

# A Trait Based Model From Scale Free Assumptions

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November 8, 2017

## 1 Model Definition

### 1.1 The Basics

We model our ecosystem by keeping track of the abundance  $N(w, w_*, t)$  of fish of weight  $w$  at time  $t$ . Here  $w_*$  denotes the weight of the fish at maturity. We suppose the maturity weight  $w_*$  represents the species of the fish, in the sense that fish of the same weight can act differently in our model, only if they have different maturity weights. We allow  $w$  and  $w_*$  to vary continuously, and to take any positive values, and hence we are assuming a continuum of species, characterized by different size traits  $w_*$ . The abundance  $N(w, w_*, t)$  is defined such that

$$\int_{w_1}^{w_2} \int_{w_{*1}}^{w_{*2}} N(w, w_*, t) . dw . dw_* \quad (1)$$

is the number of fish per unit volume which have a weight  $w \in [w_1, w_2]$  and a maturity weight  $w_* \in [w_{*1}, w_{*2}]$ .

The abundance  $N(w, w_*, t)$  of fish with a particular maturity weight  $w_*$  vary according to the McKendrick–von Foerster equation

$$\frac{\partial N(w, w_*, t)}{\partial t} = - \frac{\partial [N(w, w_*, t) g(w, w_*, t)]}{\partial w} - \bar{\mu}(w, w_*, t) N(w, w_*, t) \quad (2)$$

for time  $t \in \mathbb{R}$ , and weight  $w \in \left[ w_e(w_*), \frac{w_*}{\eta} \right]$ . In general, the growth rate  $g$  and mortality rate  $\bar{\mu}$  at time  $t$  depend integrals over abundances, and these abundances may change with time. However, for notational similitude we often neglect to notate the time dependence, and write these terms as  $g(w, w_*)$  and  $\bar{\mu}(w, w_*)$  respectively, as the only time dependence in these terms come from their dependence upon abundances. We insist that the abundance  $N(w, w_*, t)$  of fish with weight  $w \notin \left[ w_e(w_*), \frac{w_*}{\eta} \right]$  must be zero.

We further insist that our abundances  $N(w, w_*, t)$  must satisfy the boundary condition:

$$g(w_e(w_*), w_*)N(w_e(w_*), w_*, t) = \frac{\epsilon}{2w_e(w_*)} \int_0^\infty N(w, w_*, t) E_g(w) \psi(w, w_*) dw, \quad (3)$$

the left hand side of which represents the flux of new offspring into the system, and the right hand side of which represents how the amount of such new offspring is obtained by integrating over the reproducing fish of different weights. An abundance function  $N(w, w_*, t)$  is a solution to our system if and only if it satisfies (2) and (3) for each  $w_* > 0$ ,  $t \in \mathbb{R}$ , and  $w \in \left[w_e(w_*), \frac{w_*}{\eta}\right]$ , while  $w \notin \left[w_e(w_*), \frac{w_*}{\eta}\right] \Rightarrow N(w, w_*, t) = 0$ .

Here  $g(w, w_*)$  is the growth rate,  $\bar{\mu}(w, w_*)$  is the mortality rate,  $w_e(w_*)$  is the egg size of species with  $w_*$  as their maturity weight,  $\frac{w_*}{\eta}$  is the asymptotic weight of such a species,  $\epsilon$  is the reproductive efficiency,  $E_g(w)$  is the energy available for reproduction and growth, and  $\psi(w, w_*)$  is the fraction of such energy which a fish uses for reproduction. We shall describe these quantities in more detail later.

