Scaling

## Scaling—a Plenitude of Power Laws

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Scaling-at-large

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### Definitions

General observation:

Systems (complex or not) that cross many spatial and temporal scales often exhibit some form of scaling. Scaling-at-large

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#### Scaling Outline Scaling-at-large Allometry Definitions Examples History: Metabolism Measuring exponents All about scaling: History: River networks Earlier theories Geometric argument Blood networks Definitions. River networks Conclusion Examples. References How to measure your power-law relationship. Mechanisms giving rise to your power-laws.



### Definitions

### A power law relates two variables x and y as follows:

$$\prime = {\it CX}^{lpha}$$

### • $\alpha$ is the scaling exponent (or just exponent)

V

- (α can be any number in principle but we will find various restrictions.)
- c is the prefactor (which can be important!)

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### Definitions

- The prefactor c must balance dimensions.
- eg., length l and volume v of common nails are related as:

$$\ell = c v^{1/2}$$

Using [·] to indicate dimension, then

$$[c] = [I]/[V^{1/4}] = L/L^{3/4} = L^{1/4}$$

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### Looking at data

Power-law relationships are linear in log-log space:

$$y = cx^{lpha}$$

 $\Rightarrow \log_b y = \alpha \log_b x + \log_b c$ 

with slope equal to  $\alpha$ , the scaling exponent.

- Much searching for straight lines on log-log or double-logarithmic plots.
- Good practice: Always, always, always use base 10.
- Talk only about orders of magnitude (powers of 10).

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### Why is $\alpha \simeq 1.23$ ?

### Quantities (following Zhang and Sejnowski):

- ► *G* = Volume of gray matter (cortex/processors)
- W = Volume of white matter (wiring)
- T = Cortical thickness (wiring)
- S =Cortical surface area
- L = Average length of white matter fibers
- p = density of axons on white matter/cortex interface

### A rough understanding:

- $G \sim ST$  (convolutions are okay)
- $W \sim \frac{1}{2} pSL$
- $G \sim L^3 \leftarrow$  this is a little sketchy...
- Eliminate S and L to find  $W \propto G^{4/3}/T$

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### Why is $\alpha \simeq 1.23$ ?

### A rough understanding:

- We are here:  $W \propto G^{4/3}/T$
- Observe weak scaling  $T \propto G^{0.10\pm0.02}$ .
- (Implies  $S \propto G^{0.9} \rightarrow$  convolutions fill space.)
- $ightarrow W \propto G^{4/3}/T \propto G^{1.23\pm0.02}$

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### Why is $\alpha \simeq 1.23$ ?



- With V = G + W, some power laws must be approximations.
- Measuring exponents is a hairy business...



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### Good scaling:

### General rules of thumb:

- High quality: scaling persists over three or more orders of magnitude for each variable.
- Medium quality: scaling persists over three or more orders of magnitude for only one variable and at least one for the other.
- Very dubious: scaling 'persists' over less than an order of magnitude for both variables.



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### Unconvincing scaling:



Average walking speed as a function of city

from Bettencourt et al. (2007)<sup>[4]</sup>; otherwise very interesting!

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### Definitions

Power laws are the signature of scale invariance:

Scale invariant 'objects' look the 'same' when they are appropriately rescaled.

- Objects = geometric shapes, time series, functions, relationships, distributions,...
- Same' might be 'statistically the same'
- To rescale means to change the units of measurement for the relevant variables

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### Scale invariance

### Our friend $y = cx^{\alpha}$ :

• If we rescale x as x = rx' and y as  $y = r^{\alpha}y'$ ,

#### then

►

►

$$r^{lpha}y' = c(rx')^{lpha}$$

$$\Rightarrow y' = cr^{\alpha}x'^{\alpha}r^{-\alpha}$$

$$\Rightarrow$$
 y' = cx' <sup>$\alpha$</sup> 

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### Scale invariance

### Compare with $y = ce^{-\lambda x}$ :

If we rescale x as 
$$x = rx'$$
, ther

$$y = ce^{-\lambda r \lambda}$$

Original form cannot be recovered.

ightarrow  $\Rightarrow$  scale matters for the exponential.





