Supporting Information for "The dynamics of starvation and recovery"

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7 Mechanisms of Starvation and Recovery

8 Our overall goal is to understand the dynamics of starva-9 tion, recovery, reproduction, and resource competition, where 10 our framework partitions starvation and reproduction into two 11 classes of the consumer: a full class that is able to reproduce 12 and a hungry class that experiences mortality at a given rate 13 and is unable to reproduce. For the dynamics of growth, re-14 production, and resource consumption, past efforts have com-15 bined the overall metabolic rate as dictated by body size with 16 a growth rate that is dependent on resource abundance and, in 17 turn, dictates resource consumption (see Refs. (1, 2) for a brief 18 review of this perspective). This approach has been used to 19 understand a range of phenomena including a derivation of on-20 togenetic growth curves from a partitioning of metabolism into 21 maintenance and biosynthesis (e.g. (1, 3–5)) and predictions 22 for the steady-state resource abundance in communities of cells 23 (2). Here we leverage these mechanisms, combined with several 24 additional concepts, to define our nutritional state model.

We consider the following generalized set of explicit dynamics for starvation, recovery, reproduction, and resource growth and consumption

$$\dot{F}_{d} = \lambda \left(R_{d} \right) F_{d} + \rho \left(R_{d} \right) H_{d} - \sigma \left(1 - \frac{R_{d}}{C} \right) F_{d},
\dot{H}_{d} = \sigma \left(1 - \frac{R_{d}}{C} \right) F_{d} - \rho \left(R_{d} \right) H_{d} - \mu H_{d},
\dot{R}_{d} = \alpha R_{d} \left(1 - \frac{R_{d}}{C} \right) - \left[\left(\frac{\rho \left(R_{d} \right)}{Y} + P_{H} \right) H_{d} + \left(\frac{\lambda \left(R_{d} \right)}{Y} + P_{F} \right) F_{d} \right]$$
[1]

where each term has a mechanistic meaning that we detail below (we will denote the dimensional equations with $_d$ before introducing the nondimensional form that was presented in the main text). In this set of equations $\lambda\left(R_d\right)$ and $\rho\left(R_d\right)$ are the growth and recovery rates as functions of the current resource availability. Typically these can be written as $\lambda\left(R_d\right) = \lambda_{max}S\left(R_d\right)$ or $\lambda\left(R_d\right) = \lambda_{max}S\left(R_d\right)$ where λ_{max} and ρ_{max} are the maximum growth and recovery rates respectively, which scale with body size as discussed later, and $S\left(R_d\right)$ is a saturating function of resources. The saturating function could, for example, be a Michaelis-Menten or Monod function of the form $\frac{R_d}{k+R_d}$, where k is the half-saturation constant. A simplified version of the Michaelis-Menten or Monod functional form, which caps tures the essential features, is a linear function that saturates to a constant value above a certain abundance of R_d .

In the above equations Y represents the yield coefficient 41 (e.g., (6,7)) which is the quantity of resources required to build 42 a unit of organism (gram of mammal produced per gram of 43 grass consumed) and P is the specific maintenance rate of resource consumption (g resource \cdot s⁻¹ \cdot g organism). If we pick 45 F_d and H_d to have units of (g organisms \cdot m⁻²), then all of the terms of \dot{R}_d , such as $\frac{\rho(R_d)}{Y}H_d$, have units of (g resource \cdot m⁻² 47 \cdot s⁻¹) which are the units of net primary productivity (NPP),

⁴⁸ a natural choice for \dot{R}_d . This choice also gives R_d as $(g \cdot m^{-2})$ which is also a natural unit and is simply the biomass density. ⁵⁰ In this system of units α (s⁻¹) is the specific growth rate of R_d , ⁵¹ and C is the carrying capacity, or maximum density, of R_d in ⁵² a particular environment.

