The dynamics of starvation and recovery

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6 The eco-evolutionary dynamics of species are fundamentally 64 ferently in broads during periods of resource scarcity compared 7 linked to the energetic constraints of its constituent individuals. 65 to normal periods (9, 10), sometimes delaying or even foregoing s Of particular importance are the tradeoffs between reproduction of a breeding season (1, 11, 12). Even fresh-9 and the dynamics of starvation and recovery in resource-limited 10 environments. To elucidate the consequences of this tradeoff, 11 we introduce a minimal nutritional state-structured model that in-12 corporates two classes of consumer: nutritionally replete con-13 sumers that reproduce, and undernourished, non-reproducing 14 consumers that are susceptible to mortality. As a function of 15 the transition rates between these two states that are determined 72 migratory breeding grounds to reproduce after one or multiple 16 by the abundance of resources, the consumer populations can ei- 73 seasons in resource-rich environments where they accumulate 17 ther undergo cyclic dynamics or reach a steady state. We obtain 74 nutritional reserves (14-16). 18 strong constraints on starvation and recovery rates by deriving 19 allometric scaling relationships between body size and a variety 20 of traits and find that population dynamics subject to these constraints are typically driven to a steady state. Moreover, we find 22 that these rates fall within a 'refuge' in parameter space, where 23 the probability of extinction of the consumer population is min-24 imized. Thus we identify a potential mechanism that may both 25 drive and constrain the dynamics of animal populations. Our 81 other species (including humans) suffer irregular menstrual cy-26 model provides a natural framework that predicts maximum body 82 cling and higher abortion rates during periods of nutritional 27 size for mammals by determining the relative stability of an oth- 83 stress (19, 20). In the extreme case of unicellular organisms, erwise homogeneous population to a mutant population with al- $_{29}$ tered percentage of body fat. For body masses $< 1.748 \times 10^7$ g, 30 individuals with increased energetic reserves can invade resident $_{31}$ populations, and vice versa for body mass $> 1.748 \times 10^7 \mathrm{g}$, thus 32 providing a principled mechanism for a within-lineage driver of 33 Cope's rule.

34 foraging | starvation | reproduction

35 Significance Statement Energetic investment in somatic mainte-36 nance and growth vs. reproduction directly impacts the dynamics of 37 populations among species. Here, we construct a Nutritional State-38 structured Model (NSM) to assess the population-level effects of star-39 vation and recovery of a consumer population in a resource-limited en-40 vironment, and use allometric scaling relationships for mammals to es-41 tablish all timescales and rates. Our model: i. reveals that mammalian 42 energetic rates minimize the probability of stochastic extinction, ii. es-43 tablishes dynamic bounds on mammalian body size while providing 44 independent theoretical support for the energy equivalence hypothesis, 45 and iii. provides a mechanistic driver for the evolutionary trend towards 46 larger body size known as Cope's rule.

47 Introduction

51 generally manifested as tradeoffs between investing in somatic 52 maintenance and growth, or allocating energy towards repro-53 duction (1-3). The timing of these behaviors responds to se-54 lective pressure, as the choice of the investment impacts future 55 fitness (4–6). The influence of resource limitation on an or-56 ganism's ability to maintain its nutritional stores may lead to 57 repeated delays or shifts in reproduction over the course of an

The balance between (a) somatic growth and maintenance, 60 and (b) reproduction depends on resource availability (7). For 61 example, reindeer invest less in calves born after harsh winters 62 (when the mother's energetic state is depleted) than in calves 63 born after moderate winters (8). Many bird species invest dif-

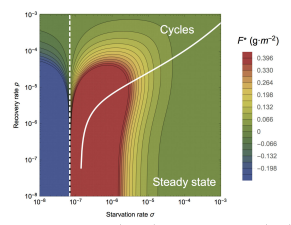
67 water and marine zooplankton have been observed to avoid 68 reproduction under nutritional stress (13), and those that do 69 reproduce have lower survival rates (2). Organisms may also 70 separate maintenance and growth from reproduction over space 71 and time: many salmonids, birds, and some mammals return to

Physiology also plays an important role in regulating repro-76 ductive expenditures during periods of resource limitation. The 77 data collected thus far has shown that diverse mammals (47 78 species in 10 families) exhibit delayed implantation, whereby 79 females postpone fetal development (blastocyst implantation) so until nutritional reserves can be accumulated (17, 18). Many 84 nutrition is unavoidably linked to reproduction because the nu-85 tritional state of the cell regulates all aspects of the cell cycle 86 (21). The existence of so many independently evolved mecha-87 nisms across such a diverse suite of organisms highlights the im-88 portance and universality of the fundamental tradeoff between 89 somatic and reproductive investment. However the general dy-90 namic implications of these constraints are unknown.

Though straightforward conceptually, incorporating the en-92 ergetic dynamics of individuals (22) into a population-level 93 framework (22, 23) presents numerous mathematical obsta-94 cles (24). An alternative approach involves modeling the 95 macroscale relations that guide somatic versus reproductive 96 investment in a consumer-resource system. For example, 97 macroscale Lotka-Volterra models assume that the growth rate 98 of the consumer population depends on resource density, thus 99 implicitly incorporating the requirement of resource availability 100 for reproduction (25).

