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Allometric scaling of production and life-history variation in vascular plants

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1958–88 (see above). Matching data from 8 trees sampled on the east end of Isle Royale, where balsam fir are not heavily or repeatedly browsed by moose¹⁰, provided a natural control for the test of our hypothesis relating climate and wolf behaviour to the dynamics of moose and fir growth. On the east end, fir growth did not correlate with the NAO index at lags of 0 ($r = 0.074$, $P = 0.69$) or 1 year ($r = 0.069$, $P = 0.71$) or with moose density at a lag of 1 year ($r = 0.054$, $P = 0.78$).

The North Atlantic Oscillation

The NAO is a meridional alternation in atmospheric mass balance between pressure centres over the Azores and Iceland. The winter NAO index is calculated on the basis of the normalized sea-level pressure difference between these two centres from December to March¹⁷. A negative difference indicates weak westerly winds across the Atlantic Ocean and unusually warm winters over north eastern North America; opposing conditions prevail during positive years¹⁷. We used data from the Climate Indices website of the National Center for Atmospheric Research, Boulder, USA (http://www.cgd.ucar.edu/cas/climind/). Although annual snow depth measures during the study period are not available for Isle Royale, the NAO index correlates negatively with snow depth in nearby Superior National Forest, USA⁹. In all regressions, degrees of freedom for tests of significance of independent variables were adjusted for autocorrelation⁹.

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A prominent feature of comparative life histories is the well documented negative correlation between growth rate and life span^{1,2}. Patterns of resource allocation during growth and reproduction reflect life-history differences between species^{1,2}. This is **particularly striking in tropical forests, where tree species can differ greatly in their rates of growth and ages of maturity but still** attain similar canopy sizes^{3,4}. Here we provide a theoretical frame**work for relating life-history variables to rates of production, d***M***/ d***t***, where** *M* **is above-ground mass and** *t* **is time. As metabolic rate** limits production as an individual grows, $dM/dt \propto M^{3/4}$. Incor**porating interspecific variation in resource allocation to wood density, we derive a universal growth law that quantitatively fits data for a large sample of tropical tree species with diverse life** histories. Combined with evolutionary life-history theory¹, the **growth law also predicts several qualitative features of tree demography and reproduction. This framework also provides a general quantitative answer to why relative growth rate** ($1/M$)(dM/df) decreases with increasing plant size ($\propto M^{-1/4}$) and **how it varies with differing allocation strategies5–8.**

Coexistence in diverse ecological communities has been thought to be due, in part, to life-history trade-offs involving allocation of resources^{6,9-13}. There is, however, no generally accepted mechanistic framework for understanding how patterns of allocation influence variation in life histories. Here we show that a general allometric growth model for trees can provide an explanation for much life-history variation. Metabolism produces the energy and materials that are used for all biological processes. A central issue of life history is how over ontogeny the products of metabolism are allocated among maintenance, growth and reproduction. Previous work has shown how the 3/4-power scaling of metabolic rate with body mass in both animals and plants results from physical and biological constraints on the distribution of resources through fractal-like vascular networks $14-16$.

In Box 1, we derive a production/growth law in terms of aboveground plant mass, *M*, or basal stem diameter, *D*. Our data set (Table 1, Fig. 1) consists of a 20-year change in *D* (*D*(20) compared with $D(0)$) for 45 species of trees in a tropical dry forest. In total, there were 2,283 individuals, all of reproductive age. Equation (6) predicts that $D^{2/3}(20) = A_i + D^{2/3}(0)$, so that a plot of $D^{2/3}(20)$ versus $D^{2/3}(0)$ for each species should yield a straight line with a universal slope of unity and an intercept, *Ai*, inversely proportional to wood density, ρ , so that A_i is proportional to ρ^{-1} . We tested this prediction by analysing the data for 45 species (Fig. 2a, Table 1). The average slope of all species, 1.04, is essentially not different from the predicted value of 1.0 (95% confidence intervals (CI): 1.01 to 1.08). Forty of the forty-five species had slopes statistically indistinguishable from the predicted value of 1 (Table 1). There was no systematic trend to deviate above or below 1; furthermore, we expect 5% (\approx 2.5 species) to differ by chance alone at the 0.05 level.

