Imperial College London

The role of resource supply in shaping ontogenetic growth and allocation in fish

DÓNAL BURNS

Imperial College London donal.burns@imperial.ac.uk

Submitted: August 27^{th} 2020

A THESIS SUBMITTED IN PARTIAL FULFILMENT OF THE REQUIREMENTS FOR THE DEGREE OF MASTER OF SCIENCE AT IMPERIAL COLLEGE LONDON

FORMATTED IN THE JOURNAL STYLE OF ECOLOGY LETTERS

SUBMITTED FOR THE MSC IN COMPUTATIONAL METHODS IN ECOLOGY AND EVOLUTION

 ${\bf Declaration}$

I declare this project as my own work. The model presented here was developed in conjunction with my

supervisor, Dr. Samraat Pawar, and Ph.D. students Tom Clegg and Olivia Morris. I was responsible

for any simulations and data presentation.

Word Count: 4835

1

Abstract

- 1 Ontogenetic growth models (OGMs) are one of the main model frameworks used to estimate and
- 2 predict the growth of organisms during ontogeny. However, they make many assumptions which are
- 3 in conflict with empirical data, in particular supply and reproduction. Recent results show that
- 4 reproduction does not scale allometrically as previously assumed, but rather hyperallometrically.
- 5 Additionally, not only OGMs but all growth models have failed to properly take variable supply
- 6 rates into account. They instead assume either optimal or proportions of optimal supply when supply
- ⁷ is known not to scale linearly. I develop a model which implements realistic supply scaling and allows
- 8 for allometric scaling of reproduction. I demonstrate that hyperallometric reproductive scaling is
- 9 dependent on metabolic scaling and that both are dependent upon supply. This shows that supply is
- 10 a factor that cannot be ignored when considering growth and reproduction.

11 Keywords:

- allometry; functional response; growth; intake; life history; metabolic theory; metabolism; reproduction;
- 13 reproductive output; supply

Acknowledgements

I would like to thank my supervisor Dr. Samraat Pawar as well as fellow lab members Tom Clegg and Olivia Moris for giving me so much of their time on weekly, and on occasion more than weekly, basis. I would also like to thank Dr. Diego Barneche for his invaluable feedback and Dr. Van Savage for his assistance with some of the initial model development.

Contents

1	Introduction							
2	Methods							
	2.1	Alteri	ng OGMs to Account for Resource Supply	8				
		2.1.1	Full Growth Equation	8				
		2.1.2	Gain	8				
		2.1.3	Loss	9				
	2.2	Calcu	lating Fitness	10				
		2.2.1	Maximising Reproduction	10				
	2.3	Sensit	ivity Analysis	11				
3	Res	Results						
	3.1	Growt	th and Maturation	12				
	3.2	3.2 Sensitivity Analysis						
		3.2.1	Resource Density	14				
		3.2.2	Metabolic Exponent	14				
		3.2.3	c Values	15				
		3.2.4	Shrinking	15				
4	Discussion 1							
	References							
	Sup	pleme	entary Information	23				
	4.1	Unit (Conversions	23				
		4.1.1	Functional Response $(f(\cdot))$	23				
		4.1.2	Metabolic Cost (B_m)	23				
	4.2	Growt	th Curves	23				
	4.3	.3 Sensitivity Analysis						
		4.3.1	Maturation Time	25				
		4.3.2	Metabolic Exponent (μ)	28				
		4.3.3	Resource Density	29				
		4.3.4	Proportion of Shrinking Allowed	31				

1 Introduction

Body mass plays a major role in determining many biological factors. For example, larger individuals are less vulnerable to predation, have lower mass specific metabolic rates and produce more offspring 15 in their lifetime (Barneche et al., 2018; Craig et al., 2006; Hixon et al., 2014; Magnhagen and Heibo, 16 2001; Marshall et al., 2006; Peters, 1983). By extension, knowing the manner in which body mass 17 changes over an organism's lifetime is the gateway to understanding how many biological rates change 18 throughout ontogeny. The reason for this is that many biological rates scale with mass (Kleiber, 1932). 19 However, despite its importance, relatively little is known about the factors which determine growth 20 trajectories (Arendt, 2011; Marshall and White, 2019). 21 In the case of fish, understanding growth, and the factors that play a role in determining it, is not only insightful from the perspective of understanding the world around us. It can also be used to better 23 manage the many fisheries and marine protected areas around the world (Heino et al., 2013; Lester 24 et al., 2009), an objective which is becoming increasingly important as the oceans' fish stocks continue 25 to be depleted by overfishing. The need to understand growth is compounded by global warming which 26 threatens to alter the structure of marine ecosystems even if left unexploited and in their "natural" 27 state (Bruno et al., 2018). It is already known that metabolic rate is dependant on temperature which 28 in turn affects fish sizes (Brown et al., 2004; Gillooly et al., 2001). This, combined with increasing 29 global temperatures, means that understanding in greater detail how increased metabolic rates may 30 affect growth is useful in population management. 31 To date, many models have been developed to predict and describe the growth of an organism 32 throughout its lifetime. The three main approaches used are the von Bertalanffy model, the dynamic 33 energy budget (DEB) model, and the ontogenetic growth model (OGM), which is the focus of this study (Bertalanffy, 1938; Kooijman, 1986; Pütter, 1920; West et al., 2001). All of these are energetic based 35 models with varying assumptions, key among which is the scaling of resource supply and metabolic 36 rate with mass. 37 One of the best known examples of an OGM is the model developed by West et al. (2001). This model is parameterised around the average energy content of animal tissue and asymptotic mass. 39 Asymptotic mass is the mass at which growth has essentially stopped due to metabolic cost and 40 energy intake equalling each other (Fig. 1a). The model hinges on the scaling between energy intake 41 (m^{0.75}, allometric sub-linear scaling) and maintenance cost (m¹, isometric linear scaling) with mass. 42 In other words, as mass increases, maintenance costs will slowly overtake the intake rate and halt 43 growth (Fig. 1a). The framework used by West et al. (2001) was later developed by Charnov et al. 44 (2001) to take the cost of reproduction into account and allow the estimation of lifetime production of offspring. Hou et al. (2008) developed West et al.'s model further by expanding maintenance cost to include the cost of feeding and digestion (specific dynamic action), synthesis of new tissue, and