In this work, we adopt an alternative approach in which we 102 explicitly account for resource limitation and the subsequent 103 effects of starvation. Namely, only individuals with sufficient 104 energetic reserves can reproduce. Such a constraint leads to 105 reproductive time lags due to some members of the population 48 The behavioral ecology of all organisms is influenced by the en- 106 going hungry and then recovering. Additionally, we incorporate 49 ergetic state of individuals, which directly influences how they 107 the idea that reproduction is strongly constrained allometrically 50 invest reserves in uncertain environments. Such behaviors are 108 (3), and is not generally linearly related to resource density. As

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sumer. These bifurcation conditions separate parameter space into unphysical, cyclic, and steady state dynamic regimes. The colors show the steady state densities for the energetically replete consumers F^* , (warmer colors denote higher densities).

109 we shall show, these constraints influence the ensuing popula-110 tion dynamics in dramatic ways.

112 Nutritional state-structured model (NSM)

113 We begin by defining a minimal Nutritional State-structured 114 population Model (NSM), where the consumer population is 115 partitioned into two states: (a) an energetically replete (full) state F, where the consumer reproduces at a constant rate λ_{160} where $A = (\lambda \xi \rho + \mu \sigma)$ and $B = (\beta \mu \xi + \delta \lambda \xi - \lambda \mu)$. The states 125 is partitioned between maintenance and growth purposes. In 169 and consumer densities are both nonzero. 126 the supplementary information (SI) we provide a fully mecha-127 nistic model for each of these dynamics and constants, and show 171 the reproduction rate λ must be less than the starvation rate 128 that the system produces a simple non-dimensional form which we describe below.

Consumers transition from the full state F to the hungry 131 state H at a rate σ —the starvation rate—and also in proportion to the absence of resources (1-R). Conversely, consumers 133 recover from state H to state F at rate $\xi \rho$ and in proportion to 134 R, where ξ represents a ratio between maximal resource con-135 sumption and the carrying capacity of the resource. Resources are eaten by the hungry consumers at rate $\rho R + \delta$, that accounts 137 for their somatic growth (ρR) and maintenance (δ) . Full con-138 sumers eat resources at a constant rate β that accounts for max-139 imal maintenance and somatic growth (see SI for mechanistic 140 derivations of these rates from resource energetics). The NSM 141 represents an ecologically motivated fundamental extension of 142 the idealized starving random walk model of foraging, which 143 focuses on resource depletion, to include reproduction and re-144 source replenishment (26–28), and is a more general formulation than previous models incorporating starvation (29).

In the mean-field approximation, in which the consumers 147 and resources are perfectly mixed, their densities evolve accord-148 ing to the rate equations

$$\frac{dF}{dt} = \lambda F + \xi \rho R H - \sigma (1 - R) F,$$

$$\frac{dH}{dt} = \sigma (1 - R) F - \xi \rho R H - \mu H,$$

$$\frac{dR}{dt} = \alpha (1 - R) R - (\rho R + \delta) H - \beta F$$
[1]

This system of nondimensional equations follows from a 150 set of first-principle relationships for resource consumption and $_{151}$ growth (see SI for a full derivation and the dimensional form). Notice that the total consumer density F+H evolves accordate in the total consumer density F+H evolves accordate in the density in the classic Lotka-Volterra motion for the predator density in the classic Lotka-Volterra 155 model (30), except that the resource density does not appear in 156 the growth term. As discussed above, the attributes of repro-Fig. 1: The transcritical (dashed) and Hopf bifurcation (solid) as a 157 duction and mortality have been explicitly apportioned to the function of the starvation rate σ and recovery rate ρ for a 100g con- 158 full and hungry consumers, respectively, so that the growth in

> Equation [1] has three fixed points: two trivial fixed points at $(F^*, H^*, R^*) = (0, 0, 0)$ and (0, 0, 1), and one non-trivial, internal fixed point at

$$F^* = (\sigma - \lambda) \frac{\alpha \lambda \mu^2 (\mu + \xi \rho)}{A(\lambda \rho B + \mu \sigma (\beta \mu + \lambda (\delta + \rho)))},$$

$$H^* = (\sigma - \lambda) \frac{\alpha \lambda^2 \mu (\mu + \xi \rho)}{A(\lambda \rho B + \mu \sigma (\beta \mu + \lambda (\delta + \rho)))},$$

$$R^* = (\sigma - \lambda) \frac{\mu}{A}.$$
[2]