### Scale invariance

### More on $y = ce^{-\lambda x}$ :

- Say  $x_0 = 1/\lambda$  is the characteristic scale.
- For  $x \gg x_0$ , y is small, while for  $x \ll x_0$ , y is large.
- $\blacktriangleright$   $\Rightarrow$  More on this later with size distributions.

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<ul> <li>fers to differential growth rates of the parts of a living organism's body part or process.</li> <li>First proposed by Huxley and Teissier, Nature, 1936 "Terminology of relative growth" <sup>[21]</sup></li> </ul>	Blood networks River networks Conclusion References
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### Definitions

### Isometry versus Allometry:

- Isometry = 'same measure'
- Allometry = 'other measure'

# Confusingly, we use allometric scaling to refer to both:

- 1. nonlinear scaling (e.g.,  $x \propto y^{1/3}$ )
- 2. and the relative scaling of different measures (e.g., resting heart rate as a function of body size)

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### A wonderful treatise on scaling:

ON SIZE AND LIFE

THOMAS A. MCMAHON AND JOHN TYLER BONNER



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McMahon and Bonner, 1983<sup>[26]</sup>

### For the following slide:

The biggest living things (left). All the organisms are drawn to the same scale. 1. The largest flying bird (albatross); 2, the largest known animal (the blue whale), 3, the largest extinct land mammal (Baluchitherium) with a human figure shown for scale; 4, the tallest living land animal (giraffe); 5, Tyrannosaurus: 6. Diplodocus: 7, one of the largest flying reptiles (Pteranodon); 8, the largest extinct snake; 9, the length of the largest tapeworm found in man; 10, the largest living reptile (West African crocodile); 11, the largest extinct lizard; 12, the largest extinct bird (Aepyornis); 13, the largest jellyfish (Cyanea); 14, the largest living lizard (Komodo dragon); 15, sheep; 16, the largest bivalve mollusc (Tridacna); 17; the largest fish (whale shark); 18, horse; 19, the largest crustacean (Japanese spider crab); 20, the largest sea scorpion (Eurypterid); 21, large tarpon; 22, the largest lobster; 23, the largest mollusc (deep-water squid. Architeuthis); 24, ostrich; 25, the lower 105 feet of the largest organism (giant sequoia), with a 100-foot larch superposed.



### The many scales of life:





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p. 2, McMahon and Bonner<sup>[26]</sup>

### For the following slide:

Medium-sized creatures (above). 1, Dog; 2, common herring; 3, the largest egg (Aepyornis); 4, song thrush with egg: 5, the smallest bird (hummingbird) with egg; 6, queen bee; 7, common cockroach; 8, the largest stick insect; 9, the largest polyp (Branchiocerianthus); 10, the smallest mammal (flying shrew); 11, the smallest vertebrate (a tropical frog); 12, the largest frog (goliath frog); 13, common grass frog; 14, house mouse; 15, the largest land snail (Achatina) with egg; 16, common snail; 17, the largest beetle (goliath beetle); 18, human hand; 19, the largest starfish (Luidia); 20, the largest free-moving protozoan (an extinct nummulite).

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### For the following slide:

Small, "naked-eye" creatures (*lower left*). 1, One of the smallest fishes (*Trimmatom nanus*); 2, common brown hydra, expanded; 3, housefly; 4, medium-sized ant; 5, the smallest vertebrate (a tropical frog, the same as the one numbered 11 in the figure *above*); 6, flea (*Xenopsylla cheopis*); 7, the smallest land snail; 8, common water flea (*Daphnia*).

The smallest "naked-eye" creatures and some large microscopic animals and cells (*below right*). 1, Vorticella, a ciliate; 2, the largest ciliate protozoan (*Bursaria*); 3, the smallest many-celled animal (a rotifer); 4, smallest flying insect (*Elaphis*); 5, another ciliate (*Paramecium*); 6, cheese mite; 7, human sperm; 8, human ovum; 9, dysentery amoeba; 10, human liver cell; 11, the foreleg of the flea (numbered 6 in the figure to the *left*).



p. 2, McMahon and Bonner<sup>[26]</sup>

### The many scales of life:





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### Non-uniform growth:



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p. 32, McMahon and Bonner <sup>[26]</sup>

years  $0 \cdot 42$ 

0 • 75

2.75

6 • 75 12 • 75

25 • 75





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p. 36, McMahon and Bonner <sup>[26]</sup>