activity. In the above OGMs, intake is assumed to scale sub-linearly to the power of 0.75. This is 48 due to the assumption that individuals are consuming at an optimal rate at all times and therefore 49 the only limitation is their ability to make use of that energy. In this case, intake should theoretically scale to the power of 0.75 (see West et al., 1997). However, this is not always the case in the field. 51 It has been shown that, for non-optimal consumption, steeper scaling can occur (Pawar et al., 2012; 52 Peters, 1983). Additionally, OGMs, like many growth and metabolic models, typically use basal or 53 resting metabolic rate to calculate metabolic cost. Resting metabolic rate is the minimal metabolic rate of an organism and is typically thought of as the metabolic rate of the organism when relaxed and 55 at rest. However, it has been shown, once factors such as movement are taken into account, scaling becomes steeper (Weibel et al., 2004). 57 The issue of non-optimal feeding is addressed somewhat by Hou et al. (2011). However, this growth was only investigated as, essentially, a proportion of optimal consumption and does not address a 59 potential change in scaling of intake rate. Another limitation of the models used in previous OGMs is 60 dependence on asymptotic mass. All other values, such as metabolic cost, are then derived in relation 61 to asymptotic mass and intake rate. However, organisms are not born with an inherent restriction 62 on the size they can attain, at least not energetically. If there is surplus energy for a given mass, the 63 organism should be able to grow. Relying on asymptotic mass to define the upper bound of attainable 64 mass does not allow for investigation of the mechanisms that underpin asymptotic mass in reality. 65 Previous OGMs have assumed that reproduction scales isometrically with mass. However, it has been 66 shown that larger fish produce far more offspring than the equivalent mass composed of smaller fish. In other words, a 2kg fish will produce more offspring than two 1kg fish, i.e. reproduction scales 68 hyperallometrically (Barneche et al., 2018). Furthermore, larger fish also use energy more efficiently 69 than multiple smaller ones per unit mass. This is due to their lower mass specific metabolic rate (Brown 70 et al., 2004; Kleiber, 1932; Peters, 1983). Additionally, larger mothers produce larger offspring which are then more likely to survive to adulthood and reproduce (Hixon et al., 2014; Marshall et al., 2006). This, combined with empirical results, has led to doubt regarding metabolic scaling. Rather than 73 metabolic scaling being steeper than supply being the cause of growth stopping (Fig. 1a), instead it is 74 thought that the onset of reproduction is what causes growth to cease (Fig. 1b) (Marshall and White, 75 2019; Sibly and Brown, 2020). 76 With two key assumptions of current OGMs, that reproduction and metabolism scale isometrically, 77 not holding in the field (Barneche et al., 2018; Pawar et al., 2012; Peters, 1983), there is a need 78 to take an unexplored approach to modelling growth. This study focuses on developing how intake 79 is described so as to better reflect the real world. To achieve this, a natural starting point is to 80 model intake as a functional response (Holling, 1959) so as to better reflect real world intake rates 81 in terms of consumed biomass over time. Non-optimal supply is a currently unexplored area within 82 growth modelling. This is likely due to the difficulty of directly measuring intake, especially in the

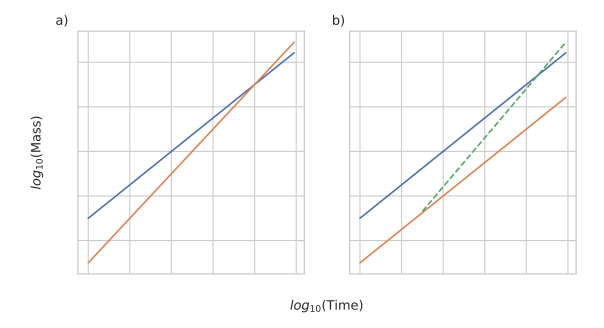


Figure 1: The effect of rate scaling exponents can be visualised within log space. The slope of the line is determined by the exponent. a) shows how maintenance cost out-scales supply in a traditional OGM. Growth only stops when maintenance (scaling exponent = 1) reaches the supply line (scaling exponent = 0.75) b) shows scaling for supply and maintenance as equal. Since scaling is equal growth will never stop until the new cost of reproduction is introduced some time during development. Blue line is supply, orange line is maintenance, green dotted line is reproduction

field. Perhaps as a result, comparatively less is known about consumption. This necessitates the use of proxy values to estimate intake, for example nutrient flux (Schiettekatte et al., 2020), or drawing 85 broad relationships to approximate consumption, as this study will do. Changing the manner in 86 which intake is defined also requires changing metabolic cost, since the two are dependent upon each 87 other in current OGMs. This can be achieved by defining metabolic rate as a value dependent on current mass rather than asymptotic mass, as has been done in OGMs up until this point. This 89 thought process is more mechanistic as an organism has no concept of "How large should I grow?", 90 but rather will acquire as much resources as possible at its current life stage and size. Taking this 91 more bottom-up mechanistic approach also allows exploration of factors which control growth, since 92 as previously mentioned, from an energetic standpoint, an organism can grow indefinitely provided 93 there is surplus energy available after costs have been paid. Of course, there are also mechanical and 94 genetic limitations upon organism size. However, once size is constrained to what is known to exist, 95 this is not an issue. This study takes the novel approach of using a mass-specific functional response and assimilation 97 efficiency to describe how intake changes both throughout ontogeny and varying levels of resource 98 availability. This study focuses on supply and growth within fish. However, the same principles can

be applied to other taxa. 100

Assuming that fish have evolved to maximise reproductive output and can adapt to find an optimal 101 strategy within the constraints of resource density, simulations can be carried out to demonstrate 102 what conditions need to be met in order to achieve hyperallometric scaling of reproduction from 103 an energetic perspective. This study shows that 1) hyperallometric reproduction is dependent upon 104 metabolic scaling exponent, 2) possible scaling of metabolism and reproduction is dependent upon 105 supply and by extension dimensionality.