116 state F, where the consumer reproduces at a constant rate Λ 160 where $A = (\Lambda \xi \rho + \mu \sigma)$ and $D = (\rho \mu \xi + \sigma \lambda \xi - \Lambda \mu)$. The sum of this fixed point is determined by the Jacobian matrix 118 cient (hungry) state H, where the consumer does not reproduce 162 \mathbf{J} , where each matrix element $J_{ij} = \partial \dot{X}_i / \partial X_j$ when evaluated 119 but dies by starvation at rate μ . The underlying resource R 163 at the internal fixed point, and \mathbf{X} is the vector (F, H, R). The 120 evolves by logistic growth with an intrinsic growth rate α and 164 parameters in Eq. [1] are such that the real part of the largest 165 consumer conscience. 121 a carrying capacity C. The rate at which consumers transition 122 between states and consume resources is dependent on their 123 overall abundance, the abundance of resources, the efficiency of 124 converting resources into metabolism, and how that metabolism 125 converting resources into metabolism, and how that metabolism 126 respect to small perturbations from the fixed point. Because 127 this fixed point is unique, it is the global attractor for all populations resources for any initial condition where the resource 128 resulting populations and between meiabolism 129 resulting populations and populations are supplied by the population of the parameters in Eq. [1] are such that the lear part of the largest 126 eigenvalue of J is negative, so that the system is stable with 127 respect to small perturbations from the fixed point. Because 128 resulting populations are particular to the largest 129 respect to small perturbations from the fixed point. Because 129 resulting populations are particular to the largest 129 respect to small perturbations from the fixed point. Because 129 respect to small perturbations from the fixed point. Because 129 respect to small perturbations from the fixed point. Because 129 respect to small perturbations from the fixed point. Because 129 respect to small perturbations from the fixed point. Because 129 respect to small perturbations from the fixed point. Because 129 respect to small perturbations from the fixed point is unique, it is the global attractor for all populations from the fixed point is unique, it is the global attractor for all populations from the fixed point is unique, it is the global attractor for all populations from the fixed point is unique.

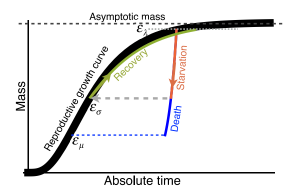


Fig. 2: The growth trajectory over absolute time of an individual organism as a function of body mass. Initial growth follows the black trajectory to an energetically replete reproductive adult mass $m = \epsilon_{\lambda} M$. Starvation follows the red trajectory to $m = \epsilon_{\sigma} \epsilon_{\lambda} M$. Recovery follows the green curve to the replete adult mass, where this trajectory differs from the original growth because only fat is being regrown which requires a longer time to reach $\epsilon_{\lambda}M$. Alternatively, death from starvation follows the blue trajectory to $m = \epsilon_{\mu} \epsilon_{\lambda} M$.

175 regime where all steady-state densities become negative after in- 242 general solution (3, 46) 176 tersecting the trivial fixed point $(F^*, H^*, R^*) = (0, 0, 0)$. The 177 biological implication of the constraint $\lambda < \sigma$ has a simple 178 interpretation—the rate at which a macroscopic organism loses mass due to lack of resources is generally much faster than the where, for $\eta < 1$, $M = (B_0/B_m)^{1/(1-\eta)}$ is the asymptotic mass, 182 isms within empirically observed body size ranges.

In the physical regime of $\lambda < \sigma$, the fixed point [2] may ei-183 184 ther be a stable node or a limit cycle (Fig.). In continuous-time $\frac{247}{248}$ it takes to reach a particular mass ϵM is given by the timescale 185 systems, a limit cycle arises when a pair of complex conjugate 186 eigenvalues crosses the imaginary axis to attain positive real 187 parts (32). This Hopf bifurcation is defined by $Det(\mathbf{S}) = 0$, with 188 S the Sylvester matrix, which is composed of the coefficients of 249 where we will define values of ϵ to describe a set of rates within 189 the characteristic polynomial of the Jacobian matrix (33). As 250 our model. For the time to reproduce, $t_{\lambda} = \tau(\epsilon_{\lambda})$, where ϵ_{λ} $_{195}$ can increase the extinction risk (35–37).

When the starvation rate $\sigma \gg \lambda$, a substantial fraction the consumers are driven to the hungry non-reproducing the rate of the 196 when the stativation recording to the hungry non-reproducing 257 The race of recording 258 acrues sufficient tissue to transition from the hungry to the 198 state. Because reproduction is inhibited, there is a low steady-199 state consumer density and a high steady-state resource den-208 increases with higher recovery rates.

210 Role of allometry

209

211 While there are no a priori constraints on the parameters in 213 the parameter space. Here we use allometric scaling relations 214 to constrain the covariation of rates in a principled and biologi-215 cally meaningful manner. Allometric scaling relations highlight 216 common constraints and average trends across large ranges in 217 body size and species diversity. Many of these relations can be 218 derived from a small set of assumptions and below we describe $a' = B_0/E'_m$ accounts for possible deviations in the 222 regime of dynamics occupied by the entire class of mammals, 279 the body fat scaling in our framework. 223 along with the key differences between the largest and smallest 280 224 mammals.

226 by consumer metabolism, which can be used to describe a vari- 283 production is impossible. For starving individuals we assume 227 ety of organismal features (39). The scaling relation between an 284 that an organism must meet its maintenance requirements by $_{228}$ organism's metabolic rate B and its body mass M at reproduc- $_{285}$ using the digestion of existing mass as the sole energy source. 229 tive maturity is known to scale as $B=B_0M^\eta$ (40), where the 286 This assumption implies the following simple metabolic balance 230 scaling exponent η is typically close to 2/3 or 3/4 for metazoans 287 $\dot{m}E'_m=-B_mm$ or $\dot{m}=-a'm/M^{1-\eta}$ where E'_m is the amount 231 (e.g., ref. 39), and has taxonomic shifts for unicellular species 288 of energy stored in a unit of existing body mass, which differs