## 1.2 Scale Free Assumption

Setting up a size spectrum model required the definition of many functions representing features of the feeding, growth, death and reproduction of fish. Often the real world nature of these functions is poorly understood, or the model becomes analytically intractable once there are modeled in sufficient detail. Therefore various assumptions are often introduced, such as the assumption of constant feeding level in Hartvig. Instead of this, our main simplifying assumption is that our model should be scale invariant, in the sense that if we apply particular kinds of rescaling upon the units of weight, maturity weight, time, and abundance, we get another solution to our system. Using this assumption gives us natural constraints on the forms of functions we should use which are often in remarkable agreement to function forms justified by other researchers for other reasons. Applying our particular type of rescaling to an abundance function  $N : \mathbb{R}_+ \times \mathbb{R}_+ \times \mathbb{R} \rightarrow \mathbb{R}$  using a scaling constant  $c > 0$  yields a new abundance function  $F_c[N] = \tilde{N}$  where  $\tilde{N} : \mathbb{R}_+ \times \mathbb{R}_+ \times \mathbb{R} \rightarrow \mathbb{R}$  is defined such that  $\tilde{N}(w, w_*, t) = c^\Lambda N(cw, c^A w_*, c^B t)$  for each  $w, w_* > 0$  and each  $t \in \mathbb{R}$ . Here  $\mathbb{R}_+$  denotes the set of positive reals. Sometimes we call  $F_C$  a symmetry transform because it represents a type of rescaling that we require the key features of our model to be invariant under. Formally, our scale free assumption is that: if the abundance function  $N$  satisfies our system, then we require the rescaled abundance function  $F_c[N]$  to also be a solution to the system, for each  $c > 0$ .

## 1.3 Prey Switching

The problem of coexistence, is the problem of making an ecological model where multiple different species competing for common resources can coexist. Coexis-

tence is not generally possible in the classical size spectrum models of [Hartvig] for example, and this can be seen by noting that there are no steady state of the McKendrick–von Foerster equation they use, where all species coexist, that satisfies the boundary conditions for each maturity weight  $w_*$ . We resolve this problem by adding a small amount of extra density dependence into the dynamics of predation and mortality. In the classical theory, the total mortality rate of individuals with mass  $w$  and maturity weight  $w_*$  is  $\mu(w, w_*)N(w, w_*, t)$ , where  $\mu(w, w_*)$  is the per capita mortality rate. In our model, the total mortality rate is instead

$$N(w, w_*, t)\bar{\mu}(w, w_*) := N(w, w_*, t)\mu(w, w_*)\hat{N}(w, w_*, t)^\chi \quad (4)$$

where  $\mu(w, w_*)$  is the unadjusted mortality rate (before we include prey switching), and the prey switching constant  $\chi \geq 0$  is some (typically small) number which abstractly represents the amount of prey switching. Also

$$\hat{N}(w, w_*, t) = N(w, w_*, t)w^\Omega w_*^\Gamma \Delta \quad (5)$$

where  $\Delta, \Omega$  and  $\Gamma$  are constants. We do not give a direct ecological interpretation to this new prey switching factor, but the result of adding the factor is that it makes the total mortality rate into a super linear function of the individuals density in a way which works well with our scale free assumption. Adding such density dependence seems justified ecologically since large groups of prey are more likely to be targeted by predators. This assumption that predation occurs at a rate superlinear to the preys density will be adjusted for when we consider the energy input to predators. Also, the unadjusted mortality rate

$$\mu(w, w_*) = \mu_p(w) + \mu_s(w) + \mu_b(w_*) \quad (6)$$

can be written as the sum of the predation mortality rate  $\mu_p(w)$ , the starvation mortality rate  $\mu_s(w)$  and the background mortality rate  $\mu_b(w_*)$ , and so we are assuming that there is also density dependence upon the death of individuals due to starvation and background effects. Only a small amount of prey switching (e.g.,  $\chi = 0.05$ ) is often required to make it possible for a solution of (2) to satisfy the boundary conditions (3).