$\mathbf{2}$ Methods

107

2.1Altering OGMs to Account for Resource Supply

In order to address the issue of supply in the context of an OGM, which can be generically described 108 as dm/dt = gain - loss, some changes need to be made to the model's terms. The first is to remove the 109 assumption of asymptotic mass and the reliance of cost upon it. Within a traditional OGM, the gain 110 term (a) and asymptotic mass are used to define the metabolic cost (b). However, since the assumption 111 of perfect intake is going to be broken, because of variable supply, this relationship no longer holds. 112 As such, both intake and metabolic cost need to be redefined. Additionally, in light of recent work 113 showing that reproduction scales allometrically and not isometrically, the reproductive cost must also 114 be modified from the form used by Charnov et al. (2001) (Barneche et al., 2018; Marshall and White, 115 2019)

2.1.1 **Full Growth Equation** 117

The general form of the model still follows that of an OGM, i.e. dm/dt = gain - loss. The gain term is represented by a functional response $(f(\cdot))$ modified by assimilation efficiency of biomass within poikilotherms (ϵ). Loss is dependent on whether the organism has reached maturity (α) or not. Prior to maturity, loss is resting metabolic rate (B_m) and results in growth as described by Eq. 1. Whereas after maturity, reproductive cost (cm^{ρ}) starts to be considered, resulting in Eq. 2.

$$\frac{dm}{dt} = \epsilon f(\cdot) - B_m \tag{1}$$

$$\frac{dm}{dt} = \epsilon f(\cdot) - B_m \qquad t < \alpha \tag{1}$$

$$\frac{dm}{dt} = \epsilon f(\cdot) - B_m - cm_t^{\rho} \qquad t \ge \alpha \tag{2}$$

2.1.2Gain

118

122

To define supply, a natural starting place is the functional response (Holling, 1959). Functional 119 responses are used to define how much an organism consumes for a given resource density and is 120 described by the following equation: 121

$$f(\cdot) = \frac{aX_r}{1 + ahX_r} \tag{3}$$

where, $f(\cdot)$ is the functional response, a is the search rate, h is handling time and X_r is resource 123 density. For a fixed mass and increasing resource density, Eq. 3 produces a sigmoidal shape with 124

intake eventually reaching an asymptote after some saturating amount of resources is reached. The functional response output is in kg/s. Therefore the units are adjusted to kg/d before use in Eq. 1 and 2 (see SI). At lower resource densities, the intake rate is primarily defined by the search rate, with higher search rates yielding higher intake rates. Conversely, at high resource densities, intake rate is approximately equal to the inverse of the handling time (h^{-1}) , where lower handling times yield higher intake rates.

An organism's functional response will not remain constant throughout its life history. Search rate and handling time are affected by both the organism's mass and how it interacts with its environment (Pawar et al., 2012). Interactions can be broken into 3D and 2D, that is whether the organism consumes from a 2D "surface" e.g. a cow grazing or a 3D "volume" e.g. a pelagic consumer which consumes prey from within the water column. As such, both search rate and handling time can be defined as Eq. 4 and Eq. 5 respectively.

$$a(m) = a_0 m_t^{\gamma} \tag{4}$$

$$h(m) = t_{h,0} m_t^{\beta} \tag{5}$$

A functional response alone is not enough to fully define intake. This is because processing of consumed resources is not one hundred percent efficient which leads to inevitable loss of consumed energy. As a result, to achieve the final gain term, a dimensionless efficiency term ϵ is applied. In poikilotherms assimilation efficiency is roughly 70% (Peters, 1983)

143 **2.1.3** Loss

137

Metabolic cost has previously been dependant upon the gain term within traditional OGMs (see Hou et al., 2008; West et al., 2001). However, for non-maximal intake the relationship will no longer hold true. As a result, this model takes previously measured values as metabolic cost (see Eq. 6 taken from Peters, 1983 and Table 1 for further details). The output of which requires conversion from J/s to kg/d (see SI).

$$B_m = 0.14 m_t^{\mu}$$
 (6)

Next, to take allometric scaling of reproduction into account, the reproductive cost term from Charnov et al. (2001) is changed from cm^1 which assumes isometric scaling to cm^{ρ} . c can be interpreted as the proportion of mass dedicated to reproduction, i.e. the gonadosomatic index of the fish (Charnov et al., 2001). Just as in Charnov et al. (2001) reproductive cost is only taken into account once maturity is reached. This means that until a length of time (α) has passed, reproductive cost is considered to be zero.

2.2 Calculating Fitness

156

164

173

At any time (t) a reproducing organism devotes some amount of energy to reproduction. This is the product of the amount of mass dedicated to reproduction (cm^{ρ}) and a declining efficiency term (h_t) which begins at maturity (α) and represents reproductive senescence (Benoît et al., 2018; Stearns, 2000; Vrtîlek et al., 2018). In addition to amount of reproduction, offspring are also subject to mortality (l_t) . By combining the two, lifetime reproductive output can be estimated and is described by the "characteristic equation" (Eq. 7) which represents reproductive output in a non-growing population (Arendt, 2011; Roff, 1993, 2001; Roff, 1984, 1986; Stearns, 1992; Tsoukali et al., 2016)

$$R_0 = \int c m_t^{\rho} h_t l_t \tag{7}$$

Mortality is experienced differently by juvenile $(t < \alpha)$ and reproducing individuals $(t \le \alpha)$. Mortality of offspring prior to maturity is described as a survival rate $l_t = e^{-Z(t)}$ which is an exponentially decreasing function bounded between zero and one. It controls how many offspring make it to maturity. After maturity, survival is again described as an exponential function which takes time to maturity into account, $l_t = e^{-Z(t-\alpha)}$. Reproductive senescence can also be estimated as an exponential function which begins after maturity and declines over time $(e^{-k(t-\alpha)})$, where k is the senescence term. When all values are inserted into the characteristic equation, it results in the equation used by Charnov et al. (2001) with the inclusion of reproductive senescence (Eq. 8).

$$R_0 = c \int_0^\alpha e^{-Z_t} dt \int_\alpha^\infty m_t^\rho e^{-(\kappa + Z)(t - \alpha)} dt$$
 (8)

In Eq. 8, Z represents instantaneous mortality. This rate has been shown to be related to time of maturation in many taxon groups, and within it follows the relationship $\alpha \cdot Z \approx 2$. This can then be rearranged to estimate instantaneous mortality, $Z \approx 2/\alpha$

