

DEB-IBM User Manual

Dynamic Energy Budget theory meets individual-based modelling: a generic and accessible implementation

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This manual explains how to use the model DEB-IBM, which is a NetLogo implementation of a generic individual-based model based on Dynamic Energy Budget (DEB) theory. It also gives a quick overview of DEB theory and its basic parameters.

The rationale of the model and its implementation are also explained in:

Martin B, Zimmer E, Grimm V, Jager T. Year. Dynamic Energy Budget theory meets individual-based modelling: a generic and accessible implementation. Journal Volume: pages.

The NetLogo implementation and the complete model description following the ODD protocol can be found here:

NetLogo implementation.

ODD model description.

We recommend reading the article and the ODD model description first.

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1. Getting Started

This manual describes how to install and use DEB-IBM, a generic individual-based model that includes DEB theory as a submodel for the individuals' bioenergetics and life-cycle. The main task of users of DEB-IBM will be to parameterize the model and to possibly adapt the underlying NetLogo program to address a specific research question. In this manual we therefore first explain the DEB parameters used in DEB-IBM and then give examples for how to revise the code. The overall purpose of this model and its implementation are explained in Martin et al. ([Year](#)), whereas the model itself is described using the ODD protocol for describing individual-based models (Grimm et al. 2006, 2010) is available at [ODD website](#).

1.1 About NetLogo

The model is implemented in NetLogo, version 4.1.1 (Wilensky 1999). NetLogo is a software platform specifically designed for implementing individual-based and agent-based models. It includes powerful "primitives" (procedures) that allow users to implement even relatively complex models with relatively few lines of code and little or no previous programming experience. Complete novices to NetLogo and programming can generally program efficiently in 2-3 days; two textbooks on individual-/agent-based modelling are in press (Railsback and Grimm, in press¹; Wilensky and Rand, in press).

1.2 About DEB

Dynamic Energy Budget theory (DEB theory, Kooijman 2010) provides a set of rules that specifies the acquisition and use of energy in an organism, and thereby the life-history traits for the whole life-cycle of a single organism. More information about DEB can be found on <http://www.bio.vu.nl/thb/>. By using the Individual Based Model (IBM) as implemented in NetLogo, it is possible to investigate the population dynamics of a species following DEB theory. This is done by inputting the specific metabolic, (DEB) parameters for the species of interest. A continuously growing library of DEB parameters for all kinds of different species can be found on the above mentioned homepage, in the DEB laboratory (see `add_my_pet` parameter section below).

1.3 Installation

NetLogo is free software and can be downloaded from:

<http://ccl.northwestern.edu/netlogo/>; versions for the operation systems Windows, MacOS, and Linux are available. Installation is straightforward and usually does not take more than five minutes. To run and use the model DEB-IBM, start NetLogo and open "DEB-IBM.nlogo" (which comes with the DEB-IBM package that can be downloaded from [DEB-IBM webpage](#)).

¹ See also www.railsback-grimm-abm-book.com

1.4 How to use the model

NetLogo comes with three tabs: interface, information, and procedures. The “interface” tab is where users can input the DEB parameters of their species, alter environmental variables, and observe individual and population level output of the model. The “procedures” tab contains the NetLogo program, or code, implementing DEB-IBM. Here users can alter model structure, create new variables to monitor, add procedures for file output, and include other aspects of importance to the population dynamic of their species such as behaviour, space, and predation.

There are two levels of use for DEB-IBM. The first requires only familiarity with the interface. On the interface users can input the DEB parameters of their species and observe various population and individual variables such as population density, size structure, and reserve levels under various feeding conditions (data of all diagrams can be exported via the diagrams’ context menus or NetLogo’s “export” primitives). This level of use requires no programming. All information that users need to use the model are the DEB parameters of their species of interest. Thus, at this level the program allows users to learn how changes in metabolic parameters alter characteristics of individual life-histories and population dynamics.

The second level of using DEB-IBM involves modifying the generic program to answer specific research questions or to adapt the model to specific species. For example, a researcher may be interested in how the population dynamics of a species is influenced by changes in land use. In this case the researcher would adapt the standard model to include space and movement behaviour of individuals, with DEB theory acting as the energetic model. This more engaged use of the model requires users to be familiar with both the interface and procedure tabs. Likewise, species may show specific behaviours that are not captured by DEB theory; these behaviours could be added to the generic model. For this second level of using DEB-IBM, basic training in modelling and NetLogo are required. Beginners in both fields would need obtaining some literacy in both fields, for example by using the textbook of Railsback and Grimm (in press).

2. DEB parameters

Our implementation of the DEB-IBM is based on the scaled DEB model (Kooijman et al. 2008) and uses compound parameters. These compound parameters are derived from the 12 primary parameters of the standard DEB model (Table 1). Implementing our model in the scaled version of DEB rather than the standard DEB model further simplifies the model: by dividing the state variables “reserve”, “maturity”, and “reproduction” by the maximum surface-area-specific assimilation rate, one parameter of the standard model is removed as well as the unit of either energy or mass (standard DEB can be based in either) from the model. Working with the scaled DEB model with compound parameters allows parameterizing the DEB model for a species without directly measuring energy or mass (you cannot estimate energy parameters without measuring energies). See Kooijman et al. 2008 for a guide for parameterizing a DEB model.

While the general principles of DEB theory are relatively simple, the formulas used in our implementation have been algebraically rearranged, reduced (using compound parameters), and scaled. Thus the resulting formulas used in DEB-IBM may not seem intuitive. For a novice to DEB theory it may be difficult to understand what processes are actually driving fluxes for each of the DEB state variables. To facilitate a better understanding of DEB theory for those interested in population dynamic applications we below provide a brief introduction to the standard DEB parameters, how the compound parameters used in DEB-IBM relate to these parameters, and how changes in the each of the parameters affects the life-history of the modelled individuals. In addition to this user manual and the ODD model description, we recommend those new to DEB theory to first read (<http://www.bio.vu.nl/thb/deb/index.html>) for a non-technical introduction to the concepts of DEB theory and Kooijman et al. (2008) and Kooijman (2010) for a more formal description. For those already familiar with compound parameters and the scaled DEB model this section can be skipped.

2.1 DEB notation

In all text (manuscript, ODD model description, and user manual) we used standard DEB notation. This notation may look somewhat cumbersome at the beginning, but has a long history and is, by itself, highly consistent. Therefore, careful attention to the notation will spare users considerable time and confusion. We recommend routinely using Table 1 in the ODD model description, which contains a comprehensive list of all parameters dealt with in both the text and in the implementation of the model.

Quantities that are expressed as unit per structural volume are surrounded with “[]”, for example $[\dot{J}_{EM}]$ is maintenance rate per unit of volume, while a symbol enclosed in “{ }” indicates a quantity that is expressed per unit of surface area; for example $\{\dot{J}_{EAm}\}$ is the maximum surface-area-specific assimilation rate. The dots above J in $\{\dot{J}_{EAm}\}$, $[\dot{J}_{EM}]$, and all other symbols indicate that the quantity is a rate per unit of time.

Because the use of DEB notation is not possible within the code of NetLogo we have to convert the notation into a code-compatible notation. The names of the parameters correspond with the standard DEB notation as follows: a rate, which is in standard DEB identified by a dot above the letter, is here identified with the extension “_rate”. For instance, energy conductance, \dot{v} , is called “v_rate_int”. The “_int” portion refers to the fact that these are the initial, or baseline, parameters for a species. Users can allow individuals to vary in their DEB parameters from the initial parameters in some way, as we do for four of the DEB parameters (see [stochasticity section](#) below). Subscripts and superscripts in DEB notation are indicated by “_” and “^”, respectively. Although NetLogo is not case sensitive we keep cases consistent with DEB notation (Kooijman 2010). When a DEB parameter contains both a super and subscript the subscript goes first. For instance, scaled maturity at birth, U_H^b , is written “U_H^b_int”.

2.2 Standard DEB parameters

The most basic version of the model can be run with just defining the eight scaled DEB parameters in the left column of input fields on the interface (Fig. 1). The “scaled” version of DEB is a simplification of the standard DEB model, which uses compound parameters that are functions of standard DEB primary parameters. These compound parameters are often easier to extract from the data (Kooijman et al. 2008). Below we first provide a brief description of each of the 12 standard DEB parameters and then how these parameters are transformed to the scaled parameter set.

Figure 1. Input fields for the eight scaled DEB parameters used by default in DEB-IBM (screenshot taken from the NetLogo program). Differences between species are primarily represented by these values.

DEB-IBM parameters

k_M_rate_int	4
g_int	37734434
U_H'b_int	26685E-6
U_H'p_int	49555E-4
v_rate_int	0.16
kap_int	0.8
kap_R_int	0.95
k_J_rate_int	4

Table 1. The 12 primary DEB parameters and their associated processes. Shown are the parameters for the unscaled model where ‘energy’ is in the dimension of mass of reserves, which has the unit of mol.