Perhaps the simplest form of prey switching would be to pick  $\Omega = \Gamma = 0$ , in which case the prey switching corresponds to a simple additional dependence upon the abundance. Similar biomass based prey switching can be encoded by setting  $\Omega = 1, \Gamma = 0$ . If one lets  $\Omega = 1, \Gamma = 1$  and  $\Delta = \delta\delta_*$  then the prey switching term approximates the number of individuals (per unit volume) with nearby weights and maturity weights, in the sense that

$$\hat{N}(w, w_*, t) \approx \int_w^{(\delta+1)w} \int_{w_*}^{(\delta_*+1)w_*} N(w', w'_*, t).dw'.dw'_* \quad (7)$$

in this case. If one lets  $\Omega = 2, \Gamma = 1$  and  $\Delta = \delta\delta_*$  then the prey switching term approximates the total biomass (per unit volume) of individuals with nearby weights and maturity weights, in the sense that

$$\hat{N}(w, w_*, t) \approx \int_w^{(\delta+1)w} \int_{w_*}^{(\delta_*+1)w_*} w' N(w', w'_*, t) dw'_* dw' \quad (8)$$

in this case.

Since adding a prey switching term  $\hat{N}(w, w_*, t)$  (of whatever form) will effect the total predation mortality rate, there will be a corresponding modification to the term  $\phi(w)$  which denotes the total amount of energy available to a size  $w$  predator. The classical expression for available energy in [Hartvig] was essentially

$$\int_0^\infty \int_0^\infty w' N(w', w'_*, t) s(w', w) dw'_* dw' \quad (9)$$

where

$$s(w, w') = \exp \left[ - \left( \ln \left( \frac{\beta w'}{w} \right) \right)^2 / (2\sigma^2) \right] \quad (10)$$

represents the preference that a predator of weight  $w$  has for a prey of weight  $w'$ . Here  $\beta$  is the preferred predator-prey mass ratio, and  $\sigma$  represents the width of sizes of prey which the predator will tolerate. When we include our prey switching, the expression for available energy changes to

$$\phi(w) = \int_0^\infty \int_0^\infty w' N(w', w'_*, t) \hat{N}(w, w'_*, t)^\chi s(w', w) dw'_* dw' \quad (11)$$

and so when prey switching is present (i.e., when  $\chi > 0$ ) the amount of food extracted from the prey goes up super-linearly with their abundance.

## 1.4 Growth Rate

The available energy (i.e., the food available)  $\phi(w)$  to a weight  $w$  predator is given by Equation (11). To get the energy encountered  $E_e(w)$  by such a predator  $\phi(w)$  must be multiplied by the volumetric search rate  $\gamma w^q$  so

$$E_e(w) = \phi(w) \gamma w^q. \quad (12)$$

The feeding level of an individual is

$$f(w) = \frac{E_e(w)}{E_e(w) + hw^n} \quad (13)$$

where  $hw^n$  is the maximum food intake rate of a weight  $w$  individual. The total energy intake  $f(w)hw^n$ , however we suppose energy is only assimilated with an efficiency  $\alpha$  and a weight  $w$  individual must spend energy  $kw^p$  in order to satisfy its metabolic needs or else it will be subject to potential starvation. It follows that the energy available for growth and reproduction is

$$E_g(w) = \alpha f(w)hw^n - kw^p. \quad (14)$$

The growth rate is now defined as

$$g(w, w_*) = \begin{cases} (1 - \psi(w, w_*)) E_g(w) & \text{if } E_g(w) > 0 \\ 0 & \text{otherwise.} \end{cases} \quad (15)$$

## 1.5 Mortality Rate

When we have taken account of prey switching the (per capita) mortality rate of individuals is  $\bar{\mu}(w, w_*) = \mu(w, w_*) \hat{N}(w, w_*, t)^\chi$ , where the unadjusted (per capita) mortality rate is

$$\mu(w, w_*) = \mu_b(w_*) + \mu_s(w) + \mu_p(w) \quad (16)$$

is the sum of the background mortality rate

$$\mu_b(w_*) = \mu_0 w_*^Q \quad (17)$$

(note in [Hartvig] they chose  $Q = n - 1$ , constraining the rescaling exponent choices  $A, B, \Lambda$ )

and the starvation mortality rate

$$\mu_s(w) = \begin{cases} \frac{-E_g(w)}{\xi} & \text{if } E_g(w) < 0 \\ 0 & \text{otherwise.} \end{cases} \quad (18)$$

