sheridan Dodds, Theoretical Biology's Buzzkill

By Mark Changizi | February 9th 2010 03:24 PM | 1 comment | Print | E-mail | Track Comments





There is an apocryphal story about a graduate mathematics student at the University of Virginia studying the properties of certain mathematical objects. In his fifth year some killjoy bastard elsewhere published a paper proving that there are no such mathematical objects. He dropped out of the program, and I never did hear where

he is today. He's probably making my cappuccino right now.

This week, a professor named Peter Sheridan Dodds published a new paper in *Physical Review Letters* further fleshing out a theory concerning why a 2/3 power law may apply for metabolic rate. The 2/3 law says that metabolic rate in animals rises as the 2/3 power of body mass. It was in a 2001 *Journal of Theoretical Biology* paper that he first argued that perhaps a 2/3 law applies, and that paper – along with others such as the one that just appeared — is what has put him in the Killipy Hall of Fame. The University of Virginia's killiov was a mere amateur.

Mark Changizi

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III Article:

ABOUT MARK

Mark Changizi is Director of Human Cognition at 2AI, and the author of *The Vision Revolution* (Benbella 2009) and *Harnessed: How.*...

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The unnecessary bafflement continues:

"Testing the metabolic theory of ecology" [33]

C. Price, J. S. Weitz, V. Savage, J. Stegen, A. Clarke, D. Coomes, P. S. Dodds, R. Etienne, A. Kerkhoff, K. McCulloh, K. Niklas, H. Olff, and N. Swenson Ecology Letters, 15, 1465-1474, 2012.

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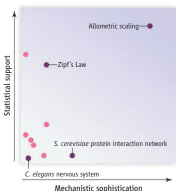






Artisanal, handcrafted silliness:

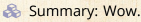
"Critical truths about power laws" [41] Stumpf and Porter, Science, 2012



How good is your power law? The chart reflects the level of statistical support—as measured in (16. 21)—and our opinion about the mechanistic sophistication underlying hypothetical generative models for various reported power laws. Some relationships are identified by name; the others reflect the general characteristics of a wide range of reported power laws. Allometric scaling stands out from the other power laws reported for complex systems.



Call generalization of Central Limit Theorem, stable distributions. Also: PLIPLO action.



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- Supply network story consistent with dimensional analysis.
- Isometrically growing regions can be more efficiently supplied than allometrically growing ones.
- Ambient and region dimensions matter (D=d versus D>d).
- Deviations from optimal scaling suggest inefficiency (e.g., gravity for organisms, geological boundaries).
- Actual details of branching networks not that important.
- Exact nature of self-similarity varies.
- 2/3-scaling lives on, largely in hiding.
- 3/4-scaling? Jury ruled a mistrial.
- The truth will out. Maybe.

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