2.2.1 Maximising Reproduction

It is assumed that evolution will converge on metabolic values which maximise fitness, with fitness 178 being defined as how much an individual is able to contribute to the gene pool (Speakman, 2008; Stearns, 2000). To this end, lifetime reproductive output is often used as a measure of fitness 180 (Audzijonyte and Richards, 2018; Brown et al., 1993; Charnov and Berrigan, 1991; Charnov et al., 181 2001, 2007; Speakman, 2008; Stearns, 2000; Tsoukali et al., 2016). Therefore, by maximising for 182 reproductive output, it should become clear what parameters will yield the highest fitness. These parameters will then show whether, within a theoretical framework, hyperallometric scaling arises. 184 To find all optimal values for reproduction would require Eq. 8 to be solved analytically. However, 185 since no such solution is possible, I simulated the problem numerically to obtain a result. This was 186 done by simulating across values of c and ρ , the parameters of interest between growth (Eq. 1 and 2) and reproductive output (Eq. 8). c was bound between 0 and 0.4, which encapsulates the values

measured within fish (Benoît et al., 2018; Fontoura et al., 2009; Lambert and Dutil, 2000; Roff, 1983; 189 Wootton, 1985) though c has been shown to reach as much as 0.7 in invertebrate (Parker et al., 2018). 190 To search for any hyperallometry within reproduction, ρ was bound between 0 and 2. The simulation 191 was then run at 0.01 value intervals in both c and ρ over a lifespan of one million days. The results 192 of each simulation were recorded and any non-viable results were discarded. A result was considered 193 non-viable if fish had "shrunk" more than 5% in order to accommodate reproductive costs. Shrinking 194 occurs in the model because the combined loss of energy to metabolism and reproduction is too much 195 for the simulated values at the mass achieved by maturation so the individual experiences a deficit 196 of energy, which is paid by loss in mass until equilibrium is achieved. Shrinking is not expected at 197 maturity in reality. Typically, maturity will occur while the organism still has room for growth. It 198 is the onset of reproduction which is considered to slow or stop growth (see Fig. S1). Shrinking can 199 be thought of as starvation in a real organism. If energetic costs are not met, then energy reserves in 200 the body, such as fat and muscle, are broken down for energy. It has been shown that some fish are 201 capable of losing up to 10% of their body mass (Berghe, 1992). However, this was during the breeding 202 season and caused by behavioural changes due to parenting. Additionally, individuals were shown to 203 rebound back to their "normal" body mass once the breeding season had ended. 204

205 2.3 Sensitivity Analysis

In order to determine the roles of metabolic exponent, maturation time and resource density within the model, sensitivity analyses were performed on each parameter with regard to c and ρ . This was done by simulating the parameters across multiple values and obtaining the optimal value for c and ρ as described above. The parameter values used in the analysis can be seen in Table 1.

Table 1: Parameters used in the model, along with values, units and sources where applicable. The units of resource density change depending on the dimension of intake. m^D represents either m^2 in 2D or m^3 in 3D

Parameter	Description	Value	Units	Range	Source
m	Mass	-	$kg day^{-1}$	-	
B_m	Metabolic Cost	$0.14m^{\mu}$	$kg day^{-1}$	-	Peters, 1983
μ	Metabolic Exponent	-	-	0.75 - 1.0	-
α	Age of maturity	-	day	-	-
c	Reproduction scaling	-	${\rm kg~day^{-1}}$	0 - 0.5	-
	constant				
ho	Reproduction scaling	-	-	0 - 1.5	-
	exponent				
Z	Rate of instantaneous	$2/\alpha$	-	-	Charnov et al.,
	mortality				2001
k	Reproductive	0.01	-	-	
	senescence				
ϵ	Assimilation Efficiency	0.70	-	-	Peters, 1983
X_r	Resource Density	-	$\mathrm{kg/m}^D$	0.11 - 30	-
γ	Search rate scaling	0.68 (2D)	-	-	Pawar et al.,
	exponent				2012
		1.05 (3D)			
a_0	Search rate scaling	$10^{-3.08} (2D)$	$m^2 s^{-1} kg^{-0.68}$	-	Pawar et al.,
	constant				2012
		$10^{-1.77} \text{ (3D)}$	$m^3 s^{-1} kg^{-1.05}$		
β	Handling time scaling	0.75	-	-	Pawar et al.,
	exponent				2012
$t_{h,0}$	Handling time scaling	$10^{3.95} (2D)$	$kg^{1-\beta}$ s	-	Pawar et al.,
	constant				2012
		$10^{3.04} (3D)$	$kg^{1-\beta}$ s		

3 Results

210

3.1 Growth and Maturation

In 3D, when the metabolic scaling exponent (μ) is 1, hyperallometry emerges in reproduction at low resources ,i.e. $\rho > 1$ (Fig. 2b). When resources are high, the value of ρ is lowered (Fig. 2d). This emerges because consumption rate scaling is higher at lower resources in 3D (see Table 1) which allows

for steeper scaling within reproduction. This same pattern occurs within 3D for $\mu = 0.75$ (Fig. 3). 214 In 2D, the opposite pattern is seen for $\mu = 0.75$, with ρ at low resources lower than at saturated 215 resources (Fig. 3a, c). This can again be explained by the difference within supply scaling at high 216 vs. low resources in 2D, since supply scaling is greater at high resources than at low resources in 2D. 217 However, when $\mu = 1$ in 2D, this pattern is reversed (Fig. 2a, c). This may be caused by the very 218 small amount of reproduction occurring at low resources, but if this were the case, the same pattern 219 would be expected to be seen for $\mu = 0.75$, which is not the case. As resources increase, but still 220 remain low, ρ does drop below that of the value at high resources, before climbing back up. However, 221 the relationship is not clear (see Fig. S12 and S14) 222

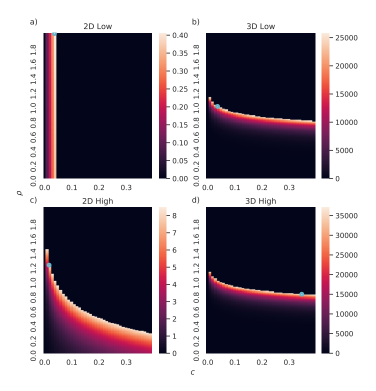


Figure 2: Metabolic exponent of 1 in 2D vs 3D at high and low resource densities. The optimal value is denoted by the blue circle. The resources used for the low resource scenario (top row) is the minimum amount of resources that allows growth with a c and ρ of 0.01. As would be expected, since scaling is steeper in 3D, it allows for growth at smaller resource densities than 2D. Low resources in 2D were $\approx 0.11 \text{kg/m}^2$ and 0.00035kg/m^3 in 3D. 100kg/m^D , where D is dependent on dimension, was used for high resources because, which although unrealistic, ensures that resources are not a limiting factor in the simulations. Hyperallometric scaling is observed in 2D at high (c) and low resources (a). Scaling in 3D is hyperallometric at low resources (b) and hypoallometric at high resources (d). Intensity of colour is determined by reproductive output in kg/s.