234 tween growth and maintenance purposes can be used to derive 291 ergy balance prescribes the mass trajectory of a non-consuming 235 a general equation for both the growth trajectories and growth $_{292}$ organism: $m(t) = Me^{-a't/M^{1-\eta}}$. The timescale for starvation 236 rates of organisms ranging from bacteria to metazoans (3, 42^- 293 is given by the time it takes m(t) to reach $\epsilon_{\sigma}M$, which gives 237 45). This relation is derived from the simple balance condition $_{238} B_0 m^{\eta} = E_m \dot{m} + B_m m$, (3, 42–45) where E_m is the energy

 $_{172}$ σ , so that the consumer and resource densities are positive. $_{239}$ needed to synthesize a unit of mass, B_m is the metabolic rate ₁₇₃ The condition $\sigma = \lambda$ thus represents a transcritical (TC) bi-₂₄₀ to support an existing unit of mass, and m is the mass of the 174 furcation (31) that demarcates a physical from an unphysical 241 organism at any point in its development. This balance has the

$$\left(\frac{m(t)}{M}\right)^{1-\eta} = 1 - \left[1 - \left(\frac{m_0}{M}\right)^{1-\eta}\right] e^{-a(1-\eta)t/M^{1-\eta}},$$
 [3]

180 rate of reproduction. As we will discuss below, this inequality $\frac{243}{244} = B_0/E_m$, and m_0 is mass at birth, itself varying allometri-181 is a natural consequence of allometric constraints (3) for organ- $\frac{244}{245}$ cally (see SI). We now use this solution to define the timescale 246 for reproduction and recovery from starvation (Fig. ; see (43) 247 for a detailed presentation of these timescales). The time that

$$\tau\left(\epsilon\right) = \ln\left[\frac{1 - (m_0/M)^{1-\eta}}{1 - \epsilon^{1-\eta}}\right] \frac{M^{1-\eta}}{a\left(1 - \eta\right)},\tag{4}$$

the characteristic polynomial of the Jacobian matrix (60). The system parameters are tuned to be within the stable regime, but close to the Hopf bifurcation, the amplitude of the transported signs constantly being perturbed (34), the onset of transient cycles, even though they decay with time in the mean-field description, risk (35–37)

250 our model. For the time to reproduce, $\iota_{\lambda} - \iota_{\lambda}(\epsilon_{\lambda})$, where ϵ_{λ} is the fraction of the asymptotic mass where an organism is 252 reproductively mature and should be close to one (typically where v is the number of offspring produced, and for any constantly being perturbed (34), the onset of transient cycles, 254 where v is the number of offspring produced, and for any constantly being perturbed (35–37)

The rate of recovery $\rho = 1/t_{\rho}$ requires that an organism 259 full state. Since only certain tissues can be digested for energy state consumer density and a high steady-state resource den-200 sity. However, if $\sigma/\lambda \to 1$ from above, the population is 261 we define the rates for starvation, death, and recovery by the 200 Sity. However, if $0/N \to 1$ from above, the F-F and above, the F-F and above, the representation of the representation of the representations and resource densities (Fig.). If the starvation rate is low and resource densities (Fig.). If the starvation rate is low and resource densities (Fig. 201) and resource densities (Fig. 202) and resource densities (Fig. 202). We define $m_{\sigma} = \epsilon_{\sigma} M$, where $\epsilon_{\sigma} < 1$ is the fraction fraction. 204 enough that the Hopf bifurcation is crossed, these oscillations
264 IZALIONS). We define mass where reproduction ceases. This fraction
265 of replete-state mass where reproduction ceases. become stable over time. This threshold occurs at higher values the range of parameter space giving rise to cyclic dynamics also the range of parameter space giving rise to cyclic dynamics also the range of parameter space giving rise to cyclic dynamics also the range of parameter space giving rise to cyclic dynamics also the range of parameter space giving rise to cyclic dynamics also the range of parameter space giving rise to cyclic dynamics also the range of parameter space giving rise to cyclic dynamics also the range of parameter space giving rise to cyclic dynamics also the range of parameter space giving rise to cyclic dynamics also the range of parameter space giving rise to cyclic dynamics also the range of parameter space giving rise to cyclic dynamics also the range of parameter space giving rise to cyclic dynamics also the range of parameter space giving rise to cyclic dynamics also the range of parameter space giving rise to cyclic dynamics also the range of parameter space giving rise to cyclic dynamics also the range of parameter space giving rise to cyclic dynamics also the range of parameter space giving rise to cyclic dynamics also the range of parameter space giving rise to cyclic dynamics also the range of parameter space giving rise to cyclic dynamics also the range of parameter space giving rise to cyclic dynamics also the range of parameter space giving rise to cyclic dynamics also the range of parameter space giving rise to cyclic dynamics also the range of parameters are range. The range of parameters are range of parameters are range of parameters are range of parameters are range. The range of parameters are range. The range of parameters are range. The range of parameters are range. The range of par 269 according to $M_{\rm fat} = f_0 M^{\gamma}$ and assuming that once this mass 270 is fully digested the organism starves, this would imply that ₂₇₁ $\epsilon_{\sigma} = 1 - f_0 M^{\gamma}/M$. It follows that the recovery timescale, t_{ρ} , While there are no a priori constraints on the parameters in the NSM, most organisms correspond to restricted portions of the NSM, most organisms correspond to restricted portions of the parameter charge. Here we use allowed a scaling relations [273] Eqs. [3] and [4] this timescale is given by simply considering 274 an adjusted starting mass of $m'_0 = \epsilon_{\sigma} \epsilon_{\lambda} M$, in which case

$$t_{\rho} = \ln \left[\frac{1 - (\epsilon_{\sigma} \epsilon_{\lambda})^{1 - \eta}}{1 - \epsilon_{\lambda}^{1 - \eta}} \right] \frac{M^{1 - \eta}}{a'(1 - \eta)}$$
 [5]