and the predation mortality rate

$$\mu_p(w) = \int_0^\infty \int_0^\infty \left( \frac{f(w') h w'^n}{E_e(w')} \right) N(w', w_*, t) s(w', w) \gamma w'^q dw' dw_* \quad (19)$$

where  $\left( \frac{f(w') h w'^n}{E_e(w')} \right)$  is the energy income of the weight  $w'$  predators, divided by the total energy encountered. This equals the fraction of encountered individuals they consume. The remaining terms inside the integral represent the amount of weight  $w$  prey encountered by predators. Rearranging (13) into an equation for  $E_e(w)$  and substituting it into (19) yields

$$\mu_p(w) = \int_0^\infty \int_0^\infty (1 - f(w')) N(w', w_*, t) s(w', w) \gamma w'^q dw' dw_* \quad (20)$$

which is the same as the expression appearing in [Hartvig].

## 2 Using Scale Invariance

The McKendrick–von Foerster equation (2) can be rewritten as

$$\frac{\partial N(w, w_*, t)}{\partial t} = - \frac{\partial [N(w, w_*, t) g(w, w_*)]}{\partial w} - (\mu_b(w_*) + \mu_s(w) + \mu_p(w)) (w^\Omega w_*^\Gamma \Delta)^\chi N(w, w_*, t)^{\chi+1} \quad (21)$$

In this section we shall demonstrate how particular choices of parameters can lead to a model that satisfies our scale free assumption. We shall assume that  $N : \mathbb{R}_+ \times \mathbb{R}_+ \times \mathbb{R}$  is a solution to our system (so it satisfies (3) and (21)). In this section, we write  $\mathcal{A} \mapsto \mathcal{B}$  to denote that expression  $\mathcal{B}$  is obtained by taking expression  $\mathcal{A}$  and replacing each  $N(w, w_*, t)$  with  $c^\Lambda N(cw, c^A w_*, c^B t)$ . Essentially the notation  $\mathcal{A} \mapsto \mathcal{B}$  denotes that expression  $A$  transforms to expression  $B$  when we apply the symmetry transform  $F_c$  for generic  $c > 0$ . We shall now demonstrate that (for appropriate parameter choices) our scale free assumption holds, in the sense that after we replace the  $N$ 's with  $F_c[N]$ 's the resulting expression still satisfies (3) and (21).

## 2.1 Time

To demonstrate the scale free assumption, we shall first consider how our symmetry transform alters the different terms of Equation (21), starting with the left hand side:

$$\frac{\partial N(w, w_*, t)}{\partial t} \mapsto \frac{\partial [c^\Lambda N(cw, c^A w_*, c^B t)]}{\partial t} = c^\Lambda \frac{\partial N(\tilde{w}, \tilde{w}_*, \tilde{t})}{\partial \tilde{t}} \frac{\partial \tilde{t}}{\partial t} = c^{\Lambda+B} \frac{\partial N(\tilde{w}, \tilde{w}_*, \tilde{t})}{\partial \tilde{t}} \quad (22)$$

where we use notation  $\tilde{w} = cw$ ,  $\tilde{w}_* = c^A w_*$  and  $\tilde{t} = c^B t$ .

## 2.2 Prey Switching Term

In order to demonstrate the scale free assumption, we wish to show that the symmetry transform has the same type of effect upon each additive term of (21). So for example, we hope that when we apply the  $F_c$  transformation upon the term  $\mu_b(w_*) (w^\Omega w_*^\Gamma \Delta)^X N(w, w_*, t)^{X+1}$  within Equation (21), we get  $c^Y \mu_b(\tilde{w}_*) N(\tilde{w}, \tilde{w}_*, \tilde{t})$ , where (we hope)  $Y = \Lambda + B$ , so that the new factors  $c^{\Lambda+B}$  we introduce into Equation (21) will cancel. Our strategy will be to show that this type of effect occurs for all terms of Equation (21), and so after we have performed our  $F_c$  transformation we are left with a new expression that satisfies an equation of the same form as Equation (21), except in terms of the new, rescaled variables  $\tilde{w} = cw$ ,  $\tilde{w}_* = c^A w_*$  and  $\tilde{t} = c^B t$ . In order to other terms change when we apply our  $F_c$  transform, and since these terms are mostly written in terms of different functions of integrals, we shall consider how the more elementary functions describing energy transfer etc. change under the  $F_c$  transform. We start by how the 'prey switching modification factor'