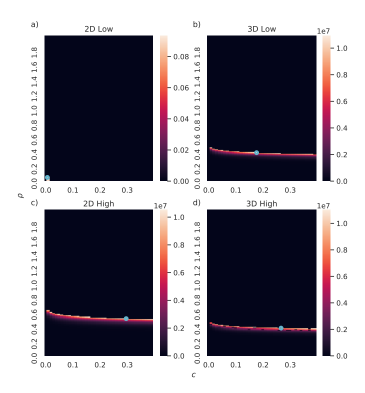


Figure 3: Metabolic exponent of 0.75 in 2D vs 3D at high and low resource densities. The optimal value is denoted by the blue circle. The resources used for the low resource scenario (top row) is the minimum amount of resources that allows growth with a c and ρ of 0.01. As would be expected, since scaling is steeper in 3D, it allows for growth at smaller resource densities than 2D. Low resources in 2D were $\approx 0.11 \text{kg/m}^2$ (a) and 0.00035kg/m^3 in 3D (b). 100kg/m^D , where D is dependent on dimension, was used for high resources because, which although unrealistic, ensures that resources are not a limiting factor in the simulations (c, d). Hypoallometric scaling is observed in all cases. Intensity of colour is determined by reproductive output in kg/s.

3.2 Sensitivity Analysis

3.2.1 Resource Density

224

The scaling relationship of ρ emerges as would be expected from the scaling of the functional response. At low resource densities, the output of the functional response will scale similarly to search rate, the scaling of which is higher in 3D (see Table 1), as resources increase and the response shifts to scaling similarly to the inverse of handling time. At this point ρ starts to take values which are higher in 2D than 3D, because of the higher normalisation constant in 2D.

3.2.2 Metabolic Exponent

The expected result is for increasing values of μ for ρ to also increase. This is because the lower values of μ will result in there being a larger gap between the scaling of intake and maintenance, which

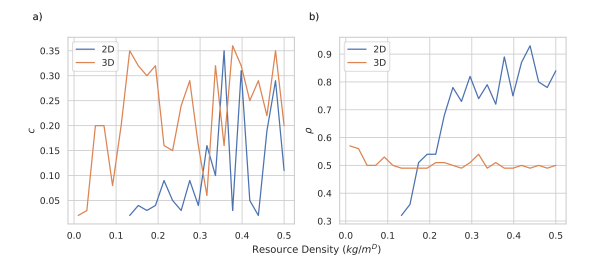


Figure 4: Effect of resource density on c and ρ where $\mu=0.75$. Demonstrates the expected trend that under limiting resources, the higher scaling of 3D search rate allows for steeper reproductive scaling (Table 1). As resources increase and supply shifts more towards being defined by the inverse of handling time, steeper scaling in 2D allows for higher ρ values. Units are kg/m^D, where D is the dimension.

allows for steeper scaling in reproduction (see Fig. 1b) (Marshall and White, 2019). However, the trend suggests that increasing μ allows for higher values of ρ . This is due to numerical instability for greater values of ρ

236 **3.2.3** *c* Values

Estimations of c are low in some cases, especially in 2D (for example Fig. S5 and Fig. S18). While this may be low compared to the $\sim 10\%$ - 35% expected (Benoît et al., 2018; Fontoura et al., 2009; Roff, 1983; Wootton, 1985), it is not unprecedented for values of 2% to be observed in some fish (Gunderson, 1997). It may be necessary for the lower bounds of c to be adjusted based on what is expected or even viable in the organisms being simulated.

242 **3.2.4** Shrinking

When $\mu = 1$, increasing proportions of shrinking allows for higher values of ρ . This is with the 243 exception of low resources in 2D (Fig. S17). This is because at limited resources, with scaling that is 244 lower than that of μ , there is no leeway for steeper scaling of reproduction. In contrast, at very low 245 resources in 3D, increased shrinking does allow for steeper reproductive scaling (Fig. S19), because of 246 the steeper scaling in supply (Table 1). 247 When $\mu = 0.75$, larger proportions of shrinking has no effect. This is because supply scaling is equal 248 to or greater than μ in all cases, meaning greater shrinking does not create "space" for ρ to scale 249 steeper. The exception to this is at low resources in 2D (Fig. S18), where some effect is seen because 250 of the low scaling of intake at low resources in 2D (Table 1). 251