Before describing the values of each of these constants, and a general nondimensionalization of the system of equations, it is important to consider the resource regimes associated with the above equations which lead to a simplification. As discussed above, the resource saturation function should be defined by a linear regime proportional to R_d when $R_d << k$, and a constant value for $R_d >> k$. Thus for hungry individuals, H_d , where $R_d << k$, we have that $\rho\left(R_d\right) \approx \rho_{max} R_d/k$, and for the full class, F_d , of organisms $\lambda\left(R_d\right) \approx \lambda_{max}$, such that the above relationships reduce to

$$\begin{split} \dot{F_d} &= \lambda_{max} F_d + \rho_{max} R_d H_d / k - \sigma \left(1 - \frac{R_d}{C} \right) F_d, \\ \dot{H_d} &= \sigma \left(1 - \frac{R_d}{C} \right) F_d - \rho_{max} R_d H_d / k - \mu H_d, \\ \dot{R_d} &= \alpha R_d \left(1 - \frac{R_d}{C} \right) - \\ & \left[\left(\frac{\rho_{max} R_d}{Y_H k} + P_H \right) H_d + \left(\frac{\lambda_{max}}{Y_F} + P_F \right) F_d \right]. \end{split}$$

We can formally nondimensionalize this system by choosing the general rescaling of $F = fF_d$, $H = fH_d$, $R = qR_d$, $t = st_d$, in which case our system of equations becomes

$$\begin{split} \dot{F} &= \frac{1}{s} \left[\lambda_{max} F + \rho_{max} \frac{R}{qk} H - \sigma \left(1 - \frac{R}{qC} \right) F \right], \\ \dot{H} &= \frac{1}{s} \left[\sigma \left(1 - \frac{R}{qC} \right) F - \rho_{max} \frac{R}{qk} H - \mu H \right], \\ \dot{R} &= \frac{1}{s} \left[\alpha R \left(1 - \frac{R}{qC} \right) - \frac{q}{f} \left[\left(\frac{\rho_{max} R}{Y_H k q} + P_H \right) H + \left(\frac{\lambda_{max}}{Y_F} + P_F \right) F \right] \right]. \end{split}$$

Reserved for Publication Footnotes

If we make the natural choice of s = 1, q = 1/C, and 110 reaching maturity given by $f = 1/Y_H k$, then we are left with

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

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

$$[4]$$

53 where we have dropped the subscripts on λ_{max} and ρ_{max} 113 (representing the cost on resources to return to the full state) 54 for simplicity, and $\xi = C/k$, $\delta = Y_H k P_H/C$, and $\beta = 114$ is given by $Y_H = M E_d/B_\rho$ where 55 $Y_H k \left(\frac{\lambda_{max}}{Y_F} + P_F \right) / C$. The above equations represent the sys-56 tem of equations presented in the main text.

 $\dot{R} = \alpha R (1 - R) - (\rho R + \delta) H - \beta F$

57 Parameter Values and Estimates

All of the parameter values employed in our model have either together, these relationships allow us to calculate ρ , δ , been directly measured in previous studies or can be estimated and β . 59 been directly measured in previous studies or can be estimated 117

74 metabolic rate and the latter represents the field metabolic rate. 75 We employ the field metabolic rate for our NSM model which 76 is appropriate for active mammals (Table 1).

An important feature of our framework is the starting size, $78 m_0$, of a mammal which adjusts the overall timescales for re-79 production. This starting size is known to follow an allometric so relationship with adult mass of the form $m_0 = n_0 M^{\nu}$ where s₁ estimates for the exponent range between 0.71 and 0.94 (see s₂ (8) for a review). We use $m_0 = 0.097 M^{0.92}$ from (9) which 83 encompasses the widest range of body sizes (8).

The energy to synthesize a unit of biomass, E_m , has been s₅ reported to vary between 1800 to 9500 (J g^{-1}) (e.g. (3-5)) in 86 mammals with a mean value across many taxonomic groups of 87 5,774 (J g⁻¹) (4). The unit energy available during starvation, ss E', could range between 7000 (J g⁻¹), the return of the total so energy stored during ontogeny (5) to a biochemical upper bound 132 Rate equations for invaders with modified body mass We allow 93 is returned during starvation (5).

99 $u_0 = 0.383$ and $\zeta = 1.00$ (13).