219 our framework to determine the covariation of timescales and 276 biosynthetic energetics during recovery (see SI). It should be 220 rates across the range of mammals for each of the key parame- 277 noted that more complicated ontogenetic models explicitly han-221 ters of our model (cf. ref. 38). We are thereby able to define the 278 dle storage (45), whereas this feature is implicitly covered by

To determine the starvation rate, σ , we are interested in the 281 time required for an organism to go from a mature adult that Nearly all of the rates described in the NSM are determined $_{282}$ reproduces at rate λ , to a reduced-mass hungry state where rebetween $\eta \approx 1$ in eukaryotes and $\eta \approx 1.76$ in bacteria (3, 41). 289 from E_m , the energy required to synthesis a unit of biomass Several efforts have shown how a partitioning of B be-290 (45). Given the replete mass, M, of an organism, the above en-

$$t_{\sigma} = -\frac{M^{1-\eta}}{a'} \ln \left(\epsilon_{\sigma} \right).$$
 [6]

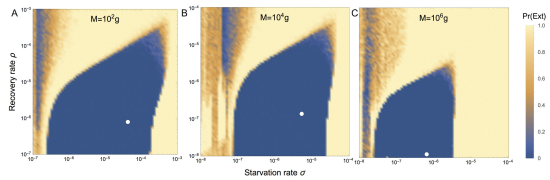


Fig. 3: Probability of extinction for a consumer with (A) $M = 10^2$ g, (B) $M = 10^4$ g, and (C) $M = 10^6$ g as a function of the starvation rate σ and recovery rate ρ , where the initial density is given as (XF^*, XH^*, R^*) , where X is a random uniform variable in [0, 2]. Note the change in scale for $M=10^4$ and $M=10^6$ g. Extinction is defined as the population trajectory falling below $0.2\times$ the allometrically constrained steady state. The white points denote the allometrically constrained starvation and recovery rate.

The starvation rate is then $\sigma=1/t_\sigma$, which scales with replete- 339 state region at some distance from the Hopf bifurcation. This 295 state mass as $1/M^{1-\eta} \ln{(1-f_0M^\gamma/M)}$. An important feature 340 result suggests that cyclic population dynamics should be rare, $_{296}$ is that σ does not have a simple scaling dependence on λ (Fig. $_{341}$ particularly in environments where resources are limiting. ??), which is important for the dynamics that we later discuss. 342 The time to death should follow a similar relation, but de- 343 Extinction risk 299 fined by a lower fraction of replete-state mass, $m_{\mu}=\epsilon_{\mu}M$ where 344 Within our model, higher rates of starvation result in a larger $_{300}$ $\epsilon_{\mu} < \epsilon_{\sigma}$. Suppose, for example, that an organism dies once it 345 flux of the population to the hungry state. In this state, repro-301 has digested all fat and muscle tissues, and that muscle tissue 346 duction is absent, thus increasing the likelihood of extinction. scales with body mass according to $M_{\rm musc} = u_0 M^{\zeta}$. This gives 347 From the perspective of population survival, it is the rate of $_{303}$ $\epsilon_{\mu}=1-\left(f_{0}M^{\gamma}+u_{0}M^{\zeta}\right)/M$. Muscle mass has been shown $_{348}$ starvation relative to the rate of recovery that determines the 304 to be roughly proportional to body mass (47) in mammals and 349 long-term dynamics of the various species (Fig.). We therefore ϵ_{μ} is merely ϵ_{σ} minus a constant. The time to go from ϵ_{μ} is merely ϵ_{σ} minus a constant. The time to go from ϵ_{μ} is merely ϵ_{σ} minus a constant. 306 starvation to death is the total time to reach $\epsilon_{\mu}M$ minus the 351 steady-state density on extinction risk, both as functions of σ 307 time to starve, or

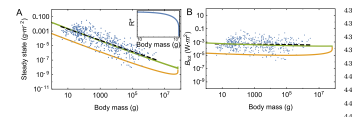
$$t_{\mu} = -\frac{M^{1-\eta}}{a'} \ln\left(\epsilon_{\mu}\right) - t_{\sigma}, \qquad [7]$$

313 10^7 g (the early Oligocene Indricotheriinae and the Miocene 361 we assessed extinction risk across a range of values for σ and ρ 362 between ca. 10^{-7} to 10^{-3} . As expected, higher rates of extinction risk across a range of values for σ and ρ 362 between ca. 10^{-7} to 10^{-3} . As expected, higher rates of extinction risk across a range of values for σ and ρ 362 between ca. 10^{-7} to 10^{-3} . As expected, higher rates of extinction risk across a range of values for σ and ρ 362 between ca. 10^{-7} to 10^{-3} . As expected, higher rates of extinction risk across a range of values for σ and ρ 362 between ca. 10^{-7} to 10^{-3} . 315 simply involve altering the metabolic exponents and scalings $\frac{1}{2}$ simply involve altering the metabolic exponents and scalings $\frac{1}{2}$ simply involve altering the metabolic exponents and scalings 316 associate with ϵ . Moreover, we emphasize that our allometric $\frac{364}{2}$ values of ρ if σ is small. For low values of σ and high values 317 equations describe mean relationships, and do not account for 365 of ρ , the increased extinction risk results from transient cycles the (sometimes considerable) variance associated with individ
366 with larger amplitudes as the system nears the Hopf bifurcation 319 ual species.