Standard DEB Parameters			
Symbol	Unit	Description	Process
$\{\dot{F}_m\}$	$\text{m}^3 \text{d}^{-1} \text{m}^{-2}$	maximum surface-area-specific searching rate	Feeding/assimilation
$\{\dot{J}_{EA_m}\}$	$\text{mol d}^{-1} \text{m}^{-2}$	surface-area-specific maximum assimilation rate	Feeding/assimilation
y_{EX}	mol mol^{-1}	yield of reserve on food	Feeding/assimilation
\dot{v}	m d^{-1}	energy conductance	Reserve dynamics
κ	-	allocation fraction	Reserve dynamics
y_{VE}	mol mol^{-1}	yield of structure on reserve	somatic growth/maintenance
$\{\dot{J}_{ET}\}$	$\text{mol d}^{-1} \text{m}^{-2}$	surface-area-specific somatic maintenance	somatic growth/maintenance
$[\dot{J}_{EM}]$	$\text{mol d}^{-1} \text{m}^{-3}$	volume-specific somatic maintenance	somatic growth/maintenance
M_H^b	mol	maturity at birth	reproduction/development
M_H^p	mol	maturity at puberty	reproduction/development
\dot{k}_J	d^{-1}	specific maturity maintenance	reproduction/development
κ_R	-	reproduction efficiency	reproduction/development

2.2.1 Feeding and assimilation related parameters: $\{\dot{F}_m\}$, $\{\dot{J}_{EA_m}\}$, y_{EX}

The maximum surface-area-specific searching rate, $\{\dot{F}_m\}$, influences the functional response for a given prey type. Earlier versions of DEB used the half saturation coefficient, K , which

relates to $\{\dot{F}_m\}$ via $K = \{\dot{J}_{EAm}\} / [y_{EX} \{\dot{F}_m\}]$. Here y_{EX} is the yield of reserves on food or, in other words, the conversion efficiency of moles of food into moles of reserve; in most bioenergetic models this is referred to as assimilation efficiency. Dividing the surface-area-specific maximum assimilation rate, $\{\dot{J}_{EAm}\}$, by this conversion efficiency (y_{EX}) gives you the surface-area-specific maximum ingestion rate, $\{\dot{J}_{XAm}\}$. The ratio between the maximum surface-area-specific ingestion rate $\{\dot{J}_{XAm}\}$ and the maximum surface-area-specific searching rate, $\{\dot{F}_m\}$, gives you the half saturation coefficient (K) in a Holling type II functional response (this response follows from the assumption that the full time budget of an organism is spent either searching or handling food).

$$K = \frac{\{\dot{J}_{XAm}\}}{\{\dot{F}_m\}}$$

In recent formulations of DEB theory $\{\dot{F}_m\}$ has replaced K as a primary parameter in the standard DEB model because it is more closely linked to the underlying mechanism. For more mechanistic details and reasoning behind the feeding process see Kooijman (2010, p. 25).

2.2.2 Reserve dynamics parameters: κ , \dot{v}

In DEB theory, assimilated energy first enters a reserve before being mobilized for somatic or development and reproduction purposes. One of the assumptions of DEB theory is “weak homeostasis”, which means that at a constant food density the ratio of reserves to structure remains constant. The derivation of reserve dynamics from this assumption is rather complex and explained in Kooijman (2010, p. 37). In a simplified case of an organism which does not grow, for reserve density to remain constant assimilation would have to equal mobilization. However, for growing organisms the mobilization flux must be lower than assimilation flux to keep reserve density constant due to dilution of reserve density via growth. The dynamics resulting from the assumption of weak homeostasis are that mobilization of reserves will be proportional to reserve density, with the proportionality constant depending on the ratio of energy conductance, \dot{v} , and the length of the individual. Higher values of energy conductance, \dot{v} , imply a lower resistance of transfer from reserves to structure along the reserve-structure interface, thus the higher the conductance, the faster reserves are depleted and mobilized for use. The maximum reserve density $[E_M]$, the maximum amount of reserves per unit of volume, is given by the maximum surface-area-specific assimilation rate, $\{\dot{J}_{EAm}\}$, and energy conductance, \dot{v} .

$$[E_M] = \frac{\{\dot{J}_{EAm}\}}{\dot{v}}$$

This results, because the equilibrium density of a stock, in this case reserve density, depends on the inflow rate, in this case dependent on $\{\dot{J}_{EAm}\}$, and the outflow that is dependent on \dot{v} .

One of the core assumptions of DEB theory is that a fixed fraction of mobilized energy is allocated to the soma, while the remainder is allocated to development and reproduction. The allocation fraction, κ , represents the fraction of mobilized reserves that are allocated to

somatic growth and maintenance, while the remainder, $(1 - \kappa)$, are allocated to development and reproduction.

2.2.3 Somatic growth and maintenance parameters: $\{\dot{J}_{ET}\}, [J_{EM}], y_{VE}$

The fraction κ of mobilized reserves is allocated to the soma, i.e. the non-reproductive parts of the organism. In the soma, maintenance costs are paid first and the remaining energy is allocated to growth. There are two basic categories of maintenance costs, those which are surface-area-specific and those which are volume-specific. Surface-area-specific costs typically relate to heat loss of endotherms, but can also represent other surface-area-related costs such as osmoregulation. In the current implementation of the model we focus on ectotherms and we assume surface-area-specific costs to be negligible, i.e. $\{\dot{J}_{ET}\} = 0$. Some species in the Add_my_pet database are endothermic, thus heat-loss related costs will not be included in the current implementation of DEB-IBM, however this will be addressed in newer editions. Volume-specific maintenance rate $[J_{EM}]$ represents the costs associated with maintaining and implementing somatic functions (maintaining concentration gradients, turnover of structure, movement). Because this is a volume-specific rate, volume-related maintenance costs of a certain individual are obtained by multiplying $[J_{EM}]$ by the individual's volume, or structural length cubed, L^3 . Thus, in the absence of surface-area-related costs, an individual two times larger in volume or weight would have double the daily maintenance costs. The remaining mobilized energy is converted to growth, with an efficiency of y_{VE} , or in other words how many moles of structure are produced from one mol of reserve. Usually, in DEB theory we consider structure in units of volumetric length, which is $L = V^{1/3}$ (see Kooijman 2010 p. 10, for explanation structural (volumetric) length in comparison to measured physical length).

To convert moles of structure into volume, L^3 , or volumetric length, L , we use $[M_V]$ which converts moles to cubic centimetres; a typical value for this parameter is 4 mmol cm^{-3} (Kooijman 2010).

2.2.4 Development and reproduction parameters: $M_H^b, M_H^p, \dot{k}_J, \kappa_R$

The fraction of mobilized energy not allocated to the soma, $(1-\kappa)$, is allocated to development or reproduction. DEB theory divides the life history of all species into three classes: embryos, juveniles, and adults. Embryos do not feed externally but use maternal reserves for growth and development (at "conception" an embryo is composed of nearly only reserves). A transition from embryo to juvenile marks a switch to exogenous feeding. Neither embryos nor juveniles reproduce. The transition from juvenile to adult marks the start of investment into reproduction (the actual reproduction may occur a little later). In DEB theory these two transitions are made after a given amount of mobilized energy has been allocated to maturity to transition from embryo to juvenile, M_H^b , and to transition from juvenile to adult, M_H^p . Unlike structure, maturity has no mass or dimensions but rather is considered "information" (maturity is quantified by the cumulative amount of reserves invested in it). Like for soma, individuals must pay costs associated with maintaining a given

level of maturity. These costs are taken proportional to the maturity level (in mol of invested reserves); the proportionality constant is the maintenance rate coefficient, \dot{k}_j , with the units of d^{-1} . Total energy spent on maintaining maturity is proportional to maturity level. When individuals reach puberty, at $M_H = M_H^p$, maturation is complete and M_H^p represents a maximum level of maturity. Once an individual reaches puberty, energy remaining after maintenance costs for maturity are paid is allocated into a reproduction buffer. The reproduction buffer is depleted during reproduction (the creation of offspring) and is converted into embryonic reserves (embryos are nearly 100% reserves) with an efficiency equal to κ_R .

2.3 Standard DEB can be expressed in mass or energy

Seven of the 12 standard DEB parameters shown in Table 1 are expressed in dimension of mass (mol). However DEB can also be expressed in the dimension of energy (Joules). In this case, different notation is used for those seven DEB parameters (Table 2).

Table 2. The parameters and units of the energy- or mass-specific parameters in standard DEB. The five standard parameters not listed are not specific to either energy or mass (Table 1). A typical value to convert between energy and mass is 550 kJ mol^{-1} , and mass can be converted to volume via $[M_v]$ (Section 2.2.3) with a typical value of 4 mmol cm^{-3} (Kooijman 2010).