4 Discussion

This study shows that supply plays a critical role in determining the growth and reproductive output 252 of an organism. I show that ρ is not only dependent upon resource density, but whether feeding occurs 253 in two or three dimensions. The results shown in Fig. 3 indicate that reproduction is hypoallometric 254 when $\mu = 0.75$. This is contrary to what should happen theoretically, since the larger "space" between 255 metabolism and supply allows for higher values of ρ . However, these results are misleading. The 256 equal scaling of metabolism and supply results in rapid growth which does not slow down without 257 other additional costs. When reproduction is introduced at such high masses $(m > 3 \times 10^{17})$, the 258 system becomes extremely unstable. Accordingly, the result cannot be taken as a "resonable" growth 259 trajectory and is discounted. Sensitivity analysis of maturation time shows that when maturation is 260 early, which effectively restricts growth, steeper reproductive scaling is possible numerically (e.g. Fig. 261 S4), however more investigation is required. The sensitivity analysis also highlights scenarios in which 262 shrinking can lead to steeper reproductive scaling. Prior to maturity metabolism and supply can be 263 scaling parallel to each other (Fig. 1b) or towards each other (Fig. 1a). At maturity, shrinking has 264 the effect of allowing mass to decrease, and for growth to effectively reverse. This opens more "space" 265 for reproduction to be steeper causing gain and loss to equal each other at the smaller mass (Fig. 1b). 266 When scaling of supply and metabolism are equal, the decrease in mass has no effect. However, if 267 maintenance has steeper scaling than supply then shrinking allows for larger values of ρ . 268 Growth within the model is fast, with asymptotic mass being reached within ~ 10 days (Fig. S3 - S8). 269 This is of course not representative of the real world, where individuals generally need several months 270 to years to reach maturity. The rapid growth may be due to several factors. First is that metabolic 271 cost may be underestimated. Similar to West et al. (2001), this study used resting metabolic rate to 272 define metabolic costs. However, this does not take other costs into account, such as digestion and 273 locomotion. This was addressed in traditional OGMs by Hou et al. (2008). However, due to the use 274 of asymptotic mass in the parameterisation of this change, the same changes could not be used in this 275 model. Resting metabolic rate and active metabolic rates do not scale in the same way with mass (Gillooly et al., 2001; Weibel et al., 2004). The additional cost of active metabolic rate would cause 277 a steeper scaling within the metabolic cost term, leading to more gradual growth (Fig. 1). As such, 278 inclusion of active metabolic rates, while challenging to measure directly and implement, is needed. 279 It is also possible that active costs will not scale constantly for all sizes of fish, as larger individuals 280 incur less drag in the water than smaller ones (Müller et al., 2000). Taking the above factors into 281 account would result in a greater metabolic cost from birth. This would reduce the speed of growth 282 and likely resolve the numerical instability within this studies results of this model at low μ values. 283 There may be behavioural or physiological factors that would lead to an altered metabolic rate. In this 284 regard, temperature plays a critical role. It is well documented that a change in temperature alters 285

many biological rates (see Peters (1983), Gillooly et al. (2001) etc.). It has been shown that growth is 286 dependent on temperature within fish. For example, a 1.5°C increase in sea temperatures could result 287 in a 15% decrease in fish lengths (Rijn et al., 2017). The functional response data used in this study 288 is standardised around 15°C (Pawar et al., 2012). Meanwhile the metabolic cost is for an unspecified 289 temperature (Peters, 1983). Using rates where the temperature effect is taken into account is crucial 290 for model accuracy, given the variation in rates that occurs over different temperatures. Work such as 291 Barneche et al. (2014) has investigated this effect. However, the estimate for metabolic rate is several 292 orders of magnitude lower than what was reported by Peters (1983), which is the rate used in this 293 study. Thus, further investigation is required. 294 In fish, metabolic rate has also been shown to drop under starvation (Cook et al., 2000). In homeotherms, 295 feeding restriction has been shown to lower body temperature, since metabolic rate and core temperature 296 are closely related in homeotherms (Ballor, 1991; Blanc et al., 2003). 297 Another possible point of error is the estimates for supply. The parameters used from Pawar et al. 298 (2012) are for a spectrum of animals from mammals to insects. It is possible by reanalysing the data 299 for only marine species, or more specifically only within taxon or species, predictions of supply could 300 be improved (Marshall and White, 2019). 301 A factor that is not taken into account in this model is that resources are not constant over time. 302 This can be implemented by varying resource density over time. The functional response will respond 303 accordingly giving intake which varies through time. One concern with implementing such a response 304 is fluctuations are not experienced by all organisms in the same way. For a fish with a small range, 305 a local fluctuation can be measured and described relatively simply. However, for a fish with a very 306 large range, there is the possibility of leaving resource poor areas in search of richer waters. The speed 307 at which growth occurs in the model limits comparisons and testing that can be done with lab or field 308 data. However, the scaling relationships and patterns demonstrated here remain true. 309 In conclusion, this study demonstrates the need for more consideration for supply within growth 310 models. Additionally, I provide direction where the model can be expanded upon that was not possible 311 with OGMs, allowing for much more controlled and detailed explanations of the factors controlling 312 growth. In contrast to all previous work, which assume optimal supply, the concept of varying supply 313 is addressed using functional responses. Additionally, qualitative evidence is provided supporting 314 hyperallometric scaling in fish using energy budget as the basis. The model can easily be applied to 315 any animal taxon, not just fish, with some simple changes. Furthermore, there are clear directions to 316

318 Code and Data Availability

317

Code is available at: https://github.com/Don-Burns/Masters_Project

be explored in order to improve the model's accuracy.

20 References

- Arendt, Jeffrey D. (2011). "Size-fecundity relationships, growth trajectories, and the temperature-size
- rule for ectotherms". In: Evolution 65.1, pp. 43-51. DOI: 10.1111/j.1558-5646.2010.01112.x.
- Audzijonyte, Asta and Shane A. Richards (2018). "The energetic cost of reproduction and its effect on
- optimal life-history strategies". In: American Naturalist 192.4, E150–E162. DOI: 10.1086/698655.
- Ballor, D. L. (1991). "Effect of dietary restriction and/or exercise on 23-h metabolic rate and body
- composition in female rats". In: Journal of Applied Physiology 71.3, pp. 801–806. DOI: 10.1152/
- jappl.1991.71.3.801.
- Barneche, D. R., M. Kulbicki, S. R. Floeter, A. M. Friedlander, J. Maina, and A. P. Allen (2014).
- "Scaling metabolism from individuals to reef-fish communities at broad spatial scales". In: Ecology
- Letters 17.9, pp. 1067-1076. DOI: 10.1111/ele.12309.
- Barneche, Diego R, D. Ross Robertson, Craig R White, and Dustin J Marshall (May 2018). "Fish
- reproductive-energy output increases disproportionately with body size". In: Science 360.6389,
- pp. 642-645. DOI: 10.1126/science.aao6868.
- Benoît, Hugues P., Douglas P. Swain, Jeffrey A. Hutchings, Derek Knox, Thomas Doniol-Valcroze,
- and Christina M. Bourne (2018). "Evidence for reproductive senescence in a broadly distributed
- harvested marine fish". In: Marine Ecology Progress Series 592, pp. 207–224. DOI: 10.3354/
- meps12532.
- Berghe, Eric P. van den (1992). "Parental care and the cost of reproduction in a Mediterranean fish".
- In: Behavioral Ecology and Sociobiology 30.6, pp. 373–378. DOI: 10.1007/BF00176171.
- 340 Bertalanffy, Ludwig von (1938). "A QUANTITATIVE THEORY OF ORGANIC GROWTH (INQUIRIES
- ON GROWTH LAWS. II)". In: *Human Biology* 10.2, pp. 181–213.
- Blanc, Stéphane, Dale Schoeller, Joseph Kemnitz, Richard Weindruch, Ricki Colman, Wendy Newton,
- Kelly Wink, Scott Baum, and Jon Ramsey (2003). "Energy expenditure of rhesus monkeys subjected
- to 11 years of dietary restriction". In: Journal of Clinical Endocrinology and Metabolism 88.1,
- pp. 16-23. DOI: 10.1210/jc.2002-020405.
- Brown, James H., James F. Gillooly, Andrew P. Allen, Van M. Savage, and Geoffrey B. West (July
- 2004). "TOWARD A METABOLIC THEORY OF ECOLOGY". In: Ecology 85.7, pp. 1771–1789.
- DOI: 10.1890/03-9000.
- Brown, James H., Pablo A. Marquet, and Mark L. Taper (Oct. 1993). "Evolution of Body Size:
- Consequences of an Energetic Definition of Fitness". In: The American Naturalist 142.4, pp. 573–
- 584. DOI: 10.1086/285558.
- Bruno, John F., Amanda E. Bates, Chris Cacciapaglia, Elizabeth P. Pike, Steven C. Amstrup, Ruben
- Van Hooidonk, Stephanie A. Henson, and Richard B. Aronson (2018). "Climate change threatens