321 Stabilizing effects of allometric constraints

323 vation and recovery rates are not independent parameters, and σ and ρ , extinction probabilities are minimized. We find that the allometrically constrained values of σ and which is constrained via covarying parameters. Given the parameters of targetical and the matter of targetical and targetical and the matter of targetical and targetical a 326 rameters of terrestrial endotherms, we find that the starvation 374 M (Fig. 3A-C, white points). These values are close enough 327 rate σ and the recovery rate ρ are constrained to lie within a 375 to the Hopf bifurcation to avoid low steady-state densities, and 328 small region of potential values (Fig. ??) for the known range 376 far enough away to avoid large-amplitude transient cycles. The 379 of body sizes M. We thus find that the dynamics for all mammalian body sizes are confined to the steady-state regime of the malian NSM and that limit-cycle behavior is precluded. Incorporations are described by the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malian body sizes are confined to the steady-state regime of the malia 332 ing uncertainty in allometric parameters (20% variation around 380 drive starvation and recovery rates within populations. Such a 381 mechanism would select for organism physiology that generates 382 appropriate σ and ρ values that serve to minimize extinction 385 that small mammals are marginally 1. that small mammals are marginally less prone to population 383 risk. This selection could occur via the tuning of body fat peroscillations—both stable limit cycles and transient cycles—than mammals with larger body size. However, starvation and respectively rates across all values of M fall squarely in the steady section risk parameter space becomes smaller (Fig. 3A-C),

 $_{352}$ and ρ . To this end, we computed the probability of extinction, 353 where we define extinction as a population trajectory falling [7] $^{354}_{355}$ below one fifth of the allometrically constrained steady state $^{355}_{355}$ at any time between $t=10^8$ and $t=10^{10}$. This procedure is 356 repeated for 50 replicates of the continuous-time system shown Although the rate equations [1] are general, here we focus 357 in Eq. 1 for organisms with mass ranging from 10^2 to 10^6 on parameterizations for terrestrial-bound endotherms, specifically mammals, which range from a minimum of $M \approx 1_{\rm g}$ 359 (XF^*, XH^*, R^*), with X a random variable that is uniformly the Etruscan shrew Suncus etruscus) to a maximum of $M \approx 1_{\rm g}$ 360 distributed in [0, 2]. By allowing the rate of starvation to vary, 361 we assessed extinction risk across a range of values for $x = 1_{\rm g}$. ₃₆₇ (Fig. 3). For high values of σ and low values of ρ , increased 368 extinction risk arises because of the decrease in the steady-state $_{369}$ consumer population density (Figs. B, 3). This interplay cre-322 As the allometric derivations of the NSM rate laws reveal, star- 370 ates an 'extinction refuge', such that for a constrained range of



Refs. 42, 43, 45).

388 more sensitive to smaller changes in physiological rates control- 454 predicted upper bound depend only on the scaling of body com-389 ling starvation and recovery. To summarize, our finding that 455 position and are independent of the resource parameters. We 390 the allometrically-determined parameters fall within this low 456 also note that the prediction of an asymptotic limit on mam-391 extinction probability region suggests that the NSM dynamics 457 malian size parallels work on microbial life where an upper and 392 may both drive—and constrain—natural animal populations.

394 Dynamic and energetic barriers to body size

 $_{\rm 395}$ Metabolite transport constraints are widely thought to place $^{\rm 461}$ 407 eages show a significant trend towards larger body size (known 473 which larger body sizes are selected for. 408 as Cope's rule) (59–62), and it is thought that within-lineage 474 extinction risk for large taxa over longer timescales (60). These 477 are entirely composed of metabolic reserves, and this occurs 412 trends are thought to be driven by a combination of climate 413 change and niche availability (62); however the underpinning 414 energetic costs and benefits of larger body sizes, and how they 415 influence dynamics over ecological timescales, have not been ex-416 plored. We argue that the NSM provides a suitable framework 417 to explore these issues.