Standard Parameters in mass and energy			
Mass (moles)		Energy (Joules)	
Symbol	Unit	Symbol	Unit
$\{J_{EAm}\}$	$\text{mol d}^{-1}\text{m}^{-2}$	$\{\dot{p}_{Am}\}$	$\text{J d}^{-1}\text{m}^{-2}$
y_{EX}	mol mol^{-1}	κ_X	- (assimilation efficiency)
y_{VE}	mol mol^{-1}	$[E_G]$	J m^{-3}
$\{J_{ET}\}$	$\text{mol d}^{-1}\text{m}^{-2}$	$\{\dot{p}_T\}$	$\text{J d}^{-1}\text{m}^{-2}$
$[J_{EM}]$	$\text{mol d}^{-1}\text{m}^{-3}$	$[\dot{p}_M]$	$\text{J d}^{-1}\text{m}^{-3}$
M_H^b	mol	E_H^b	J
M_H^p	mol	E_H^p	J

Table 3. Symbols used for the state variables in the dimensions of mass and energy, and in the dimensionless scaled DEB used in DEB-IBM.

DEB state variables			
State variable	Mass	Energy	Scaled
Length	L	L	L
Reserves	M_E	E_E	U_E
Maturity	M_H	E_H	U_H
reproduction buffer	M_R	E_R	U_R

2.4 From standard to compound parameters

As mentioned earlier it is often convenient to work with compound parameters as they require less data to parameterize. These compound parameters represent combinations of primary parameters that are grouped together in the differential equations for the standard DEB model. In the current implementation we used two compound parameters: energy investment ratio, g (dimensionless), and specific somatic maintenance rate, \dot{k}_M (rate, t^{-1}). The former, g , is derived from the standard primary parameters using units of mass:

$$g = \frac{[M_V] \dot{v}}{\kappa \{ \dot{J}_{EAm} \} y_{VE}}$$

or energy:

$$g = \frac{[E_G] \dot{v}}{\kappa \{ \dot{p}_{Am} \}}.$$

Because maximum reserve density

$$[E_M] = \frac{\{ \dot{p}_{Am} \}}{\dot{v}},$$

we can think of g as the cost to create a unit of structure relative to the maximum reserve density which would be allocated to the soma:

$$g = \frac{[E_G]}{\kappa [E_M]}$$

\dot{k}_M is derived from the standard parameters using units of mass:

$$\dot{k}_M = \frac{[J_{EM}] y_{VE}}{[M_V]}$$

or energy

$$\dot{k}_M = \frac{[\dot{p}_M]}{[E_G]}$$

The reason why we see $[M_V]$ in the formulas for g and \dot{k}_M in the mass parameterization and not in the energy parameterization of the standard DEB model is that $[E_G]$ converts energy in reserves to growth in the dimension of length, while y_{VE} converts moles of reserves to moles of structure and $[M_V]$ is needed to convert from mass to structural length.

2.5 From standard DEB to scaled DEB

Using the two compound parameters g and \dot{k}_M in place of the primary parameters we get a simplified version of the standard DEB differential equations for reserves:

$$\frac{d}{dt}M_E = (\{J_{EAm}\}fL^2 - j_{EC})$$

with

$$j_{EC} = \{J_{EAm}\}L^2 \frac{ge}{g+e} \left(1 + \frac{L\dot{k}_M}{\dot{v}}\right)$$

and

$$e = v \frac{\dot{M}_E}{L^3 \{J_{EAm}\}}.$$

e is the scaled reserve density; the “scaled” is in reference to the amount of reserves per unit of structure (reserve density) relative to the maximum amount of reserves per unit of structure. Remembering that maximum reserve density

$$[E_M] = \frac{\{J_{EAm}\}}{\dot{v}}$$

we see that e is the total mass of the reserves divided by volume and maximum reserve density and will have a value between 0 and 1. The changes in DEB’s three state variables (Table 3), length, maturity, and reproduction buffer can then be calculated as follows:

$$\frac{d}{dt}L = \left(\frac{(\kappa j_{EC} - [J_{EM}]L^3)y_{VE}}{[M_V]} \right)^{1/3}$$

$$\frac{d}{dt}M_H = ((1-\kappa)j_{EC} - \dot{k}_J M_H) \text{ for } M_H < M_H^p \text{ else } \frac{d}{dt}M_H = 0$$

$$\frac{d}{dt}M_R = ((1-\kappa)j_{EC} - \dot{k}_J M_H^p) \text{ for } M_H > M_H^p \text{ else } \frac{d}{dt}M_R = 0.$$

As we mentioned earlier, our model is implemented in the scaled version of DEB. By this we mean that to remove the unit mol (or Joule if using the energy parameterization of DEB) we divide all state variables in moles (M_E , M_H , and M_R) by the maximum surface-area-specific assimilation rate $\{J_{EAm}\}$ (or $\{\dot{p}_{Am}\}$ if working with energy) to get scaled reserve U_E , scaled maturity U_H , and scaled reproduction buffer U_R (Table 3). Dividing both sides of the three differential equations by $\{J_{EAm}\}$ gives:

$$\frac{d}{dt}U_E = (fL^2 - S_C)$$

$$\text{With } S_C = L^2 \frac{ge}{g+e} \left(1 + \frac{L\dot{k}_M}{\dot{v}} \right)$$

$$\frac{d}{dt}U_H = ((1-\kappa)S_C - \dot{k}_J U_H) \text{ for } U_H < U_H^p \text{ else } \frac{d}{dt}U_H = 0$$

$$\frac{d}{dt}U_R = ((1-\kappa)S_C - \dot{k}_J U_H^p) \text{ for } U_H > U_H^p \text{ else } \frac{d}{dt}U_R = 0.$$

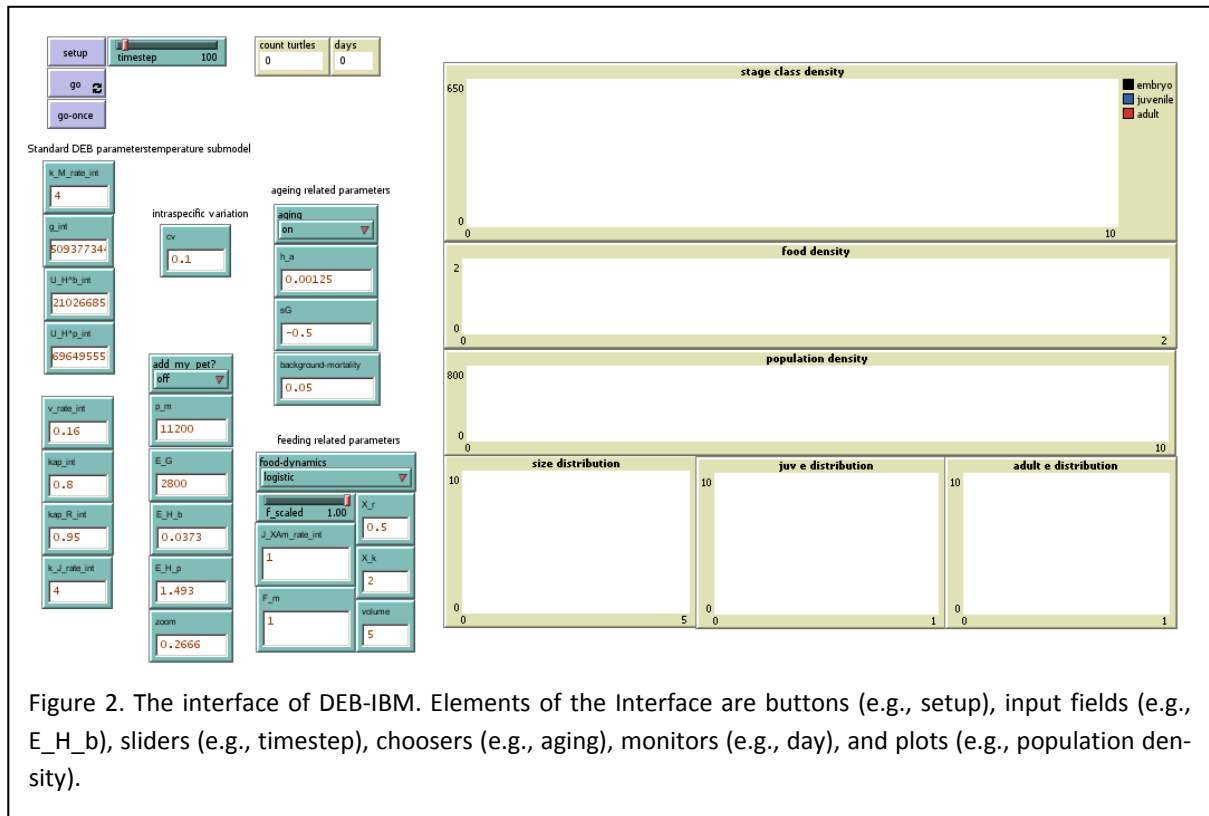
Notice that we now use scaled U_H^p and U_H^b for parameters related to life-stage transitions, which are equal to M_H^p and M_H^b divided by $\{\dot{J}_{EAm}\}$.