### The allometry of nails:

### A buckling instability?:

- Physics/Engineering result: Columns buckle under a load which depends on d<sup>4</sup>/l<sup>2</sup>.
- To drive nails in, resistive force  $\propto$  nail circumference =  $\pi d$ .
- Match forces independent of nail size:  $d^4/\ell^2 \propto d$ .
- Leads to  $d \propto \ell^{2/3}$ .
- ► Argument made by Galileo<sup>[13]</sup> in 1638 in "Discourses on Two New Sciences." (⊞) [pdf] Also, see here (⊞).
- Also see McMahon, "Size and Shape in Biology," Science, 1973.<sup>[24]</sup>

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### Rowing: Speed $\propto$ (number of rowers)^{1/9}

No. of

oarsmen

30

25 20 Speed (km/hr)

10

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### Scaling in Cities:

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•	"Growth, innovation, scaling, and the pace of life in cities"
	Bettencourt et al., PNAS, 2007. <sup>[4]</sup>
	Quantified levels of
	<ul> <li>Infrastructure</li> <li>Wealth</li> <li>Crime levels</li> </ul>
	► Disease
	<ul> <li>Energy consumption</li> </ul>
	as a function of city size N (population).
# Scaling in Cities:

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Iabi	e i		calling	expon	ients i	01.1	urban	inuical	lors	vs. (	JUY	size

Y	β	95% CI	Adj-R <sup>2</sup>	Observations	Country-year	Allometry
New patents	1.27	[1.25, 1.29]	0.72	331	U.S. 2001	Definitions
Inventors	1.25	[1.22,1.27]	0.76	331	U.S. 2001	Examples
Private R&D employment	1.34	[1.29,1.39]	0.92	266	U.S. 2002	History: Metabolism
"Supercreative" employment	1.15	[1.11,1.18]	0.89	287	U.S. 2003	Measuring exponents
R&D establishments	1.19	[1.14,1.22]	0.77	287	U.S. 1997	History: River networks
R&D employment	1.26	[1.18,1.43]	0.93	295	China 2002	Earlier theories
Total wages	1.12	[1.09,1.13]	0.96	361	U.S. 2002	Geometric argument
Total bank deposits	1.08	[1.03,1.11]	0.91	267	U.S. 1996	Blood networks
GDP	1.15	[1.06,1.23]	0.96	295	China 2002	River networks
GDP	1.26	[1.09,1.46]	0.64	196	EU 1999–2003	Conclusion
GDP	1.13	[1.03,1.23]	0.94	37	Germany 2003	References
Total electrical consumption	1.07	[1.03,1.11]	0.88	392	Germany 2002	
New AIDS cases	1.23	[1.18,1.29]	0.76	93	U.S. 2002-2003	
Serious crimes	1.16	[1.11, 1.18]	0.89	287	U.S. 2003	
Total housing	1.00	[0.99,1.01]	0.99	316	U.S. 1990	
Total employment	1.01	[0.99,1.02]	0.98	331	U.S. 2001	111
Household electrical consumption	1.00	[0.94,1.06]	0.88	377	Germany 2002	1 in
Household electrical consumption	1.05	[0.89,1.22]	0.91	295	China 2002	-( )-
Household water consumption	1.01	[0.89,1.11]	0.96	295	China 2002	-15%
Gasoline stations	0.77	[0.74,0.81]	0.93	318	U.S. 2001	いに、
Gasoline sales	0.79	[0.73,0.80]	0.94	318	U.S. 2001	Y
Length of electrical cables	0.87	[0.82,0.92]	0.75	380	Germany 2002	S
Road surface	0.83	[0.74,0.92]	0.87	29	Germany 2002	<b>e</b>

Data sources are shown in SI Text. CI, confidence interval; Adj-R<sup>2</sup>, adjusted R<sup>2</sup>; GDP, gross domestic product.

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# Scaling in Cities:

#### Intriguing findings:

- Global supply costs scale sublinearly with N ( $\beta < 1$ ).
  - Returns to scale for infrastructure.
- Total individual costs scale linearly with N ( $\beta = 1$ )
  - Individuals consume similar amounts independent of city size.
- Social quantities scale superlinearly with N ( $\beta > 1$ )
  - Creativity (# patents), wealth, disease, crime, ...