The NSM correctly predicts that species with smaller $_{419}$ masses have larger steady-state population densities (Fig. 3A). 420 Moreover, we show that the NSM provides independent the-421 oretical support for the energy equivalence hypothesis and 422 Damuth's Law (63-65). The energy equivalence hypothesis ar-423 gues that the total energy use, B_{tot} , of a population is constant 424 independent of species size (63–65). This hypothesis is based on 425 observations showing that the steady state abundance, N^* , of a ⁴²⁶ species is proportional to the inverse of individual metabolism, ⁴²⁷ such that $N^* \propto M^{-3/4}/B_0$ (64, 65). This relationship im-⁴²⁸ plies that $B_{\rm tot} = N^*B(M) = Q$, where Q is a constant, and 429 has been shown to hold in both mammalian and vascular plant 430 communities (63–65). Figure 3A shows that both F^* and H^* $_{431}$ scale as $M^{-\eta}$ over a wide range of organism sizes and Figure 3B $_{432}$ shows that F^*B is nearly constant over this same range. This 433 result is remarkable because it illustrates that the steady state 434 values of the NSM combined with the derived timescales natu-

435 rally give rise to energy equivalence. Our model shows that the 436 equivalence breaks down at the maximum observed body sizes 437 for mammals, suggesting that this maximum is a hard limit 438 where deviations outside of this range are energetically subop-439 timal. In the framework of our model, the total metabolic rate $_{440}$ of F and H becomes infinite at a finite mass, and occur at the 441 same scale where the steady state resources vanish (Fig. 3). 442 This asymptotic behavior is governed by body sizes at which 443 ϵ_{μ} and ϵ_{λ} equal zero causing the timescales to become infinite $_{444}$ (see Equation 7) and the rates μ and λ to equal zero. The Fig. 4: (A) Consumer steady states F^* (green) and H^* (orange) as a $_{445}$ $\mu = 0$ asymptote occurs first when $f_0 M^{\gamma-1} + u_0 M^{\zeta-1} = 1$, function of body mass. Inset: Resource steady state R^* as a function of 446 and corresponds to $(F^*, H^*, R^*) = (0, 0, 0)$. This point preconsumer body mass. (B) Total energetic use B_{tot} of consumer popula₄₄₇ dicts a strong upper bound on mammalian body size and octions at the steady state as a function of body mass. The data are from Damuth (63) and have been converted to total population metabolism using the allometric relationships for metabolic rate (please see SI and 449 determined by the population-level consequences of energetic 450 constraints, is close to the maximum body size observed in the 451 North American mammalian fossil record (59), as well as the 452 mass predicted from an evolutionary model of body size evolu-387 suggesting that the population dynamics of larger organisms are 453 tion (60). It should be noted that the asymptotic behavior and 458 lower bound on bacterial size, and an upper bound on single cell 459 eukaryotic size, is predicted from similar growth and energetic 460 scaling relationships (3, 66).

We contend that the NSM provides a mechanistic under-396 strict boundaries on biological scaling (39, 55, 56) and thereby 462 standing of the energetic dynamics that give rise to both ob-397 lead to specific predictions on the minimum possible body size 463 served limitations on mammalian body size, as well as the ob-398 for organisms (57). Above this bound, a number of energetic 464 served trend towards larger body size over evolutionary time. and evolutionary mechanisms have been explored to assess the 465 The NSM predicts that the steady state resource density R^* 400 costs and benefits associated with larger body masses, partic-466 decreases with increasing body size of the consumer popula- $_{401}$ ularly for mammals. One important such example is the fast- $_{467}$ tion (Fig. 3C), and classic resource competition theory predicts 402 ing endurance hypothesis, which contends that larger body size, 468 that the species surviving on the lowest resource abundance will with consequent lower metabolic rates and increased ability to 469 outcompete others (67–69). Thus, the combined NSM steady-404 maintain more endogenous energetic reserves, may buffer or- 470 state dynamics and allometric timescales predict that larger 405 ganisms against environmental fluctuations in resource avail-471 mammals have an intrinsic competitive advantage given a com-406 ability (58). Over evolutionary time, terrestrial mammalian lin- 472 mon resource, but does not offer a within-lineage mechanism by

To examine whether the NSM could provide such a mecha-409 drivers generate selection towards an optimal upper bound of 475 nism, we begin by noting that a theoretical upper bound on 410 roughly 10^7 grams (59), a value that is likely limited by higher 476 mammalian body size is given by $\epsilon_{\sigma}=0$, where mammals

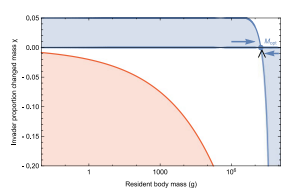


Fig. 5: Invasion feasibility for organisms with a proportional change in mass χ against a population with a resident body mass M. The blue region denotes proportions of modified mass χ resulting in successful invasion. The red region denotes values of χ that result in a mass that is below the starvation threshold and are thus infeasible. Arrows point to the predicted optimal mass from our model $M_{\rm opt} = 1.748 \times 10^7$ which may serve as an evolutionary attractor for body mass. The black wedge points to the largest body mass known for terrestrial mammals (Deinotherium spp.) at $1.74 \times 10^7 \text{g}$ (61).