The length dynamics simplify to:

$$\frac{d}{dt}L = \frac{1}{3} \left(\frac{\dot{v}}{gL^2} S_C - \dot{k}_M L \right).$$

2.6 DEB-IBM parameters

We have now derived the equations used in DEB-IBM. If food conditions are constant we on-



ly need eight parameters to run simulations. To run DEB-IBM at constant food conditions, set the “food-dynamics” chooser to “constant” and set scaled assimilation rate to the desired value. The input boxes for the eight parameters needed to run DEB-IBM at constant food conditions are on the left-most side of the interface (Fig. 1). The bottom four parameters in the column are primary DEB parameters (\dot{v} , κ , κ_R , and \dot{k}_J). The top four parameters include

the two compound parameters g and \dot{k}_M , in addition to the two life-stage transition parameters, U_H^b and U_H^p .

3. The interface

In Fig. 2, you see what the interface looks like after opening the model in NetLogo.

3.1 Running the basic model

The first thing you should know is how to run the model. In Fig. 3, you see the most important buttons on the interface: The “setup” and “go” buttons. By pressing the “setup” button, you initialize the system, for instance, individuals are created and obtain values for their state variables, for example their DEB parameters. By pressing the

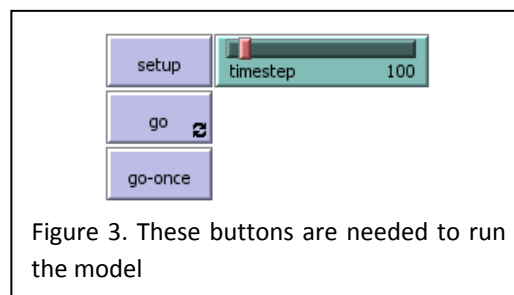


Figure 3. These buttons are needed to run the model

“go” button, the simulation starts; the simulation will run until you press “go” again. (Pressing “go-once” makes the program execute one timestep.) In the current implementation of the model the parameters for a species are set for individuals during the setup procedure. Thus altering a parameter value in the interface will not result in a change in the DEB individuals’ parameter values unless the setup procedure is run after the changes were made. This could be altered to allow “mid-simulation” modification of parameter values. In the basic version of the model, all of the standard DEB parameters are derived from the “Add my pet” database for the water flea *Daphnia magna*,

(http://www.bio.vu.nl/thb/deb/deblab/add_my_pet/index.php).

The timestep slider allows the user to control the length of a timestep. The value selected on the slider bar represents how many timesteps a day is divided into. Thus all of the DEB parameters which are input into the model should be daily rates. Because the model is a discrete implementation of differential equations (Euler method), the timestep needs to be small enough for the equations to function properly. How small a timestep needs to be is dependent on the parameter values of a species. Fast-growing species need shorter time steps.

3.2 Getting parameters

Here we give a brief discussion of how DEM-IBM is compatible with the Add_my_pet data base and for the many species not currently in the Add_my_pet data base we direct users to existing guides and a tool for model parameterization.

3.2.1 Add my pet parameters

A growing database of parameter sets for species can be found at: <http://www.bio.vu.nl/thb/deb/deblab>. There are many parameters listed in the Add_my_pet data base, but for use in DEB-IBM we will only need 9 of these (in code notation): v, kap,

kap_R, p_M, E_G, k_J, E_Hb, E_Hp, z (zoom factor). Additionally the two ageing parameters h_a and s_G are also given.

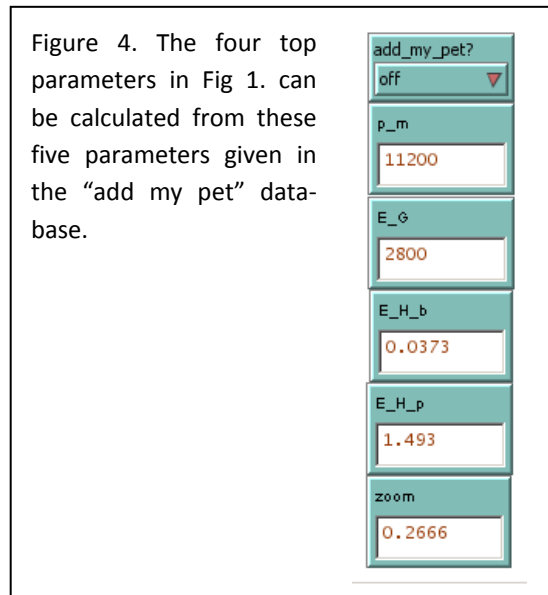
The parameter sets contained in the “add my pet” database are the primary DEB parameters in energy. Thus we need to convert the primary DEB parameters from the add_my_pet database to those used in our implementation. On the interface of DEB-IBM are input boxes for five add_my_pet parameters (Fig. 4). By selecting “on” in the “add_my_pet” chooser and clicking the setup button, DEB-IBM automatically converts these five add_my_pet parameters to the four DEB-IBM parameters at the top of the scaled DEB-IBM parameters column (Fig. 1). The bottom four are primary DEB parameters require no conversion from those listed in the add_my_pet database. The code for the conversion is in the procedure “convert-parameters” within the “setup” procedure. In the following paragraph, we explain how these parameters are calculated.

Noticeably missing from the add_my_pet database is the maximum surface-area-specific assimilation parameter, $\{\dot{p}_{Am}\}$. This is because $\{\dot{p}_{Am}\}$ is food type specific. Some food types are of higher energetic value than others and thus organisms raised on ad-libidum concentrations of varying quality food can assimilate, and thus grow at different rates. However, we can use the “zoom” factor to estimate the value of $\{\dot{p}_{Am}\}$. The zoom factor is the maximum volumetric length, L_M , of an organism in centimetres. It is called the zoom factor because DEB theory makes several predictions for scaling of DEB parameters inter-specifically with body size, and the scaling of these parameters leads to many observable covariations such as growth rate, respiration, and life span (van der Meer 2006; Kooijman 2010, chapter 8). In DEB theory maximum length is a function of three primary parameters: maximum assimilation rate, kappa, and volume-specific maintenance costs.

$$L_M = \kappa \frac{\{\dot{p}_{Am}\}}{[\dot{p}_M]}.$$

When L_M is expressed in centimeters it is equal to the “zoom factor” (z). Because add_my_pet gives values for all the parameters in this equation other than $\{\dot{p}_{Am}\}$, we can rearrange the formula to determine its value. Once the value of $\{\dot{p}_{Am}\}$ is determined, all other conversions are straightforward.

For U_H^b and U_H^p , we need to convert maturation threshold for birth and puberty to scaled maturity at birth and puberty by dividing by the surface-area-specific maximum assimilation rate:



$$U_H^b = E_H^b / \{p_{Am}\} \text{ And } U_H^p = E_H^p / \{p_{Am}\} \text{ where } \{p_{Am}\} = \frac{[\dot{p}_M]z}{\kappa}$$

For the two compound parameters we just use the formulas consisting of primary DEB parameters:

$$\dot{k}_M = \frac{[\dot{p}_M]}{[E_G]} \text{ and } g = \frac{[E_G]\dot{v}}{\{\dot{p}_{Am}\}\kappa}$$

3.2.2 How to get your own DEB parameters

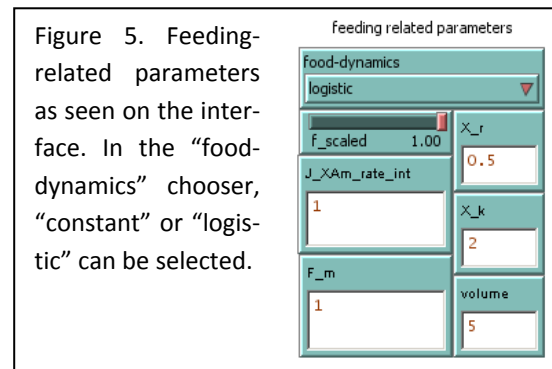
There are currently two reviews on parameterizing a DEB model (van der Meer 2006; Kooijman et al. 2008). A DEB model can be parameterized with varying levels of data, but generally if more data are available fewer assumptions need to be made. For use in DEB at constant food densities only the 8 scaled parameters need to be estimated (figure 1). Data of growth and reproduction over the lifespan (or some large portion of it) of individuals at more than one food level is enough to fully parameterize the model (Kooijman et al. 2008). There is also a tool (DEBtool <http://www.bio.vu.nl/thb/deb/deblab/debtool/>) available in Matlab or Octave which can be used to estimate parameters from available data sets. Kooijman et al. (2008) indicates which DEBtool procedures to use depending on the level of data available.

3.3 Optional submodels

3.3.1 Food dynamics

In the basic version of the model, the scaled functional response, f , (f_scaled in DEB-IBM) is set to 1 (ad-libidum feeding conditions). Keeping the food-dynamics constant, you can change the food supply by changing f_scaled on the slider. This can be done while the model is running without a new setup.