We conclude that production within species scales as $M^{3/4}$. There is, however, considerable interspecific variation in the intercepts. From Box 1, the intercepts should be inversely proportional to wood density, ρ , and this is confirmed by regression analysis $(r = -0.468$, exponent = -0.934 , 95% CI: -1.631 to -0.237 , intercept $= 0.6196$, $n = 29$). Weighting each species by multiplying $D^{2/3}$ by ρ collapses the production relationships for each species onto a line with a universal slope of unity and a common intercept (see Fig. 2b and equation (6)). Essentially, there is a trade-off in growth rate (change in stem diameter) with allocation to tissue density: species that allocate less biomass to their stems (light woods) increase in basal diameter faster than species that allocate more to stems (dense woods).

The trees depicted in Fig. 2a are reproducing, so some proportion of production, λ , must be allocated to reproduction rather than growth. The derivation in Box 1 (resulting in equation (7)) and measured empirical reproductive allometries both suggest that λ is independent of size within a species. To obtain the convergence of allometries implied in Fig. 2b requires the stronger constraint that λ be the same across species. Studies reporting relationships between production of reproduction mass and stem diameter indicate that whereas measured exponents usually approximate to the predicted value of 2, there are differences in normalization constants, and hence in λ , among species^{17–20}. Such variation will not change the predicted slope of unity when data are plotted as in Fig. 2a. If this variation in λ is independent of ρ , however, it will show up as residual variation in plots such as Fig. 2b.

Production data for seven species of temperate trees provide additional evidence that dM/dt is proportional to $M^{3/4}$. Instead of measuring change in diameter, Whittaker and Woodwell²¹ recorded total annual mass production of leaves and twigs, and bark and stem wood as a function of trunk diameter. Relating trunk diameter to annual production for twig and leaf mass, and wood and bark mass, shows that both scale with exponents essentially indistinguishable from the predicted $M^{3/4}$. So mass production in both tropical and temperature trees scales as *M*3/4.

Relative growth rate (1/*M*)(d*M*/d*t*) decreases with increases in individual size and varies across species^{$5-8$}. Although this decrease has been qualitatively attributed to several mechanisms⁶, a quantitative rule emerges as a direct consequence of equations (2) and (3): $(1/M)(dM/dt) \propto M^{-1/4}$. This rule is predicted to hold even across species growing in the same environment for comparisons when

Box 1

The growth law

As trees continue to increase in size throughout life, and this growth must be fuelled by metabolism, it is reasonable to assume that the growth rate (the rate at which its mass, *M*, increases over time), is directly proportional to metabolic rate, *B*, (the rate of gross photosynthesis). Thus, at any time *t*,

$$
\frac{dM}{dt} = C_{\rm G} B \tag{2}
$$

where C_G is a proportionality constant that can be time dependent. It has been shown that *B* is proportional to $V^{3/4}$, and stem diameter, *D*, is proportional to $V^{3/8}$, where *V* is the total volume of the plant¹⁴. If $\rho = (MV)$ is the tissue- or species-specific wood density, then, at any time *t*, these can be expressed as

$$
B = C_B \left(\frac{M}{\rho}\right)^{3/4} \qquad D = C_D \left(\frac{M}{\rho}\right)^{3/8} \tag{3}
$$

where C_B and C_D are corresponding proportionality constants. Implicit in these results is the assumption that the ratio E/ρ , where E is the Young's modulus of elasticity, is constant for all plants¹⁸. Here, we relax the restriction implicit in ref. 14 that ρ is constant and allow it to differ among plant species and to vary with time. Equations (2) and (3) can be combined to give

$$
\frac{dD}{dt} = \left(\frac{3C}{2\rho}\right)D^{1/3} \tag{4}
$$

where $C \equiv 1$ /4 $C_{\rm G}C_{\rm B}C_{\rm D}^{2/3}$. This can be integrated to give

$$
D(t)^{2/3} - D(t_0)^{2/3} = \int_{t_0}^t \frac{C(t)}{\rho(t)} dt
$$
 (5)

where the time dependence of all variables has been made explicit and the integration has been carried out from some initial time, t_0 , up to some arbitrary time, *t*. Thus, regardless of any possible time dependence of either the proportionality constants or the density, a plot of $D^{2/3}(t)$ versus $D^{2/3}(t_0)$ for fixed times *t* and t_0 for any species should yield a straight line with a universal slope of unity but with an intercept that depends on the time interval and the species. If, however, ρ of a given plant is taken to be independent of time, but allowed to vary across species, this can be re-expressed as

$$
\rho[D(t)^{2/3} - D(t_0)^{2/3}] = \int_{t_0}^t C(t) dt
$$
 (6)

The intercept of the production relationship should therefore be inversely proportional to ρ . If C_G , C_B , and C_D are independent of time then the intercept, namely the righthand side of equation (6), is given by $C(t - t_0)$. Furthermore, if *C* does not vary among species as implicitly assumed in ref. 14, and the time interval is the same for all species, then weighting $D^{2/3}(t)$ and $D^{2/3}(t_0)$ by ρ should collapse all species onto a universal line of unit slope. The intercept should no longer depend upon the species but only upon the time interval and *C* (however, see below for the reproductive period).