Typically the value of $\xi = C/k$ should roughly be 2. The $_{143} \chi_{\rm max} \approx 0.05$. value of ρ , λ , σ , and μ are all simple rates (note that we have 144 $_{102}$ not rescaled time in our nondimensionalization) as defined in $_{145}$ unchanged, the rate of starvation from the modified full state to 103 the maintext. Given that our model considers transitions over 146 the starved state, the rate of recovery from the starved state to 104 entire stages of ontogeny or nutritional states the value of Y 147 the modified full state, and the maintenance rates of both, will must represent yields integrated over entire life stages. Given $_{148}$ be different, such that $\sigma' = \sigma(M')$, $\rho' = \rho(M')$, $\beta' = \beta(M')$, $_{106}$ an energy density of $E_d = 18200$ (J g⁻¹) for grass (14) the $_{149}$ $\delta' = \delta(M')$. Rates of starvation and recovery for the invading maintenance value is given by $P_F = B_0 M^{3/4}/ME_d$, and the 150 population are easily derived by adjusting the starting or end-108 yield for a full organism will be given by $Y_F = ME_d/B_\lambda$ (g 151 ing state before and after starvation and recovery, leading to ₁₀₉ individual · g grass $^{-1}$), where B_{λ} is the lifetime energy use for ₁₅₂ the following timescales:

111 Similarly, the maintenance for hungry individuals is P_H = $_{112} B_0(\epsilon_{\sigma}M)^{3/4}/(\epsilon_{\sigma}M)E_d$, and the yield for hungry individuals

$$B_{\rho} = \int_{\tau(\epsilon_{\sigma}\epsilon_{\lambda})}^{t_{\lambda}} B_{0} m(t)^{\eta} dt.$$
 [6]

Finally, the value of α can be roughly estimated by the from combining several previous studies. Below we outline pre
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NPP divided by the corresponding biomass densities. From from combining several previous studies. Below we outline previous measurements and simple estimates of the parameters.

Metabolic rate has been generally reported to follow an exponent close to $\eta = 0.75$ (e.g., (3, 4) and the supplement of (5)).

We make this assumption in the current paper, although alternate exponents, which are know to vary between roughly 0.25
and 1.5 for single species (4), could be easily incorporated into rour framework, and this variation is effectively handled by the 20% variations that we consider around mean trends. The exponent not only defines several scalings in our framework, but also the value of the metabolic normalization constant, B_0 , given a set of data. For mammals the metabolic normalization congruence of the metabolic normalization congruence of the standard by the corresponding biomass densities. From the data in (15) we estimate the value of α to range between 120 2.81 × 10⁻¹⁰ (s⁻¹) and 2.19 × 10⁻⁸ (s⁻¹) globally. It should be noted that the value of α sets the overall scale of the F^* and H^* standard the standard problem of A to range between 120 2.81 × 10⁻¹⁰ (s⁻¹) and 2.19 × 10⁻⁸ (s⁻¹) globally. It should be noted that the value of α sets the overall scale of the F^* and H^* standard the standard problem of A sets the overall scale of the A with the data from Damuth (16). We find that the best fit is range above. However, two points are important to note here:

125 first, our framework predicts the overall scaling of A and A is also independent of A and this correctly matches data, and second, and this correctly matches data, and second, A that is, our prediction of the maximum mammal size does and 1.50 for A that is, our prediction of the maximum mammal size does and 1.51 the data in (15) we estimate the value of A that is, our prediction of the maximum mammal size does and 1.52 the data in (15) we estimate the value of A the total A the data in (15) we estimate the value of A to range developed A a

Table 1: Parameter values for mammals

Parameter	Value	References
$\overline{\eta}$	3/4	(e.g. (3-5))
E_m	$5774 (J \text{ gram}^{-1})$	(3-5)
E'_m	7000	(5, 10)
B_0	$0.047 \; (W \; g^{-0.75})$	(5)
γ	1.19	(12)
f_0	0.02	(12)
ζ	1.00	(13)
u_0	0.38	(13)