If you want to simulate a scenario with a dynamic food source, you can change the food-dynamics to “logistic” (just click on it). In the built-in scenario, a logistically growing prey population is depleted by the DEB population via predation. This is a very simple scenario, and it may be more realistic to let the DEB animal feed on another DEB organism but this is too specific for this implementation. For “logistic” prey dynamics, two new parameters are needed for the DEB individuals $\{j_{XAm}\}$ and $\{\dot{F}_M\}$. $\{\dot{F}_M\}$ is a primary DEB state variable for maximum surface-area-specific searching rate and $\{j_{XAm}\}$ is maximum surface-area-specific ingestion rate. $\{j_{XAm}\}$ differs from $\{j_{EAm}\}$ in that the latter only considers assimilated energy or $\{j_{EAm}\} = \{j_{XAm}\} y_{EX}$.



The prey population is characterized by the state variable density, X , and two parameters describing population growth rate, r (X_r), and carrying capacity K (X_k), respectively. For details see the prey dynamics submodel in the ODD model description. In the default model we assume feeding takes place in a three-dimensional environment (e.g. aquatic filter-feeders). However this can be modified to model feeding over two-dimensional surfaces. The parameter “volume” represents the size of the environment. The feeding submodel is only connected to the standard DEB model via the dimensionless scaled assimilation rate, f . Therefore, the units of X and volume can be user-defined (e.g. energy liter⁻¹, mg cm⁻³, cells per mm⁻³) as long as they are consistent with each other.

2 3.3.2 Ageing

4 In the basic version of the model, the ageing
6 submodel is turned “on”. Individuals age as described
8 in Kooijman (2010) and the ageing submodel section
10 of the ODD. If the ageing submodel is turned off,
12 animals have a daily background mortality rate.
14

Figure 6. Parameters as seen on the interface which affect the ageing submodel.

ageing related parameters

aging	on
h_a	0.00125
sG	-0.5
background-mortality	0.05

16 3.3.3 Stochasticity

18 In the standard DEB model the only inherent
20 source of stochasticity comes from the ageing
22 submodel. This can lead to extreme population
24 fluctuations because life-histories are exactly

Figure 7. parameter which controls the coefficient of variation. If set to 0, there is no intraspecific variation in DEB parameters

intraspecific variation

cv	0.1
----	-----

25 the same for all individuals which leads to synchronisation. One likely reason natural systems
26 do not always exhibit such drastic fluctuations is that stochastic processes and heterogeneity
27 among individuals prevent strong synchronization of life histories. One way of incorporating
28 stochasticity is to allow individuals to vary in some of their DEB parameters. This method is
29 justified because experiments often find that repeated physiological measurements of individuals
30 are less variable than those between individuals.

31 We followed the method outlined in Kooijman (1989) where individuals have a
32 random component in the maximum surface-area-specific ingestion rate, $\{J_{EAm}\}$. In our
33 implementation of the scaled DEB model there is no parameter $\{J_{EAm}\}$ because we scaled it
34 out of our model, but changing $\{J_{EAm}\}$ affects other parameter values indirectly. Both values
35 of the life-stage transition parameters will be affected because they are both scaled by
36 $\{J_{EAm}\}$. The maximum surface-area-specific ingestion rate will be influenced
37 $\{J_{XAm}\} = \{J_{EAm}\} / y_{EV}$, which further influences the half-saturation coefficient K as $K = \{J_{EAm}\} /$
38 $[y_{EX} \{F_m\}]$, and finally it affects g as $\{J_{EAm}\}$ is in the denominator of this formulation.

39 Thus, variation in all of these parameters is included by multiplying (for $\{J_{XAm}\}$) or
40 dividing (for g , U_H^b , and U_H^p) by a “scatter-multiplier” which is a log-normally distributed
41 number with user-defined standard deviation “cv”. Users can select the value of cv in the
42 “cv” input box. One way of estimating how large cv should be for your species is to compare

43 the spread of individual growth and reproduction trajectories over time to existing data. This
44 method can also be used to assess the appropriateness of the default pattern of intraspecific
45 variation, for example if no value of cv can be found which replicates the pattern of
46 intraspecific variation in growth and reproduction data, then some other pattern of
47 intraspecific variation may be more appropriate to use. If data on individual growth and
48 reproduction is available, the DEB model could be fit to individuals separately, and the
49 variation in DEB parameters could be used as an input into the model. Entering a value of 0
50 results in all individuals having the same parameters.

51 Obviously there are many other sources of stochasticity in natural systems. However
52 the sources of stochasticity to be included in the model are likely to be system-specific and
53 should be carefully considered by the researcher.

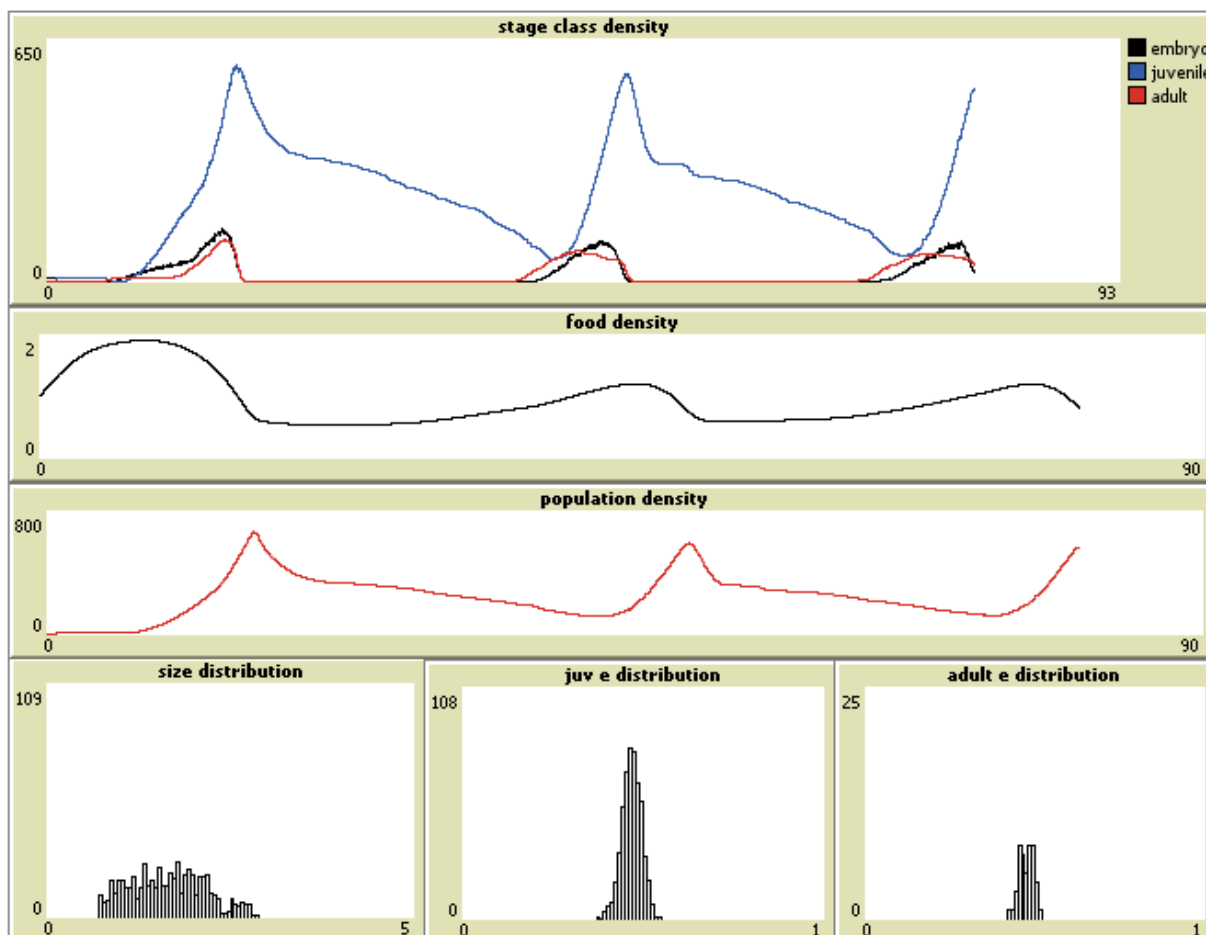
54 3.4 Plots, Histograms and Monitors

55 In the default model we include several output plots and histograms to monitor the
56 population dynamics of the modelled species. A plot of the population density, a plot of their
57 prey density, and a plot of the stage class density with the x axis in days (embryo, juvenile,
58 and adult). Additionally there are three histograms: the frequency distribution of length, L ,
59 and scaled reserve density, e , for both juveniles and adults. Ultimately the user can record
60 and plot any individual- or population-level variable of interest. For information on
61 generating output data, plots, monitors, and histograms, see the NetLogo User manual and
62 programming guide.

63 3.5 Using BehaviorSpace for analyzing the model

64 NetLogo contains a useful tool for model analysis, “BehaviorSpace”, which can be used for
65 testing population dynamics in different environmental conditions or characteristics of
66 individual via changes in parameter values. BehaviorSpace can also be used to perform
67 sensitivity analysis. The use of BehaviorSpace is well explained in the NetLogo User Manual;
68 additional guidance for using BehaviorSpace for, e.g., sensitivity analysis is given in Railsback
69 and Grimm (2011).