479 phant. Next we examine to what extent a more realistic upper 546 associated with the dynamics of starvation and recovery. 480 bound to body mass may serve as an evolutionary attractor, 481 thus providing a suitable within-lineage mechanism for Cope's 482 rule. We directly assess the susceptibility of an otherwise ho-483 mogeneous population to invasion by a mutated subset of the 547 References 484 population (denoted by ') where individuals have a modified $^{548}_{549}$ 485 proportion of body fat $M'=M(1+\chi).$ For the allowable values 550 $_{486}$ of χ the adjusted mass should exceed the amount of body fat, $_{552}^{551}$ $_{487}$ $1+\chi > \epsilon_{\sigma}$, and the adjusted time to reproduce must be positive, $_{553}$ $_{488}$ which given Equation 4, implies that $1 - \epsilon_{\lambda}^{1-\eta} \left(1+\chi\right)^{1-\eta} > 0$. $_{555}^{554}$ 489 Together these conditions imply that $\chi \in (-f_0 M^{\gamma-1}, 1/\epsilon_{\lambda} - 1)$ 555 556 $_{\rm 490}$ where the upper bound approximately equals 0.05. The modi- $_{\rm 557}^{\rm 557}$ $_{491}$ fied mass adjusts our model via the altered rates of starvation $_{559}^{500}$ $_{492} \sigma(M')$, recovery $\rho(M')$, and the maintenance of both starv- $_{560}$ 493 ing $\delta(M')$ and full consumers $\beta(M')$. Importantly, ϵ_{σ} , which $_{562}^{561}$ 494 determines the point along the growth curve that defines the 563 $_{495}$ body composition of starved foragers, is assumed to remain un- $_{565}^{564}$ 496 changed for the invader population (see SI for detailed derivations of invader rates).

To assess the susceptibility of the resident population to 569 499 invasion, we determine which consumer has a lower steady- 570 $_{500}$ state resource density for a given value of χ , again with the $_{572}^{571}$ $_{12}^{11}$. 501 expectation that populations able to survive on lower resource 573 $_{502}$ densities have a competitive advantage (67). We find that for $_{575}^{574}$ 13 . the solution of the state of t $_{505}$ invader has an intrinsic competitive advantage over the resident $_{579}^{516}$ 506 population. However, for $\hat{M} > 1.748 \times 10^7 \, \mathrm{g}$, leaner individuals 580 16. $_{507}$ (χ < 0) have lower resource steady state densities, switching $_{582}^{581}$ the advantage for higher values of M.

The observed switch in susceptibility as a function of χ at $^{584}_{585}$ $^{18.}_{19.}$ $M_{\rm opt}=1.748\times 10^7{\rm g}$ thus serves as an attractor, such that $^{586}_{586}$ $^{19.}$ $_{511}$ the NSM predicts organismal mass to increase if $M < M_{\rm opt}$ $_{588}^{587}$ $_{20.}$ and decrease if $M > M_{\rm opt}$. This value is close to but smaller $_{589}^{589}$ 513 than the asymptotic upper bound for terrestrial mammal body 590 21. $_{514}$ size predicted by the NSM, however it is remarkably close to $_{592}^{591}$ 515 independent estimates of the largest land mammals, the early 593 516 Oligocene Indricotherium at ca. 1.5×10^7 g and the late Miocene $^{594}_{595}$ 23. $_{517}$ Deinotherium at ca. $1.74\times10^7\mathrm{g}$ (61). Additionally, our calcula- $_{596}$ $_{24}$. 518 tion of $M_{\rm opt}$ as a function of mass-dependent physiological rates $_{598}^{597}$ (20) $_{598}^{598}$ 25. 519 is similar to theoretical estimates of maximum body size (60), 599 $_{520}$ and provides independent theoretical support for the observa- $_{600\ 26.}$ $_{520}$ and provides independent theoretical support for the observable tion of a 'maximum body size attractor' for North American $_{602}^{601}$ $_{27.}$ 522 mammals outlined by Alroy (59). While the state of the envi-523 ronment, as well as the competitive landscape, will determine 604 28. 524 whether specific body sizes are selected for or against (62), we 606 29. 525 propose that the dynamics of starvation and recovery described 607 $_{526}$ in the NSM provide a general within-lineage mechanism for the $_{609}^{\circ\circ\circ}$ $_{30.}$ evolution of larger body size among terrestrial mammals.

The energetics associated with somatic maintenance, $\frac{611}{612}$ 529 growth, and reproduction are important elements that influence 613 32. $_{530}$ the dynamics of all populations (11). The NSM is a minimal $_{615}^{614}$ $_{33.}$ ₅₃₁ and general model that incorporates the dynamics of starvation ₆₁₆ $_{532}$ and recovery that are expected to occur in resource-limited en- $_{617}^{617}$ 34 . vironments. By incorporating allometric relations between the $^{618}_{619}$ 35. 534 rates in the NSM, we found: (i) different organismal masses 620 $_{535}$ have distinct population dynamic regimes, (ii) allometrically- $_{621}^{621}$ $^{36.}$ 536 determined rates of starvation and recovery appear to minimize 623 37. $_{537}$ extinction risk, and (iii) the dynamic consequences of these rates $_{625}^{624}$ $^{38.}$ $_{538}$ may introduce additional drivers and hard boundaries on the $_{626}^{626}$ $_{39.}$ $_{539}$ evolution of maximum body size. We suggest that the NSM $_{538}^{627}$ $_{540}$ offers a means by which the dynamic consequences of energetic $_{629}^{\circ\circ\circ}$ 541 constraints can be assessed using macroscale interactions be-630 $_{542}$ tween and among species. Future efforts will involve exploring $_{632}^{631}$ 41 . 543 the consequences of these dynamics in a spatially explicit frame- $_{544}$ work, thus incorporating elements such as movement costs and $_{635}^{634}$ $^{42}\!\!.$

 $_{478}$ at $M=8.3\times10^8$, or 120 times the mass of a male African ele- $_{545}$ spatial heterogeneity, which may elucidate additional tradeoffs

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