$$\hat{N}(w, w'_*, t)^X = (N(w, w_*, t) w^\Omega w_*^\Gamma \Delta)^X \quad (23)$$

gets transformed under the  $F_c$  symmetry transform, this gives

$$(24)$$

$$\begin{aligned}
\hat{N}(w, w'_*, t)^\chi &\mapsto (c^\Lambda N(cw, c^A w_*, c^B t) w^\Omega w_*^\Gamma \Delta)^\chi \\
c^\Lambda N(cw, c^A w_*, c^B t) (w^\Omega w_*^\Gamma \Delta)^\chi &= \left( c^\Lambda N(\tilde{w}, \tilde{w}_*, \tilde{t}) \left( \frac{\tilde{w}}{c} \right)^\Omega \left( \frac{\tilde{w}_*}{c^A} \right)^\Gamma \Delta \right)^\chi \\
&= c^{\chi\Lambda - \chi\Omega - \chi\Gamma} \hat{N}(\tilde{w}, \tilde{w}_*, \tilde{t})^\chi \\
&= c^r \hat{N}(\tilde{w}, \tilde{w}_*, \tilde{t})^\chi,
\end{aligned}$$

where we define  $r = \chi\Lambda - \chi\Omega - \chi\Gamma$ .

We shall use expression (24) for the transformation of the prey switching modification factor a few times. For example, we can use it to determine how the factor

$$(w^\Omega w_*^\Gamma \Delta)^\chi N(w, w_*, t)^{\chi+1} = N(w, w_*, t) \hat{N}(w, w'_*, t)^\chi \quad (25)$$

(which multiplies the specific mortality rates in Equation (21)) is transformed under the  $F_c$  operation. In this case we see

$$(w^\Omega w_*^\Gamma \Delta)^\chi N(w, w_*, t)^{\chi+1} = N(w, w_*, t) \hat{N}(w, w'_*, t)^\chi \mapsto c^{\Lambda+r} N(\tilde{w}, \tilde{w}_*, \tilde{t}) \hat{N}(\tilde{w}, \tilde{w}_*, \tilde{t})^\chi \quad (26)$$

where

$$c^{\Lambda+r} N(\tilde{w}, \tilde{w}_*, \tilde{t}) \hat{N}(\tilde{w}, \tilde{w}_*, \tilde{t})^\chi = c^{\Lambda+r} (\tilde{w}^\Omega \tilde{w}_*^\Gamma \Delta)^\chi N(\tilde{w}, \tilde{w}_*, \tilde{t})^{\chi+1}. \quad (27)$$

### 2.3 Background Mortality

Applying the symmetry transform to the background mortality rate gives

$$\mu_b(w_*) \mapsto \mu_0 w_*^Q = \mu_0 \frac{\tilde{w}_*^Q}{c^{AQ}} = c^{-AQ} \mu_b(\tilde{w}_*). \quad (28)$$

We can use hence use expressions (26) and (28) together to get the following expression for how the total backfound mortality is transformed under  $F_c$  :

$$\mu_b(w_*) (w^\Omega w_*^\Gamma \Delta)^\chi N(w, w_*, t)^{\chi+1} \mapsto c^{\Lambda+r-AQ} \mu_b(\tilde{w}_*) (\tilde{w}^\Omega \tilde{w}_*^\Gamma \Delta)^\chi N(\tilde{w}, \tilde{w}_*, \tilde{t})^{\chi+1} \quad (29)$$