#### Density doesn't seem to matter...

 Surprising given that across the world, we observe two orders of magnitude variation in area covered by agglomerations of fixed populations. Scaling-at-large

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#### Scaling Ecology—Species-area law: $N_{\rm species} \propto A^{\beta}$ Scaling-at-large Allometry Definitions Examples History: Metabolism Measuring exponents History: River networks Earlier theories Allegedly (data is messy): Geometric argument Blood networks River networks Conclusion • On islands: $\beta \approx 1/4$ . References • On continuous land: $\beta \approx 1/8$ .



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### A focus:

His

- How much energy do organisms need to live?
- And how does this scale with organismal size?

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# Animal power

Fundamental biological and ecological constraint:

 $P = c M^{\alpha}$ 

P = basal metabolic rate

M = organismal body mass







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# $P = c M^{\alpha}$

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

39–41 <i>°C</i>
36–38° <i>C</i>
34–36° <i>C</i>
30–31 <i>°C</i>

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# What one might expect:

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

 Dimensional analysis suggests an energy balance surface law:

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

#### Lognormal fluctuations:

Gaussian fluctuations in log *P* around log  $cM^{\alpha}$ .

Stefan-Boltzmann law (⊞) for radiated energy:

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

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#### The prevailing belief of the church of Scaling quarterology Scaling-at-large Allometry Definitions Examples History: Metabolism Measuring exponents History: River networks Earlier theories Geometric argument $\alpha = 3/4$ Blood networks River networks Conclusion References $P \propto M^{3/4}$ Huh? UNIVERSITY

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# Related putative scalings:

- number of capillaries  $\propto M^{3/4}$
- time to reproductive maturity  $\propto M^{1/4}$
- heart rate  $\propto M^{-1/4}$
- cross-sectional area of aorta  $\propto M^{3/4}$
- population density  $\propto M^{-3/4}$

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#### History Scaling Scaling-at-large Allometry Definitions Examples 1840's: Sarrus and Rameaux<sup>[33]</sup> first suggested $\alpha = 2/3$ . History: Metabolism Measuring exponents History: River networks Earlier theories Geometric argument Blood networks River networks Conclusion References

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#### Scaling History Scaling-at-large Allometry Definitions Examples 1883: Rubner<sup>[31]</sup> found $\alpha \simeq 2/3$ . History: Metabolism Measuring exponents History: River networks Earlier theories Geometric argument Blood networks River networks Conclusion References UNIVERSITY ∽ < < > 51 of 126

# 1930's: Brody, Benedict study mammals. <sup>[7]</sup> Found $\alpha \simeq 0.73$ (standard).



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1950/1960: Hemmingsen <sup>[18, 19]</sup> Extension to unicellular organisms.  $\alpha = 3/4$  assumed true.



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1964: Troon, Scotland: <sup>[5]</sup> 3rd symposium on energy metabolism.  $\alpha = 3/4$  made official . . .



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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

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- But—much controversy...
- See 'Re-examination of the "3/4-law" of metabolism' Dodds, Rothman, and Weitz<sup>[11]</sup> and ensuing madness...



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Passerine vs. non-passerine...