This framework allows us to recast the mass-production law as a function of basal stem diameter, *D*,

d*M*

$$
\frac{dM}{dt} = \left(\frac{C_{\rm G}C_B}{C_D^2}\right) D^2 \tag{7}
$$

This gives the new prediction that across species, d*M*/d*t*, for trees of the same diameter is explicitly independent of wood density ρ

The derivation so far has assumed that the plant is not reproducing, so that all production is given to growth. Growth must, however, slow with the onset of reproduction¹, as some fraction of production, (λ)dM/dt, is then devoted to reproduction rather than to individual growth, $(1 - \lambda)dM/dt$. The scaling rule for reproductive allocation and growth after the onset of reproduction depends upon the behaviour of λ . Life-history theory^{1,2} predicts the age (size) course of this reproductive allocation. The simplest model is to take λ as a constant both within and across species. A constant λ within species is supported by several studies $17-20$ that have measured allometries for the biomass of reproductive organs, $M_{\rm R}$, versus stem diameter, *D*, in both angiosperms and gymnosperms. M_R reflects the rate of allocation to reproduction. Since published studies show that M_R is generally proportional to D^2 , it follows from equation (7) that M_{R} is proportional to dM/dt indicating that λ is indeed approximately a constant within species and independent of size. This is an important result, as animals have λ increasing from 0 to 1 over the reproductive period^{1,2}. The fact that trees appear to have nearly constant values of λ probably reflects the constraints of resource harvesting and biomechanics on plant growth and development. Note that if λ is approximately constant, all of the previous growth results are true for all time periods $(t - t_0)$ after the onset of reproduction, with C_G redefined to be $(1 - \lambda)C_G$.

wood density is held constant.

The simplest life-history theory assumes determinate growth: at the onset of reproduction all production $(\lambda = 1)$ is given to reproduction and growth ceases¹. If we approximate plant growth

in this way and assume a pre-reproductive production rate $dM/dt = C_G B = C_G C_B (M/\rho)^{3/4}$ as in equations (2) and (3), and if mortality rate is independent of plant size (over a range which includes possible sizes at maturation), well known arguments¹ for

Figure 1 Representative variation of $D^{2/3}(0)$ versus $D^{2/3}(20)$ for six species of tropical trees. When sample sizes are large, the apparent variation is exaggerated because of the overlap of many points. Values for slopes and r^2 for these species are given in Table 1.

Figure 2 Growth rates of multiple species of tropical forest trees. **a**, $D^{2/3}(0)$ versus $D^{2/3}$ (20) for 45 species of dry forest trees. If production rate scales as $M^{3/4}$ (equation (2)), a slope of 1 (equation (5)) should be obtained. Note that all species have slopes that are essentially indistinguishable from 1, represented by the dashed line of unity. Statistics for each species are reported in Table 1. **b**, Weighting each species production

 $D(20)^{2/3} = A_1 + D(0)^{2/3}$ by wood density, ρ , collapses its production relationship onto a common line with unit slope and intercept A (see equation (6)). The dotted box shows the much greater range of variation represented in Fig. 2a. The coefficient of variation of variation for intercepts is reduced, as predicted, from 65.8 to 54.33.

Note that 95% confidence intervals for essentially all species include the predicted slope of 1 indicating that production $\alpha M^{3/4}$.

the optimal life history yield the approximate rule

$$
E_{\alpha} \approx \frac{(1.3)\rho^{3/4} M_{\alpha}^{1/4}}{C_{\rm G} C_B} \tag{1}
$$

for the average adult lifespan, E_α , is related to size at maturation, M_α . Equation (1) states that between species with similar M_{α} , E_{α} is proportional to $\rho^{3/4}$. If wood density is constant, this also implies that E_α is proportional to $M_\alpha^{1/4}$. Although we are not aware of any data sets precise enough to test this prediction, qualitative support is provided by several studies showing a generally positive relationship between wood density, lifespan and age to reproduction^{4,11,19,22-24}. In the future, one could relax the assumption $\lambda = 1$ and explore the general case.