Hence the effect of the  $F_c$  transformation is to convert the additive terms  $\frac{\partial N(w, w_*, t)}{\partial t}$  and  $\mu_b(w_*) (w^\Omega w_*^\Gamma \Delta)^\chi N(w, w_*, t)^{\chi+1}$  of Equation 21 into the additive terms  $c^{\Lambda+B} \frac{\partial N(\tilde{w}, \tilde{w}_*, \tilde{t})}{\partial \tilde{t}}$  and  $c^{\Lambda+r-AQ} \mu_b(\tilde{w}_*) (\tilde{w}^\Omega \tilde{w}_*^\Gamma \Delta)^\chi N(\tilde{w}, \tilde{w}_*, \tilde{t})^{\chi+1}$  respectively. In order for the transformed function to satisfy 21, with respect to the rescaled variables  $\tilde{w} = cw$ ,  $\tilde{w}_* = c^A w_* \tilde{t} = c^B t$  the new factors, multiplying these terms, that our symmetry transform introduced, must cancel out. In other words, we must have  $c^{\Lambda+B} = c^{\Lambda+r-AQ}$ , for each  $c > 0$ , so we must have  $Q = \frac{r-B}{A}$ .

## 2.4 Available Energy

Before we consider how the rest of the mortality terms transform, we shall consider how the available energy transforms under  $F_c$  as follows

(30)

$$\begin{aligned}\phi(w) &\mapsto \int_0^\infty \int_0^\infty w' c^{\Lambda+r} N(\tilde{w}', \tilde{w}'_*, \tilde{t}) \hat{N}(\tilde{w}', \tilde{w}'_*, \tilde{t})^\chi s(w', w) dw' dw'_* \\ &= \int_0^\infty \int_0^\infty \frac{\tilde{w}'}{c} c^{\Lambda+r} N(\tilde{w}', \tilde{w}'_*, \tilde{t}) \hat{N}(\tilde{w}', \tilde{w}'_*, \tilde{t})^\chi s(\tilde{w}', \tilde{w}) \frac{d\tilde{w}'}{c} \frac{d\tilde{w}'_*}{c^A} \\ &= c^{\Lambda+r-2-A} \phi(\tilde{w})\end{aligned}$$

where we changed our integration variables to  $\tilde{w}' = cw'$  and  $\tilde{w}'_* = c^A w'_*$  and used the fact  $s(w', w) = s(cw', cw) = s(\tilde{w}', \tilde{w})$ .

## 2.5 Encountered Energy

The encountered energy  $E_e(w)$  transforms under  $F_c$  as

$$E_e(w) \mapsto c^{\Lambda+r-2-A} \phi(\tilde{w}) \gamma w^q = c^{\Lambda+r-2-A-q} E_e(\tilde{w}) = c^{\hat{e}} E_e(\tilde{w}), \quad (31)$$

where we define  $\hat{e} = \Lambda + r - 2 - A - q$ .

## 2.6 Feeding level

The feeding level  $f(w)$  transforms under  $F_c$  as

$$f(w) \mapsto \frac{c^{\hat{e}} E_e(\tilde{w})}{c^{\hat{e}} E_e(\tilde{w}) + c^{-n} h \tilde{w}^n}, \quad (32)$$

and by looking at where  $f(w)$  appears in our different terms, we come to the conclusion that our scale free assumption implies that we must have  $f(w) \mapsto c^Z f(\tilde{w})$  for some  $Z$ . However, by looking at the transformed expression (32), we see that this can only be so if all our multipliers cancel out. In other words, we must have  $\hat{e} = -n$ , and this implies that we simply have that  $f(w) \mapsto f(\tilde{w})$ .