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#### Linear regression

#### Important:

- Ordinary Least Squares (OLS) Linear regression is only appropriate for analyzing a dataset {(x<sub>i</sub>, y<sub>i</sub>)} when we know the x<sub>i</sub> are measured without error.
- 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. [32, 30]

```
(aka Reduced Major Axis = RMA.)
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#### Measuring exponents Scaling Scaling-at-large Allometry Definitions Examples History: Metabolism Measuring exponents For Standardized Major Axis Linear Regression: History: River networks Earlier theories Geometric argument Blood networks standard deviation of y data **Biver networks** $slope_{sma} =$ Conclusion standard deviation of x data

- Very simple!
- Scale invariant.



# Measuring exponents

Relationship to ordinary least squares regression is simple:

$$slope_{SMA} = r^{-1} \times slope_{OLS y \text{ on } x}$$
$$= r \times slope_{OLS x \text{ on } y}$$

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, 19	91 (3	91 Mammals)	Scaling
	range of M	N	â	Scaling-at-large
	$\leq$ 0.1 kg	167	$0.678\pm0.038$	Definitions Examples History: Metabolism Measuring exponents
	$\leq$ 1 kg	276	$0.662\pm0.032$	History: River networks Earlier theories Geometric argument Blood networks
	$\leq$ 10 kg	357	$0.668\pm0.019$	River networks Conclusion References
	$\leq$ 25 kg	366	$0.669\pm0.018$	
	$\leq$ 35 kg	371	$0.675\pm0.018$	
	$\leq$ 350 kg	389	$0.706\pm0.016$	N.
	$\leq$ 3670 kg	391	0.710 ± 0.021	

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Bennett a	Ind Harve	ey, 19 ⊢ <i>№</i>	987 (398 birds) â	Scaling
	< 0.032	162	0.636 ± 0.103	Scaling-at-large Allometry Definitions Examples
	≤ 0.1	236	$0.602\pm0.060$	History: Metabolism Measuring exponents History: River networks Earlier theories
	≤ 0.32	290	$0.607\pm0.039$	Geometric argument Blood networks River networks Conclusion
	<u>≤</u> 1	334	$0.652\pm0.030$	References
	<b>≤ 3.2</b>	371	$0.655\pm0.023$	
	<u>≤</u> 10	391	$0.664\pm0.020$	
	≤ 32	396	$0.665\pm0.019$	
	≤ 100	398	$0.664\pm0.019$	Deco 65 of 126

#### Hypothesis testing

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

 $H_0: \alpha = \alpha' \text{ and } H_1: \alpha \neq \alpha'.$ 

- Assume each B<sub>i</sub> (now a random variable) is normally distributed about α' log<sub>10</sub> M<sub>i</sub> + log<sub>10</sub> c.
- Follows that the measured α for one realization obeys a t distribution with N – 2 degrees of freedom.
- 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."<sup>[8]</sup>

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Re	visiting tl	ne pa	ıst—m	ammals	3	Scaling
Fu	II mass ran	ge: N	$\hat{lpha}$	<i>p</i> <sub>2/3</sub>	<b>P</b> <sub>3/4</sub>	Allometry Definitions Examples History: Metabolism Measuring exponents History: Metabolism
	Kleiber	13	0.738	< 10 <sup>-6</sup>	0.11	Earlier theories Geometric argument Blood networks River networks
	Brody	35	0.718	< 10 <sup>-4</sup>	< 10 <sup>-2</sup>	Conclusion
	Heusner	391	0.710	< 10 <sup>-6</sup>	< 10 <sup>-5</sup>	
a	Bennett nd Harvey	398	0.664	0.69	< 10 <sup>-15</sup>	N. S.
						Deversity S

Revisitin	g th	e past-	—mam	mals	Scaling
<i>M</i> ≤ 10 kg:	N	â	p <sub>2/3</sub>	P <sub>3/4</sub>	Scaling-at-large
Kleiber	5	0.667	0.99	0.088	History: Metabolism Measuring exponents History: River network
Brody	26	0.709	< 10 <sup>-3</sup>	< 10 <sup>-3</sup>	Earlier theories Geometric argument Blood networks Blong actuation
Heusner	357	0.668	0.91	< 10 <sup>-15</sup>	Conclusion
<i>M</i> ≥ 10 kg:	N	$\hat{lpha}$	<i>p</i> <sub>2/3</sub>	<i>p</i> <sub>3/4</sub>	
Kleiber	8	0.754	$< 10^{-4}$	0.66	
Brody	9	0.760	< 10 <sup>-3</sup>	0.56	J. J.
Heusner	34	0.877	< 10 <sup>-12</sup>	< 10 <sup>-7</sup>	

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#### Analysis of residuals

- 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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# Analysis of residuals

We use the spiffing Spearman Rank-Order Correlation Cofficient (⊞)

#### Basic idea:

Þ

- Given {(x<sub>i</sub>, y<sub>i</sub>)}, rank the {x<sub>i</sub>} and {y<sub>i</sub>} separately from smallest to largest. Call these ranks R<sub>i</sub> and S<sub>i</sub>.
- Now calculate correlation coefficient for ranks, r<sub>s</sub>:

$$r_{s} = \frac{\sum_{i=1}^{n} (R_{i} - \bar{R})(S_{i} - \bar{S})}{\sqrt{\sum_{i=1}^{n} (R_{i} - \bar{R})^{2}} \sqrt{\sum_{i=1}^{n} (S_{i} - \bar{S})^{2}}}$$

Perfect correlation: x<sub>i</sub>'s and y<sub>i</sub>'s both increase monotonically. Scaling

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### Analysis of residuals

#### We assume all rank orderings are equally likely:

- *r<sub>s</sub>* is distributed according to a Student's