70



71
72 Figure 8. The three default plots and three histograms displayed on the DEB-IBM interface. The plot “stage class
73 class density” shows the density of each life stage (embryo, juvenile, and adult) over time. The plots “food density”
74 and “population density” show the density of the food (X) and total population. The “size distribution” histo-
75 gram shows the distribution of lengths (L) of the population. The histograms “juv e distribution” and “adult e
76 distribution” give the distribution of scaled reserve density (e) of juveniles and adults.

77 4. Procedures

78 In this tab the structure of the model can be modified and other aspects relevant to
79 population dynamics can be included. Here we discuss the basic structure of the code and
80 suggest how to modify it.

81 4.1 Basic code structure

82 The first section of the code declares the variables and specifies which type of variables they
83 are. DEB individuals are referred to as “turtles”, which is the NetLogo term for “agent” or
84 “individual”. Turtle variables are state variables characterizing the state of a certain turtle,
85 i.e. L , U_E and U_H . Additionally, because we allow some of the DEB parameters to differ
86 between individuals, we made the entire set of DEB parameters turtle variables (Grimm et
87 al. 2010).

88 In NetLogo, the spatial arena consists of square grid cells, called patches. The default
89 model is non-spatial and therefore consists of only one patch (updating the view of the
90 model world, or “view”, is therefore deactivated in the program). Patch variables are the

91 state variables of a patch. In our model the density of prey is a patch variable. This allows
92 one to easily make the model spatially explicit by defining a grid of patches, each with their
93 own states, e.g. prey and turtle density. Local predator-prey interactions are then easy to
94 include, e.g. feeding of DEB predators on a patch reduces the prey density only on that
95 patch.

96 Finally, “globals” are global variables which are known, and can be changed, any-
97 where in the program. In NetLogo programs, they are often used to implement model pa-
98 rameters, that need to be “known” by all individuals (turtles) or spatial units (patches).
99 “Globals” can either be declared in the procedures tab or created on the interface tab, in
100 which case they are not declared in the “globals-own[]”. The DEB parameters which do not
101 vary between individuals could have been made global variables but we chose to make them
102 turtle variables so that users could allow individuals to vary in any DEB parameter with little
103 programming effort. Note that on the interface, you can only use global variables, no turtle
104 or patch variables. Therefore, all eight DEB parameters on the interface (Fig. 1) are distin-
105 guished from the turtles variables by the suffix “_int”; the top four parameters are then
106 made to vary between individuals (see section 3.2.4).

107 The remainder of the code includes two major procedures: setup and go. The setup
108 procedure involves all processes required to initialize the model. In the setup procedure
109 some initial individuals are created and their state variables and parameters are specified. A
110 detailed description of the initialization is given in the ODD model description. The go
111 procedure runs the population model. An overview of the model processes and their
112 scheduling and a detailed description of each submodel are given in the ODD model
113 description.

114 4.2. Guide for adapting the model

115 For most applications the default model will need to be adapted in some way to address a
116 specific research question. These alterations may be either to adapt the standard DEB model
117 to reflect the life-history of the species of interest (ex. modifying the reproduction sub-
118 model) or to adapt the model to address a specific research question (ex. Including spatial
119 dynamics or more complex prey dynamics). Below we provide some examples of how the
120 default model can be adapted. In each example we show the major code changes needed to
121 implement each model adaptation, however the complete code for each example is given on
122 the website.

123 4.2.1 Feeding

124 In the standard DEB model, the assimilation rate depends on the surface area of the
125 predator and the density of the prey. These two variables are often sufficient to describe
126 feeding rates in controlled laboratory settings. Usually, however, varying environmental
127 conditions strongly influences foraging success. For example light intensity, turbidity, and
128 turbulence strongly influences encounter rate and capture success for most visual predators
129 in aquatic environments. Different types of habitat provide varying degrees of refuge for
130 prey species thus influencing the foraging rate of predators. These influences can be

131 incorporated into DEB-IBM via a mechanistic foraging submodel or a simple modification of f
 132 as a function of important environmental variables.

133 Users can also input a data file to specify f over the course of the simulation (see File
 134 I/O section in the NetLogo User Manual).

135 4.2.2 Reproduction

136 Differences between species are for the most part characterized by differences in their set of
 137 DEB parameters. However, species also exhibit differences in behaviour which are important
 138 for population dynamics. In the context of the DEB model, the most notable variation in
 139 behaviour is the reproduction strategy of a species. The default reproduction strategy in the
 140 model is for mature individuals to check if they have enough energy to reproduce; if they do
 141 they produce one embryo. Altering the reproductive strategy of the DEB individual to
 142 produce clutches of offspring requires a minor modification of the code. Below we give an
 143 example of how to modify the reproduction behaviour of the DEB animal.

144 DEB theory assumes that mothers in good conditions (higher scaled reserve density)
 145 produce higher quality offspring (offspring with higher scaled reserve density). This has been
 146 observed for many species, but there are exceptions (Kooijman 2010). Thus in the standard
 147 DEB model, mothers invest enough energy in an embryo so that when the embryo hatches
 148 ($U_H = U_{H_B}$) its scaled reserve density will be equal to its mothers scaled energy density.
 149 In the default version of DEB-IBM, mature individuals reproduce when they have enough
 150 energy to produce a single embryo with enough reserves to meet the condition noted above.

151 However, the water flea *Daphnia magna* does not produce one offspring at a time,
 152 but rather mature daphnids produce new broods every 2-3 days and the release of a brood
 153 coincides with molting. Time between reproduction events for *Daphnia* is dependent on
 154 temperature, but is independent of food. Because we are considering a situation where
 155 temperature is constant, we will assume that some internal clock triggers molting, and
 156 subsequently reproduction, at fixed intervals. To accomplish this, we need to give *Daphnia*
 157 individuals a new state variable (“repro-time”) to keep track of time since the last
 158 reproduction event, which increases by 1 each timestep (remember “timestep” represents
 159 how many timesteps one day is broken up into). We also need to create a global variable
 160 “days-between-repro”, which is a parameter representing how many days are between
 161 reproductive events; we will set this value to 2.5 days.

162 We update the reproduction part of the “go” procedure as follows:

```

163 [
164   if U_H >= U_H^p
165     [
166       set repro-time repro-time + (1 / timestep)
167       if repro-time > days-between-repro
168         [
169           calc-lay-eggs
170           if lay-egg? = 1
171             [
172               calc-embryo-reserve-investment
173               lay-eggs
174             ]
175         ]

```

```

176     ]
177   ]
178 ]

```

As we see above, individuals only reproduce when their time since last reproduction is greater than the new parameter “days-between-repro” which represents the time between reproduction events. “Calc-lay-eggs” is the next procedure which makes sure the individual has enough energy in the repro buffer to create at least one embryo. If not, repro-time will be set back to 0, and the reproduction buffer remains unchanged. The individual will then continue to accumulate energy in the reproduction buffer for another 2.5 days and then reproduce.

```

186
187 to calc-lay-eggs
188   set L_embryo L_0
189   set U_E_embryo U_R * kap_R
190   set U_H_embryo 0
191
192 loop [
193   ...
194   if U_H_embryo > U_H^b * 1 [ set lay-egg? 1 stop]
195   if e_scaled_embryo < e_scaled [set repro-time 0 stop]
196 ]
197 end
198

```

Once the energy required to create one offspring is determined, the individual will produce as many offspring as it has reserves for, each with the initial reserves equivalent to the value determined using the bisection method in the “calc-embryo-reserve-investment” procedure (see http://en.wikipedia.org/wiki/Bisection_method). The bisection method determines initial reserves via adaptive trial and error. Each estimation is the mean (therefore the name of this method, “bisection”) of upper and lower bounds set for the possible values of “initial reserves”. In the first estimation the upper bound is U_R / kap_R (this is because this is the highest value a mother can invest in an offspring), and a lower bound of 0. A simulation of the embryonic life stage is then run, and if the embryo matures with too much energy remaining in its reserves when it reaches energy for birth, the upper bound is then set to the previous “estimation”. We can do this because we know if the estimation was too large then all values larger than estimation will be too large and thus we can exclude those values from the range of possible values. If the value set for initial energy results in too little reserves left when the embryo reaches maturity needed to hatch, or the embryo has too little energy to reach the maturity threshold for hatching, the lower bound is then set to the “estimation” of initial reserves used in the simulation. This process repeats itself until the reserve density of the embryos’ matches that of the mothers within some acceptable range of error. In this simulation we allow 5% deviation between the embryo’s and mother’s reserve density.

```

217
218 to lay-eggs
219 hatch floor (U_R / estimation)
220 [
221   set die? 0
222   set mother-id id
223   set id who
224   set scatter-multiplier e ^ (random-normal 0 cv)

```

```

225     ...
226   ]
227   set lay-egg? 0
228   set repro-time 0
229   set U_R U_R - floor (U_R / estimation) * estimation
230
231 end

```

232 Notice that we also created two new state variables, “mother-id” and “id”. This section of
 233 code sets the mother-id of a new turtle to the id of the mother and the id of the new turtle
 234 to “who”: a built-in state variable of each turtle which is a unique identity number. We
 235 create these state variables because *Daphnia* carry their broods internally; thus if the mother
 236 dies, so do her offspring. We then have to modify the “update” procedure so that when a
 237 mother dies, the program checks to see if she is carrying any embryos (if she is, they die too).

```

238 to update
239 ; individuals update their state variables based on the calc_state variable
240 ; processes
241 ask turtles
242 [
243   ...
244   if die? = 1 and U_H >= U_H^p
245   [
246     let m-id id
247     let offspring turtles with [mother-id = m-id and U_H < U_H^b]
248     if any? offspring [ask offspring [die] ]
249     die ; the mother then dies
250   ]
251   if die? = 1 [die] ;
252 ]
253 ...
254 end

```

255 An important thing to remember is that all new state variables must be declared, either as
 256 “turtles-own” or global variables. Here, all variables except for “days-between-repro” (which
 257 is a global variable that we declared in the interface) are turtle variables. Additionally, in the
 258 setup procedure you have to tell turtles to “set id who” within the brackets following the
 259 hatch primitive where turtles are created.