## 2.7 Energy For Growth And Reproduction

The energy for growth and reproduction  $E_g(w)$  transforms under  $F_c$  as

$$E_g(w) \mapsto \alpha f(\tilde{w}) h \frac{\tilde{w}^n}{c^n} - k \frac{\tilde{w}^p}{c^p} \quad (33)$$

and so our scale invariance assumption implies that we must have  $n = p$  and

$$E_g(w) \mapsto c^{-n} E_g(\tilde{w}) \quad (34)$$



## 2.8 Growth Rate

The growth rate  $g(w, w_*)$  is zero when  $E_g(w) \leq 0$  and otherwise, the growth rate has the form  $g(w, w_*) = (1 - \psi(w, w_*)) E_g(w)$ . In order for this to satisfy our scale invariance assumption, the reproduction function  $\psi(w, w_*)$  must transform under  $F_c$  so that

$$\psi(w, w_*) \mapsto \psi(\tilde{w}, \tilde{w}_*) \quad (35)$$

and it follows that our growth rate transforms so (in general, irrespective of the sign of  $E_g(w)$ ), we have that the growth rate transforms under  $F_c$  so that

$$g(w, w_*) \mapsto c^{-n} g(\tilde{w}, \tilde{w}_*). \quad (36)$$

We can now see that the term of our central equation (21) involving a weight derivative transforms under  $F_c$  as

$$\frac{\partial[g(w, w_*)N(w, w_*, t)]}{\partial w} \mapsto c^{\Lambda-n} \frac{\partial[g(\tilde{w}, \tilde{w}_*)N(\tilde{w}, \tilde{w}_*, t)]}{\partial \tilde{w}} = c^{\Lambda-n+1} \frac{\partial[g(\tilde{w}, \tilde{w}_*)N(\tilde{w}, \tilde{w}_*, t)]}{\partial \tilde{w}} \quad (37)$$

We want the new factor  $c^{\Lambda-n+1}$  that is introduced here to cancel with the factor  $c^{\Lambda+B}$  that was introduced when we transformed the time derivative (22), and hence we require  $n = 1 - B$ .

## 2.9 Starvation Mortality Rate

The per capita starvation mortality rate

$$\mu_s(w) = \begin{cases} 0 & \text{if } E_g(w) > 0 \\ -\frac{E_g(w)}{\xi w} & \text{otherwise} \end{cases} \quad (38)$$

this transforms under  $F_c$  as

$$\mu_s(w) \mapsto \begin{cases} 0 & \text{if } E_g(\tilde{w}) > 0 \\ -\frac{c^{-n} E_g(\tilde{w})}{\xi} \frac{c}{\tilde{w}} & \text{otherwise} \end{cases} \quad (39)$$

and so we have

$$\mu_s(w) \mapsto c^{1-n} \mu_s(\tilde{w}) \quad (40)$$

So the total starvation rate appearing as a term in (21)

$$\mu_s(w) (w^\Omega w_*^\Gamma \Delta)^\chi N(w, w_*, t)^{\chi+1} = \mu_s(w) N(w, w_*, t) \hat{N}(w, w'_*, t)^\chi$$

transforms under  $F_c$  as

$$(41)$$

$$\begin{aligned}\mu_s(w)N(w, w_*, t)\hat{N}(w, w'_*, t)^\chi &\mapsto c^{1-n}\mu_s(\tilde{w})c^{\Lambda+r}N(\tilde{w}, \tilde{w}_*, \tilde{t})\hat{N}(\tilde{w}, \tilde{w}_*, \tilde{t})^\chi \\ &= c^{\Lambda+r+1-n}\mu_s(\tilde{w})\left(\tilde{w}^\Omega \tilde{w}_*^\Gamma \Delta\right)^\chi N(\tilde{w}, \tilde{w}_*, \tilde{t})^{\chi+1}\end{aligned}$$

and so, since the same factor must be introduced here as in our considerations of the time derivative (Equation (22)), we must have  $\Lambda + r + 1 - n = \Lambda + B$ , it is interesting to compare this to our previous observation that  $c^{\Lambda+B} = c^{\Lambda+r-AQ}$ , since this means  $Q = \frac{r-B}{A}$ , I dare say that if  $A = 1$ , and  $Q = n - 1$ , this implies  $r = 0$ . In other words, unless we change the exponents in either the background mortality rate, or the starvation rate, we cannot incorporate prey switching.