The framework developed above highlights the central roles of allometric scaling and wood density in the life histories of trees. The tropical tree species studied here varied from fast-growing disturbance specialists with low-density wood and short life spans to slow growing emergent trees with dense wood and long life spans. Despite all of this variation, production scaled as *M*3/4, the same as in animals^{1,14}. Although all of the tree species produced new biomass at nearly the same rates (Fig. 2b), differences in wood density resulted in substantial differences in growth rates as measured as change in basal diameter (Fig. 2a). Wood density is frequently cited as being a 'principal determinant' of life history variation in woody plants^{11,19,22-26}. It is correlated with stem water storage and transport capacity, resistance to decay and leaf characteristics such as toughness and deciduousness^{4,11,1922-26}. Although the size of trees powerfully constrains rates of carbon

fixation, differences in carbon allocation strategies facilitate the coexistence of multiple species in tropical forests owing to variation in growth rate, lifespan and reproductive effort¹⁰.

We have shown that a general allometric framework incorporating the specifics of vascular transport and allocation can account for numerous features of biological diversity¹⁴⁻¹⁶. Because of the availability of data to evaluate the model, we have focused on scaling of growth and reproduction in trees; however, the model could easily be modified for plants with different growth forms (herbs) and life histories (monocarpy). Many details of plant anatomy, physiology, and ecology (for example, water and nutrient availability, plant density) can be incorporated into the allometric coefficients (the *C*s) to link pattern and process across multiple scales of biological organization. Interspecific variation in allocation to roots or defences should also be reflected in the *C* values. Production is ultimately limited by the physical and biological constraints that limit transportation through the vascular system. However, species differ in how they allocate production. These differences in rates of growth, life span, time until reproduction, wood density and other variables can be mechanistically linked to a general allometric framework of allocation. More importantly, this framework shows how size-dependent variation in life-history strategies can be derived from a set of general allometric principles. \Box

Methods

Study area

Measurements of diameters at breast height (dbh) of trees were recorded within a permanently marked study plot of seasonally dry tropical forest $(10^{\circ}$ 45' N, 85° 30' W)

within Sector Santa Rosa, Area de Conservacion, Guanacaste (ACG), of northwest Costa Rica²⁷. In 1976, all stems \geq 3 cm dbh were mapped within a continuous 680 m \times 240 m (16.32 Ha) area of forest²⁰ by S. P. Hubbell. Using an identical mapping protocol, a second remap of the San Emilio forest was completed between 1995 and 1996. In total, 46,833 individuals have been surveyed, 26,960 in 1976 and 19,873 in 1996. Together, the two surveys document 20 yr of growth and population change for about 150 species. The plot is composed of secondary growth forest and is heterogeneous with respect to age, topography and degree of deciduousness.

Calculation of individual tree growth

In 1976, most trees greater than 10 cm dbh were tagged with aluminum tree markers and given a unique identification number. Because few smaller individuals were given aluminum tags in 1976, tree growth was usually followed only for those trees greater than 10 cm dbh. Growth was calculated by monitoring changes in dbh for each individual. To ensure an accurate estimate of growth, a species was included only if a minimum representation of seven individuals had initial stem diameters \geq 10 cm, and the diameter range of all individuals \geq 20 cm. As the minimum diameter cut off for individuals was 10 cm, this imposed a minimum size range of 30 cm. Only individuals experiencing positive growth in the 20-year period were used for the calculation of allometric equations. In some cases, individuals experienced no change or even a decrease in diameter over time. This was usually due to partial death, loss of the main trunk or measuring errors. The 45 species meeting the above criteria are listed in Table 1. Production equations for each species were generated by plotting $D^{2/3}(0)$ versus $D^{2/3}(20)$ on linear axes. Because dbh was measured identically in 1976 and 1996, measurement error is likely to be equally distributed across the *x* and *y* axes. For these reasons, allometric slopes were determined using Model II RMA regression^{1,28,29}. Equations and statistics for each species are also reported in Table 1.

Species-specific wood density

The specific wood density, ρ , is a simple measure of the total dry mass per unit volume of wood (g cm⁻³). The specific density of wood is closely related to mechanical properties of strength, such as elastic moduli, which describe resistance to static and impact bending, compression and tension²⁸. For 29 of the 45 species reported in this study, values of specific wood density, ρ , in g cm⁻³, were taken from the literature^{24,26,30}. If more than one study reported a different value for a species, then the average value was used (Table 1).