   *t*-distribution (⊞) 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.<sup>[29]</sup>

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#### Recap:

- So: The exponent α = 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<sup>[12]</sup>
- But see later: non-isometric growth leads to lower metabolic scaling. Oops.

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### The widening gyre:

#### Now we're really confused (empirically):

- White and Seymour, 2005: unhappy with large herbivore measurements<sup>[40]</sup>. Pro 2/3: Find α ≃ 0.686 ± 0.014.
- White ... <sup>[39]</sup>
- Glazier, BioScience (2006) <sup>[16]</sup>: "The 3/4-Power Law Is Not Universal: Evolution of Isometric, Ontogenetic Metabolic Scaling in Pelagic Animals."
- Glazier, Biol. Rev. (2005)<sup>[15]</sup>: "Beyond the 3/4-power law': variation in the intra- and interspecific scaling of metabolic rate in animals."
- Savage et al., PLoS Biology (2008) <sup>[34]</sup> "Sizing up allometric scaling theory" Pro 3/4: problems claimed to be finite-size scaling.
- Mori et al.<sup>[?]</sup>
- Add more.



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### Basic basin quantities: $a, I, L_{\parallel}, L_{\perp}$ :



- a = drainage basin area
- length of longest (main) stream
- $L = L_{\parallel} =$ longitudinal length of basin

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#### **River networks**

 1957: J. T. Hack<sup>[17]</sup>
 "Studies of Longitudinal Stream Profiles in Virginia and Maryland"

 *l* ~ a<sup>h</sup>

$$h \sim 0.6$$

- 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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#### Earlier theories

#### Building on the surface area idea...

 Blum (1977)<sup>[6]</sup> speculates on four-dimensional biology: d

$$P \propto M^{(d-1)/2}$$

So we need another dimension...

Obviously, a bit silly... <sup>[35]</sup>

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Blood networks **Biver networks** 



#### Earlier theories

#### Building on the surface area idea:

- McMahon (70's, 80's): Elastic Similarity <sup>[24, 26]</sup>
- Idea is that organismal shapes scale allometrically with 1/4 powers (like nails and trees...)
- Appears to be true for ungulate legs...<sup>[25]</sup>
- Metabolism and shape never properly connected.

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#### Nutrient delivering networks:

- 1960's: Rashevsky considers blood networks and finds a 2/3 scaling.
- ▶ 1997: West et al. [38] use a network story to find 3/4 scaling.



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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:

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

- networks are fractal
- quarter powers everywhere

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#### Impedance measures:

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):

$$Z\propto\sum_{k=0}^{N}rac{h_{k}^{1/2}}{r_{k}^{5/2}N_{k}}$$

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#### Not so fast ....

#### Actually, model shows:

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

#### Do find:

Murray's cube law (1927) for outer branches: <sup>[28]</sup>

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

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

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#### Scaling Connecting network structure to $\alpha$ 1. Ratios of network parameters: Scaling-at-large $R_n = \frac{n_{k+1}}{n_k}, \ R_\ell = \frac{\ell_{k+1}}{\ell_k}, \ R_r = \frac{r_{k+1}}{r_k}$ Allometry Definitions Examples History: Metabolism 2. Number of capillaries $\propto P \propto M^{\alpha}$ . Measuring exponents History: River networks $\Rightarrow \alpha = -\frac{\ln R_n}{\ln R_r^2 R_\ell}$ Earlier theories Geometric argument Blood networks **Biver networks** (also problematic due to prefactor issues) References Soldiering on, assert: • area-preservingness: $R_r = R_n^{-1/2}$ • space-fillingness: $R_{\ell} = R_n^{-1/3}$

 $\Rightarrow \alpha = 3/4$ 



### Data from real networks

Network	R <sub>n</sub>	$R_{r}^{-1}$	$R_\ell^{-1}$	$-\frac{\ln R_r}{\ln R_n}$	$-\frac{\ln R_{\ell}}{\ln R_{n}}$	α	Scaling-at-large
West <i>et al.</i>	-	-	-	1/2	1/3	3/4	Definitions Examples History: Metabolism
rat (PAT)	2.76	1.58	1.60	0.45	0.46	0.73	Measuring exponents History: River networks Ceometric argument Biodn networks River networks Conclusion References
cat (PAT) (Turcotte <i>et al.</i> <sup>[37]</sup> )	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	N.
human (PAT) human (PAT)	3.03 3.36	1.60 1.56	1.49 1.49	0.42 0.37	0.36 0.33	0.83 0.94	

## Simple supply networks









 Banavar et al., Nature, (1999)<sup>[1]</sup>

- 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

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