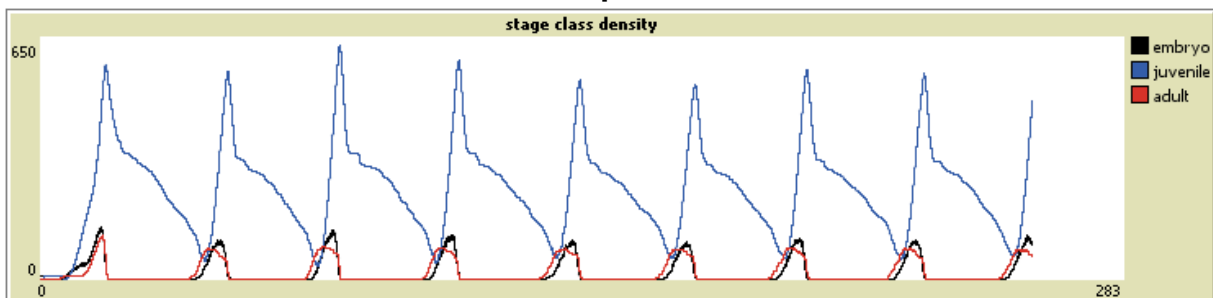
260 In Figure 9, we show comparisons of the population dynamics under logistic prey dy-
 261 namics, where on the top frame the population uses the *Daphnia* reproduction behaviour
 262 and the bottom frame is the default reproduction behaviour. *Daphnia* parameters were tak-
 263 en from the add_my_pet database. Below we see that the two reproduction strategies over-
 264 all do not result in widely different population dynamics.

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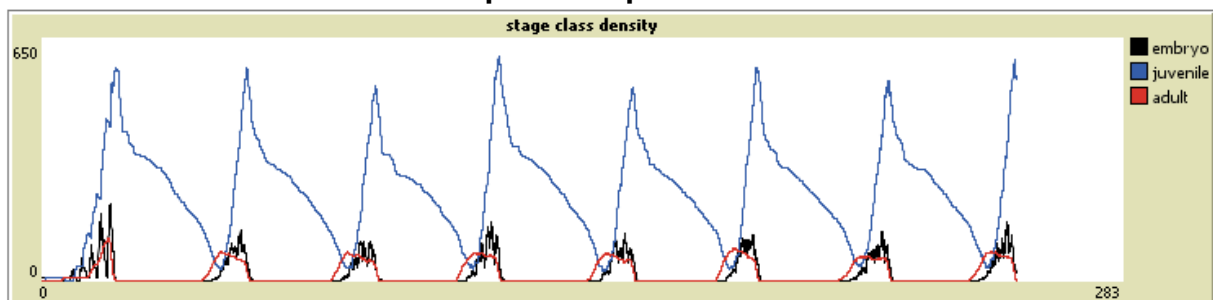
Figure 9. Density of 3 life-stages: embryo (black), juvenile (blue), and adult (red), under the default reproduction strategy (reproduce when enough energy to create one embryo), and the daphnia reproduction strategy (release broods at fixed intervals, 2.5 days). In the default strategy eggs are laid externally and survival is not dependant on the mother's survival, in the daphnia reproductive strategy if the mother dies while carrying embryos, those embryos also die. The predation submodel parameters of: $\{\dot{J}_{XAm}\} = 1$, $F_m = 1$, $X_r = 0.5$, $X_k = 2$, and volume = 5.

Default reproduction



285
286
287

"Daphnia" reproduction



288

289 4.2.3 Starvation

290 Starved individuals follow standard reserve dynamics until their scaled reserve density, e ,
291 falls below their scaled length L / L_m . Under this condition, individuals no longer mobilize
292 enough reserve to the soma to pay somatic maintenance costs, and thus must alter energy
293 allocation in some way. Continued starvation beyond this condition requires some alteration
294 of the reserve dynamics or its allocation. By default in DEB-IBM, individuals will no longer
295 grow, but divert just enough mobilized energy from reproduction and development to the
296 soma to pay maintenance costs. The remainder of mobilized energy is then allocated to
297 reproduction and development. When scaled reserve density falls below $\kappa L / L_m$, an individual
298 no longer mobilizes enough energy to pay somatic maintenance costs and thus dies. A
299 technical description of the starvation submodel is given in the ODD model description.

300 However, species differ in their response to starvation conditions. For example, an
301 individual may stop allocation to reproduction altogether when starved, reduce maintenance
302 costs, stop paying maturity maintenance, burn structure to pay maintenance costs, or

303 allocate all available energy into a final reproduction bout (emergency reproduction). How
 304 individuals respond to periods of starvation is likely driven by the fitness benefits associated
 305 with different strategies under the environmental conditions in which their genotype has
 306 evolved. Even within species the response to periods of starvation can vary depending on
 307 environmental conditions. For example, the energy allocation strategies of the pond snail
 308 depend on day length (Zonneveld and Kooijman 1989). Below, we give an example of how to
 309 modify reserve dynamics to an alternate starvation strategy. For a more thorough discussion
 310 of possible starvation strategies see Kooijman (2010, p. 118).

311 One possible starvation strategy is for individuals to stop growth, reproduction, and
 312 the payment of maintenance costs when $e < L/L_M$ and alter reserve dynamics to only
 313 mobilize enough energy for paying somatic maintenance. This starvation strategy was found
 314 to be appropriate for pond snails kept under short day conditions (12 hrs light / 12 hrs dark).
 315 In the unscaled version of the model this would be easy to implement, just by setting
 316 mobilization:

$$317 \dot{p}C = [\dot{p}_M]L^3$$

320 (Note that we are dealing with the DEB in energy and $\dot{p}C$ is analogous to $\{J_{EC}\}$.)

321 However, DEB-IBM is scaled and uses compound parameters which require some re-
 322 arrangement.

$$323 SC = \frac{\dot{p}C}{\{\dot{p}_{Am}\}} = \frac{[\dot{p}_M]L^3}{\{\dot{p}_{Am}\}}$$

326 because

$$328 [E_G] = g\kappa\{\dot{p}_{Am}\}/\dot{v}$$

330 thus:

$$332 SC = \frac{\dot{k}_M g \kappa}{\dot{v}} L^3 .$$

333
 334 Modifying the model is rather easy from here. The starvation strategy is coded in the “calc-
 335 dL” procedure. We need to modify the code to set dU_H or dU_R to 0 (depending on
 336 whether and individual is a juvenile or an adult) and then alter the mobilization flux to its
 337 new formula. An individual then dies when its scaled reserve density falls below zero.

```
338
339 to calc-dL ; calculate change in structural length
340   set dL  ((1 / 3) * (((V_rate / ( g * L ^ 2 )) * S_C) - k_M_rate * L))
341   if e_scaled < L / (V_rate / ( g * K_M_rate))
342     [
343       set dl 0
344       ifelse U_H < U_P_H
345         [set dU_H 0]
346         [ set dU_R 0]
```



```

347     set S_C (k_M_rate * kap * g * L ^ 3) / v_rate
348     set dU_E S_A - S_C
349     if e_scaled =< 0 [die]
350   ]
351 end

```

352 4.2.4 Spatial dynamics

353 DEB-IBM is non-spatial. However population dynamics can be influenced by the spatial dis-
354 tribution of resources. The model can be made spatial by clicking on the settings button on
355 the interface tab and setting the “max-pxcor” and “max-pycor” coordinates to the desired
356 size. (If you do so, make sure to deactivate the primitive “no-display” in the “setup” proce-
357 dure.) Files with coordinates of resource distribution, or GIS files can be input into NetLogo
358 to model real landscapes. However, to include spatial dynamics the DEB species of interest
359 should likely have some movement or dispersal capability which requires including a disper-
360 sal submodel in the procedure tab. Below we present an example of how to include spatial
361 dynamics into the current version of the model. This example is meant only to demonstrate
362 how to technically link DEB-IBM to spatial dynamics; the demonstration model was not de-
363 signed to answer any specific research question.