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Optimizing the success of random searches

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We address the general question of what is the best statistical strategy to adapt in order to search efficiently for randomly located objects ('target sites'). It is often assumed in foraging theory that the flight lengths of a forager have a characteristic scale: from this assumption gaussian, Rayleigh and other classical distributions with well-defined variances have arisen. However, such theories cannot explain the long-tailed power-law distributions^{1,2} of flight lengths or flight times³⁻⁶ that are observed **experimentally. Here we study how the search efficiency depends on the probability distribution of flight lengths taken by a forager that can detect target sites only in its limited vicinity. We show that, when the target sites are sparse and can be visited any number of times, an inverse square power-law distribution of** flight lengths, corresponding to Lévy flight motion, is an optimal **strategy. We test the theory by analysing experimental foraging data on selected insect, mammal and bird species, and find that they are consistent with the predicted inverse square power-law distributions.**

Lévy flights are characterized by a distribution function

$$
P(l_j) \sim l_j^{-\mu} \tag{1}
$$

with $1 < \mu \leq 3$, where l_i is the flight length. The gaussian is the stable distribution for the special case $\mu \geq 3$ owing to the centrallimit theorem, while values $\mu \leq 1$ do not correspond to probability distributions that can be normalized². This generalization, equation (1) , introduces a natural parameter μ such that we essentially have a

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Correspondence and requests for materials should be addressed to M.G.R. (e-mail: mgr@indiana.bio.purdue.edu). Coordinates of the connector (accession number 1FOU) and pRNA (accession number 1FOQ) have been deposited with the Protein Data Bank.

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errata

Logical computation using algorithmic self-assembly of DNA triple-crossover molecules

Chengde Mao, Thomas H. LaBean, John H. Reif & Nadrian Seeman

Nature 407, 493-496 (2000).

In Fig. 2 of this paper, y_2 should have equalled 0 in Calculation 1; and x_2 , y_3 and y_4 should have equalled 0 in Calculation 2.

Acceleration of global warming due to carbon-cycle feedbacks in a coupled climate model

Peter M. Cox, Richard A. Betts, Chris D. Jones, Steven A. Spall & Ian J. Totterdell

Nature 408, 184-187 (2000).

In this paper, part of the labelling of Figs $2-4$ was printed incorrectly. In Fig. 2, the year 2950 should have been 2050. In Figs 2 and 3, the year 1050 should have been 1950. In Fig. 4, the numbers on the y axis should have been labelled sequentially from -200 to $+400$; in addition, in Fig. 4a the y axis should have been labelled 'Changes in vegetation carbon $(Gt C)$ '.

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correction

Allometric scaling of production and life-history variation in vascular plants

Brian J. Enquist, Geoffrey B. West, Eric L. Charnov & James H. Brown

Nature 401, 907-911 (1999).

In the last sentence of the abstract the relative growth rate equation, presented as $(1/M)$ (dM/df), should read $(1/M)$ (dM/dt). Also, the sentence following that containing equation (5) should read "Thus, regardless of any possible time dependence of either the proportionality constants or the density, a plot of $M^{1/4}$ versus $M^{1/4}_0$ for fixed times t and t_0 for any species should yield a straight line with a universal slope of unity but with an intercept that depends on the time interval and the species.''

In other words, the primary relationship is in terms of mass, M and not diameter, D. Equations (4) and (5), which relate the dependence of trunk diameter, D, on time, t, are only valid if the proportionality constant $A = C_D/\rho^{3/8}$ is time-independent; A is the proportionality constant in the allometric relation $D = AM^{3/8}$ of equation (3). This is needed because data are typically given in terms of D rather than M. The other proportionality constants, C_G and C_B , occurring, respectively, in the growth equation (2) and allometric equation for metabolic rate (3), can be time-dependent.

As our analysis of the data assumed time-independence of all of these coefficients as well as of wood density, ρ (see below equation (6)), this oversight does not affect our results or conclusions. If A were slowly varying, it would contribute a small correction to the unit slope prediction given by $(t - t_0)d$, where d is the logarithmic derivative of A. This is expected to be very small, especially as the time interval $(t - t_0)$ is small relative to the lifespan of the species sampled. Furthermore, any time dependence in A is almost certainly smaller than its variation across species and effects arising from neglecting maintenance costs in the growth equation (2). We thank J. Banavar, J. Damuth, A. Maritan and A. Rinaldo for bringing this oversight to our attention.