- the world's marine protected areas". In: Nature Climate Change 8.6, pp. 499-503. DOI: 10.1038/ 354 s41558-018-0149-2. 355
- Charnov, Eric L. and David Berrigan (1991). "Evolution of life history parameters in animals with 356 indeterminate growth, particularly fish". In: Evolutionary Ecology 5.1, pp. 63-68. DOI: 10.1007/ 357

BF02285246.

358

361

- Charnov, Eric L., Thomas F. Turner, and Kirk O. Winemiller (2001). "Reproductive constraints and 359 the evolution of life histories with indeterminate growth". In: Proceedings of the National Academy 360 of Sciences of the United States of America 98.16, pp. 9460-9464. DOI: 10.1073/pnas.161294498.
- Charnov, Eric L., Robin Warne, and Melanie Moses (2007). "Lifetime reproductive effort". In: American 362 Naturalist 170.6. DOI: 10.1086/522840. 363
- Cook, J. T., A. M. Sutterlin, and M. A. McNiven (2000). "Effect of food deprivation on oxygen 364 consumption and body composition of growth-enhanced transgenic Atlantic salmon (Salmo salar)". 365 In: Aquaculture 188.1-2, pp. 47-63. DOI: 10.1016/S0044-8486(00)00333-1. 366
- Craig, J. Kevin, Brian J. Burke, Larry B. Crowder, and James A. Rice (2006). "Prey growth and size-367 dependent predation in juvenile estuarine fishes: Experimental and model analyses". In: Ecology 368 87.9, pp. 2366-2377. DOI: 10.1890/0012-9658(2006)87[2366:PGASPI]2.0.C0;2. 369
- Fontoura, Nelson F., Aloísio S. Braun, and Paulo Cesar C. Milani (2009). "Estimating size at first 370 maturity (L50) from Gonadossomatic Index (GSI) data". In: Neotropical Ichthyology 7.2, pp. 217– 371 222. DOI: 10.1590/S1679-62252009000200013. 372
- Gillooly, J. F., J. H. Brown, G. B. West, V. M. Savage, and E. L. Charnov (2001). "Effects of size and 373 temperature on metabolic rate". In: Science 293.5538, pp. 2248-2251. DOI: 10.1126/science. 374 1061967. 375
- Gunderson, Donald R. (1997). "Trade-off between reproductive effort and adult survival in oviparous 376 and viviparous fishes". In: Canadian Journal of Fisheries and Aquatic Sciences 54.5, pp. 990–998. DOI: 10.1139/f97-019. 378
- Heino, Mikko et al. (July 2013). "Can fisheries-induced evolution shift reference points for fisheries 379 management?" In: ICES Journal of Marine Science 70.4, pp. 707-721. DOI: 10.1093/icesjms/ 380 fst077. 381
- Hixon, Mark a, Darren W Johnson, and Susan M Sogard (Oct. 2014). "BOFFFFs: on the importance 382 of conserving old-growth age structure in fishery populations". In: ICES Journal of Marine Science 383 71.8, pp. 2171-2185. DOI: 10.1093/icesjms/fst200. 384
- Holling, C. S. (1959). "Some Characteristics of Simple Types of Predation and Parasitism". In: The 385 Canadian Entomologist 91.7, pp. 385-398. DOI: 10.4039/Ent91385-7. 386
- Hou, Chen, Kendra M. Bolt, and Aviv Bergman (2011). "A general model for ontogenetic growth under 387 food restriction". In: Proceedings of the Royal Society B: Biological Sciences 278.1720, pp. 2881-388 2890. DOI: 10.1098/rspb.2011.0047. 389