 $_{90}$ of $E'=36,000~(\mathrm{J~g^{-1}})$ for the energetics of palmitate (5, 10). $_{133}$ an invading subset of the resident population with mass M to For our calculations we use the measured value for bulk tissues $^{133}_{134}$ have an altered mass $M'=M(1+\chi)$ where χ varies between of 7000 which assumes that the energy stored during ontogeny $_{135}$ $chi_{\min} < 0$ and $chi_{\max} > 0$, where $\chi < 0$ denotes a leaner 136 invader and $\chi > 0$ denotes an invader with additional endoge-For the scaling of body composition it has been shown that 137 nous reserves. Importantly, we assume that the invading and fat mass follows $M_{\rm fat} = f_0 M^{\gamma}$, with measured relationships following $0.018 M^{1.25}$ (11), $0.02 M^{1.19}$ (12), and $0.026 M^{1.14}$ (13). The state of the same proportion of non-fat tisgues the values from (12) which falls in the middle of this that $\chi_{\rm min} = -f_0 M^{\gamma-1}$. Similarly, we assume that the invading $_{98}$ range. Similarly, the muscle mass follows $M_{\rm musc}=u_0M^{\zeta}$ with $_{141}$ organisms do not add endogenous reserves above the asympty-142 otic mass of the species M, such that $(1 + \chi)\epsilon_{\lambda}M < M$, and

Although the starved state of invading organisms remains

$$t_{\sigma'} = \frac{-M^{1/4}}{B_0/E'_m} \log\left(\frac{\epsilon_\sigma}{\chi + 1}\right),$$
 [7] 178
$$t_{\rho'} = \frac{-4M^{1/4}}{B_0/E'_m} \log\left(\frac{1 - (\epsilon_\lambda(\chi + 1))^{1/4}}{1 - (\epsilon_\lambda\epsilon_\sigma)^{1/4}}\right).$$
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The maintenance rates for the invading population require 184 $_{154}$ more careful consideration. First, we must recalculate the yields $_{185}$ $_{155}$ Y, as they must now be integrated over life stages that have also $_{186}$ $_{156}$ been slightly modified by the addition or subtraction of endoge- $_{187}$ $_{157}$ nous reserves. Given an energy density of $E_d=18200~(\mathrm{J~g^{-1}})_{188}$ $_{158}$ for grass (14) the maintenance value of the invading population $_{189}$ 159 is given by $P_F = B_0(1+\chi)M^{3/4}/(1+\chi)ME_d$, and the yield 190 for a full organism will be given by $Y_F = (1+\chi)ME_d/B_\lambda'$ (g 191 individual · g grass $^{-1}$) where B_λ' is the lifetime energy use for 192 162 the invading population reaching maturity given by

$$B'_{\lambda} = \int_{0}^{t_{\lambda'}} B_0 m(t)^{\eta} dt.$$
 [8] 194

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163 where

$$t_{\lambda'} = \frac{-4M^{1/4}}{B_0/E_m} \log \left(\frac{1 - (m_0/M)^{1/4}}{(1 - \epsilon_{\lambda}(1 + \chi))^{1/4}} \right).$$
 [9] $^{198}_{199}$ 10.

164 Note that we do not use this timescale to determine the repro-165 ductive rate of the invading consumer—which is assumed to re- 202 $_{166}$ main the same as the resident population—but only to calulate 203 167 the lifetime energy use. Similarly, the maintenance for hungry ²⁰⁴ 12. 168 individuals $P_H' = B_0 (\epsilon_\sigma (1+\chi)M)^{3/4}/(\epsilon_\sigma (1+\chi)M)E_d$ and the ²⁰⁵ 169 yield for hungry individuals (representing the cost on resources ²⁰⁶ 170 to return to the full state) is given by $Y_H' = (1+\chi)ME_d/B_\rho'$ ²⁰⁷ 13. 171 where 171 where

$$B'_{\rho} = \int_{\tau(\epsilon_{\sigma}\epsilon_{\lambda})}^{t_{\lambda'}} B_0 m(t)^{\eta} dt.$$
 [10] $^{209}_{210}$ 14.

172 Finally, we can calculate the maintenance of the invaders as

$$\delta' = P'_{H}Y'_{H}/\xi$$
 [11] ²¹⁴

$$\beta' = \left(\frac{\lambda_{\text{max}}}{Y'_{F}} + P'_{F}\right)Y'_{H}/\xi$$
 ²¹⁶
²¹⁷

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