364 First you have to set the “max-pxcor” and “max-pycor” coordinates to the desired
365 world size. For this, you need to decide on what grid cell size would be appropriate for your
366 question. Note that within grid-cells, spatial relationships are often ignored, for example all
367 individuals within a grid cell might compete “globally” for the resources within the grid cell.
368 Grid cell size usually is chosen to represent typical distances of local competition (for further
369 aspects of choosing appropriate spatial and temporal scales, see Grimm and Railsback, 2005,
370 and Railsback and Grimm [in press]).

371 We arbitrarily picked a 80 by 80 grid. We then included a submodel in the “go”
372 procedure after all the DEB procedures. We implemented a simplified version of the
373 movement heuristic used in Hancock (2006). Once each day the individuals make a
374 probabilistic decision whether to stay in their current patch or move to one of their eight
375 neighbouring patches, where the probability of staying on its current patch or moving to a
376 neighbour patch is proportional to the relative amount of resources in each patch. To
377 accomplish this, at the first time step of every day the food (x) on the eight neighbouring
378 patches and in the turtle’s current patch are summed. The probability of the individual of
379 moving to (or staying on) patch i of the nine patches is determined by:

380

$$381 \quad \Pr(P_i) = \frac{x_i}{\sum_j x_j}$$

382

383 Below we show the entire movement submodel procedure. This procedure is run for each
384 individual. We use “p-” to denote “probability”, followed by the coordinates of the patch of
385 interest relative to the patch the turtle currently occupies: a (above), b (below), r (right), and
386 l (left), and h (here, patch the individual is on). Combinations of two letters denote the
387 patches located diagonally from the current patch, e.g. “p-ar” is the patch above and to the

388 right. Then, we choose the target patch by drawing a uniformly distributed random number
 389 from the interval [0, 1] and assigning target patches according to intervals within [0, 1] that
 390 correspond to the target patches probability of being chosen, i.e. $\Pr(P_i)$.

```

391
392 to movement-submodel
393   ask turtles with [U_H > U_H^B] [if ticks mod timestep = 0
394     [
395       let scale sum [x] of neighbors + [x] of patch-here
396       let p-a [x] of patch-at 0 1 / scale
397       let p-b [x] of patch-at 0 -1 / scale
398       let p-r [x] of patch-at 1 0 / scale
399       let p-l [x] of patch-at -1 0 / scale
400       let p-ar [x] of patch-at 1 1 / scale
401       let p-br [x] of patch-at 1 -1 / scale
402       let p-al [x] of patch-at -1 1 / scale
403       let p-bl [x] of patch-at -1 -1 / scale
404       let p-h [x] of patch-at 0 0 / scale
405       let random-number random-float 1
406
407       if random-number < p-a
408         [move-to patch-at 0 1]
409       if random-number >= p-a and
410         random-number < p-a + p-b
411         [move-to patch-at 0 -1]
412       if random-number >= p-a + p-b
413         and random-number < p-a + p-b + p-r
414         [move-to patch-at 1 0]
415       if random-number >= p-a + p-b + p-r
416         and random-number < p-a + p-b + p-r + p-r
417         [move-to patch-at -1 0]
418       if random-number >= p-a + p-b + p-r + p-l
419         and random-number < p-a + p-b + p-r + p-l + p-ar
420         [move-to patch-at 1 1]
421       if random-number >= p-a + p-b + p-r + p-l + p-ar
422         and random-number < p-a + p-b + p-r + p-l + p-ar + p-br
423         [move-to patch-at 1 -1]
424       if random-number >= p-a + p-b + p-r + p-l + p-ar + p-br
425         and random-number < p-a + p-b + p-r + p-l + p-ar + p-br + p-al
426         [move-to patch-at -1 1]
427       if random-number >= p-a + p-b + p-r + p-l + p-ar + p-br + p-al
428         and random-number
429           < p-a + p-b + p-r + p-l + p-ar + p-br + p-al + p-bl
430         [move-to patch-at -1 -1]
431       if random-number >= p-a + p-b + p-r + p-l + p-ar + p-br + p-al + p-bl
432         [move-to patch-at 0 0]
433     ]
434
435

```

435 Additionally to show the spatial distribution of prey density we can implement the following
 436 code in the go statement following the movement-submodel procedure.

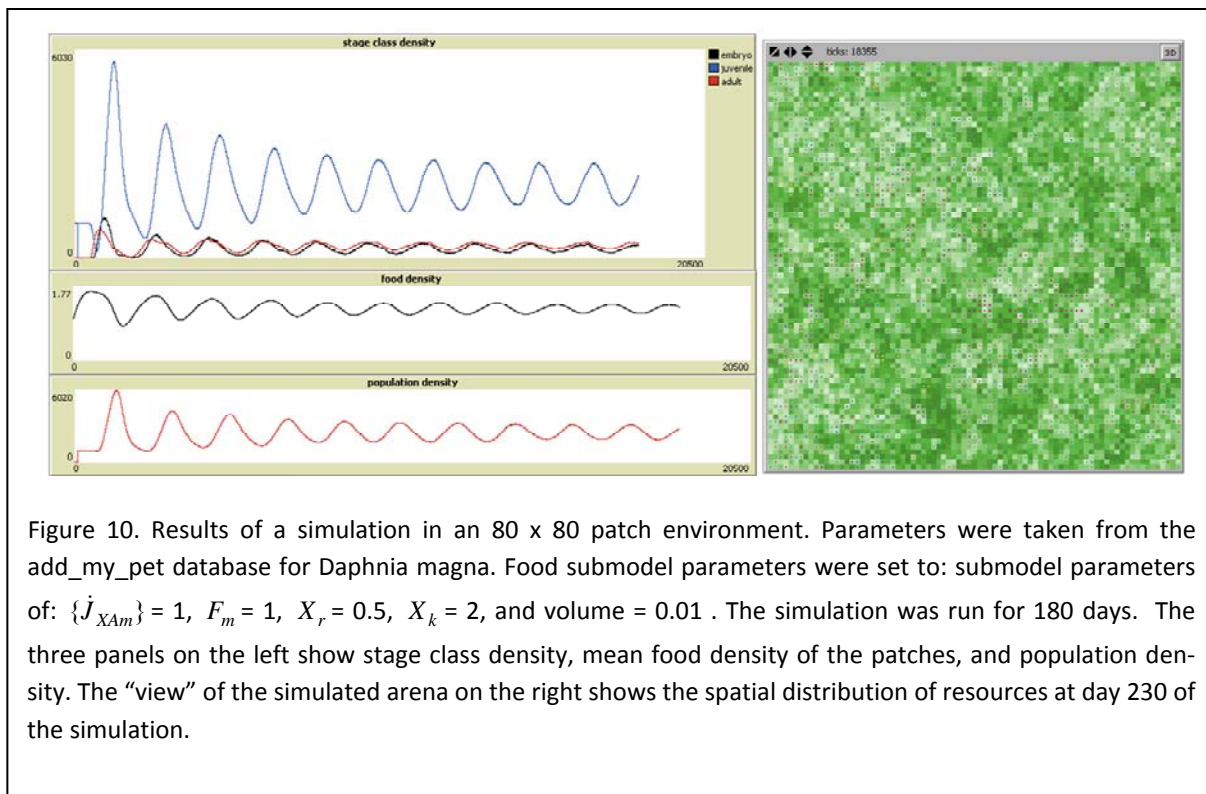
```

437
438 ask patches ask patches [ set pcolor scale-color green X 2 0]
439

```

440 This line of modified code does not affect the model run but scales the color of each patch to
 441 its level of food density (see primitive “scale-color” in color subsection of NetLogo
 442 programming guide). Because food density is a patch state variable, each patch has an

443 independent food density, which undergoes logistic growth. Predation is local, as DEB
 444 predators reduce only the density of prey in the patch they currently occupy.



445 Notice that in Fig. 10 the population initially shows large fluctuations but eventually, these
 446 fluctuations dampen dramatically. It is interesting to note that here population oscillations
 447 are much smaller than in the non-spatial model. Like many other models DEB population
 448 models typically exhibit the phenomenon known as “paradox of enrichment”. When carrying
 449 capacity of the food is much higher than the half-saturation coefficient, which in our feeding
 450 model is given by: $\{\dot{J}_{XAm}\} / F_m$, the population will often show large fluctuations, and if the
 451 carrying capacity of the food is much higher the populations will even collapse. The above
 452 example shows how the inclusion of spatial dynamics and the movement behaviour of indi-
 453 viduals can lead to a resolution of the paradox of enrichment. However, this is just one
 454 mechanism that can stabilize populations, and many other processes can also stabilize popu-
 455 lations at high food carrying capacities such as: inducible defences of the prey type, multiple
 456 prey types, interference competition, environmental heterogeneity and stochasticity. The
 457 importance of these mechanisms should be considered when modelling in a population con-
 458 text, and the inclusion of one or more of these mechanisms may be required to replicate re-
 459 alistic population dynamics.

460

461

462

463

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