▶ *d* = 3:

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

- Consider a 3 g shrew with  $V_{\text{blood}} = 0.1 V_{\text{body}}$
- ►  $\Rightarrow$  3000 kg elephant with  $V_{\text{blood}} = 10 V_{\text{body}}$

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- "Optimal Form of Branching Supply and Collection Networks." Dodds, Phys. Rev. Lett., 2010.<sup>[9]</sup>
- Consider one source supplying many sinks in a d-dim. volume in a D-dim. ambient space.
- 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:





- Q: how does the number of sustainable sinks N<sub>sinks</sub> scale with volume V for the most efficient network design?
- Or: what is the highest  $\alpha$  for  $N_{\text{sinks}} \propto V^{\alpha}$ ?

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Have d length scales which scale as

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

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

 For allometric growth, we must have at least two of the {γ<sub>i</sub>} being different



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Best and worst configurations (Banavar et al.)



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



#### Minimal network volume:





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.

(2006) Gastner and Newman<sup>[14]</sup>: "Shape and efficiency in spatial distribution networks"



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#### Minimal network volume:

Approximate network volume by integral over region:

min 
$$V_{
m net} \propto \int_{\Omega_{d,D}(V)} 
ho \, ||ec{x}|| \, \mathrm{d}ec{x}$$

$$\rightarrow \rho V^{1+\gamma_{\max}} \int_{\Omega_{d,D}(c)} (c_1^2 u_1^2 + \ldots + c_k^2 u_k^2)^{1/2} \mathrm{d}\vec{u}$$

 $\propto 
ho V^{1+\gamma_{
m max}}$ 

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#### Scaling Geometric argument General result: Scaling-at-large min $V_{\rm net} \propto \rho V^{1+\gamma_{\rm max}}$ Allometry Definitions Examples History: Metabolism Measuring exponents History: River networks Earlier theories • If scaling is isometric, we have $\gamma_{max} = 1/d$ : Geometric argument Blood networks **Biver networks** min $V_{\text{net/iso}} \propto \rho V^{1+1/d} = \rho V^{(d+1)/d}$ References • If scaling is allometric, we have $\gamma_{max} = \gamma_{allo} > 1/d$ : and min $V_{ m net/allo} \propto ho V^{1+\gamma_{ m allo}}$ 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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#### **Blood networks**

• Material costly  $\Rightarrow$  expect lower optimal bound of  $V_{\text{net}} \propto \rho V^{(d+1)/d}$  to be followed closely.

- For cardiovascular networks, d = D = 3.
- Blood volume scales linearly with body volume <sup>[36]</sup>, V<sub>net</sub>  $\propto$  V.
- Sink density must ... decrease as volume increases:

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

Density of suppliable sinks decreases with organism size.

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#### **Blood networks**

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

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

For d = 3 dimensional organisms, we have







Prefactor:

►





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:

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

• Measured for  $M \le 10$  kg:

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

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River networks



#### **River networks**

View river networks as collection networks.

- Many sources and one sink.
- Assume ρ is constant over time:

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

- Network volume grows faster than basin 'volume' (really area).
- It's all okay:

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

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

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#### Hack's law

- Volume of water in river network can be calculated by adding up basin areas
- Flows sum in such a way that

$$V_{\rm net} = \sum_{\rm all \ pixels} a_{\rm pixels}$$

Hack's law again:

$$\ell \sim a^h$$

Can argue

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

where h is Hack's exponent.

... minimal volume calculations gives

$$h = 1/2$$

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# Yet more theoretical madness from the Quarterologists:

- Banavar et al., 2010, PNAS:
   "A general basis for quarter-power scaling in animals." <sup>[2]</sup>
- "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."
- Cough, cough, cough, hack, wheeze, cough.

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#### Conclusion

- 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.

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