- Hou, Chen, Wenyun Zuo, Melanie E. Moses, William H. Woodruff, James H. Brown, and Geoffrey B.
- West (2008). "Energy uptake and allocation during ontogeny". In: Science 322.5902, pp. 736–739.
- DOI: 10.1126/science.1162302.
- ³⁹³ Kleiber, Max (1932). "California Agricultural Experiment Station Body Size and Metabolism". In:
- Hilgardia Journal of Agricultural Science 6.11, pp. 315–353.
- Kooijman, S. A.L.M. (1986). "Energy budgets can explain body size relations". In: Journal of Theoretical
- Biology 121.3, pp. 269–282. DOI: 10.1016/S0022-5193(86)80107-2.
- Lambert, Yvan and Jean Denis Dutil (2000). "Energetic consequences of reproduction in Atlantic cod
- (Gadus morhua) in relation to spawning level of somatic energy reserves". In: Canadian Journal
- of Fisheries and Aquatic Sciences 57.4, pp. 815–825. DOI: 10.1139/f00-022.
- Lester, Sarah E., Benjamin S. Halpern, Kirsten Grorud-Colvert, Jane Lubchenco, Benjamin I. Ruttenberg,
- Steven D. Gaines, Satie Airamé, and Robert R. Warner (2009). "Biological effects within no-take
- marine reserves: A global synthesis". In: Marine Ecology Progress Series 384, pp. 33–46. DOI:
- 403 10.3354/meps08029.
- Magnhagen, C. and E. Heibo (2001). "Gape size allometry in pike reflects variation between lakes
- in prey availability and relative body depth". In: Functional Ecology 15.6, pp. 754–762. DOI: 10.
- 406 1046/j.0269-8463.2001.00576.x.
- 407 Marshall, C. Tara, Coby L. Needle, Anders Thorsen, Olav Sigurd Kjesbu, and Nathalia A. Yaragina
- (2006). "Systematic bias in estimates of reproductive potential of an Atlantic cod (Gadus morhua)
- stock: Implications for stock-recruit theory and management". In: Canadian Journal of Fisheries
- and Aquatic Sciences 63.5, pp. 980–994. DOI: 10.1139/F05-270.
- Marshall, Dustin J. and Craig R. White (2019). "Have We Outgrown the Existing Models of Growth?"
- In: Trends in Ecology and Evolution 34.2, pp. 102-111. DOI: 10.1016/j.tree.2018.10.005.
- Müller, U. K., E. J. Stamhuis, and J. J. Videler (2000). "Hydrodynamics of unsteady fish swimming
- and the effects of body size: Comparing the flow fields of fish larvae and adults". In: Journal of
- Experimental Biology 203.2, pp. 193–206.
- Parker, Geoff A., Steven A. Ramm, Jussi Lehtonen, and Jonathan M. Henshaw (2018). "The evolution
- of gonad expenditure and gonadosomatic index (GSI) in male and female broadcast-spawning
- invertebrates". In: *Biological Reviews* 93.2, pp. 693–753. DOI: 10.1111/brv.12363.
- Pawar, Samraat, Anthony I. Dell, and Van M. Savage (June 2012). "Dimensionality of consumer
- search space drives trophic interaction strengths". In: Nature 486.7404, pp. 485–489. DOI: 10.
- 421 1038/nature11131.
- Peters, Robert Henry (Oct. 1983). The Ecological Implications of Body Size. Cambridge University
- Press. DOI: 10.1017/CB09780511608551.

- Pütter, August (Dec. 1920). "Studien über physiologische Ähnlichkeit VI. Wachstumsähnlichkeiten".
- In: Pflügers Archiv für die Gesamte Physiologie des Menschen und der Tiere 180.1, pp. 298–340.
- DOI: 10.1007/BF01755094.
- Rijn, Itai van, Yehezkel Buba, John DeLong, Moshe Kiflawi, and Jonathan Belmaker (2017). "Large
- but uneven reduction in fish size across species in relation to changing sea temperatures". In: Global
- *Change Biology* 23.9, pp. 3667–3674. DOI: 10.1111/gcb.13688.
- Roff, D. A. (1983). "An allocation model of growth and reproduction in fish." In: Canadian Journal
- of Fisheries and Aquatic Sciences 40.9, pp. 1395–1404. DOI: 10.1139/f83-161.
- Roff, Derek (1993). The Evolution of Life Histories: Theory and Analysis. Springer Science \& Business
- 433 Media.
- (2001). "Life History, Evolution of". In: Encyclopedia of Biodiversity. Elsevier, pp. 631–641. DOI:
- 10.1016/B978-0-12-384719-5.00087-3.
- Roff, Derek A. (1984). "The evolution of life history parameters in teleosts". In: Canadian Journal of
- Fisheries and Aquatic Sciences 41.6, pp. 989–1000. DOI: 10.1139/f84-114.
- 438 (1986). "Predicting Body Size with Life History Models". In: BioScience 36.5, pp. 316–323. DOI:
- 10.2307/1310236.
- Schiettekatte, Nina M.D. et al. (2020). "Nutrient limitation, bioenergetics and stoichiometry: A new
- model to predict elemental fluxes mediated by fishes". In: Functional Ecology November 2019,
- pp. 1–13. DOI: 10.1111/1365-2435.13618.
- 443 Sibly, R. M. and J. H. Brown (2020). "Toward a physiological explanation of juvenile growth curves".
- In: Journal of Zoology 1, pp. 1–5. DOI: 10.1111/jzo.12770.
- Speakman, John R. (2008). "The physiological costs of reproduction in small mammals". In: *Philosophical*
- Transactions of the Royal Society B: Biological Sciences 363.1490, pp. 375–398. DOI: 10.1098/
- rstb.2007.2145.
- Stearns, Stephen C (1992). The evolution of life histories. 575 S81.
- 449 (2000). "Life history evolution: Successes, limitations, and prospects". In: Naturwissenschaften
- 450 87.11, pp. 476–486. DOI: 10.1007/s001140050763.
- Tsoukali, Stavroula, Karin H. Olsson, Andre W. Visser, and Brian R. Mackenzie (2016). "Adult lifetime
- reproductive value in fish depends on size and fecundity type". In: Canadian Journal of Fisheries
- and Aquatic Sciences 73.9, pp. 1405-1412. DOI: 10.1139/cjfas-2015-0378.
- ⁴⁵⁴ Vrtílek, Milan, Jakub Žák, Radim Blažek, Matej Polačik, Alessandro Cellerino, and Martin Reichard
- (2018). "Limited scope for reproductive senescence in wild populations of a short-lived fish". In:
- *Science of Nature* 105.11-12. DOI: 10.1007/s00114-018-1594-5.
- Weibel, Ewald R., Leonardo D. Bacigalupe, Beat Schmitt, and Hans Hoppeler (2004). "Allometric
- scaling of maximal metabolic rate in mammals: Muscle aerobic capacity as determinant factor". In:
- Respiratory Physiology and Neurobiology 140.2, pp. 115-132. DOI: 10.1016/j.resp.2004.01.006.

- West, Geoffrey B., James H. Brown, and Brian J. Enquist (1997). "A general model for the origin of
- allometric scaling laws in biology". In: Science 276.5309, pp. 122-126. DOI: 10.1126/science.
- 462 276.5309.122.
- (2001). "A general model for ontogenetic growth". In: Nature 413.6856, pp. 628–631. Doi: 10.
- 1038/35098076.
- Wootton, R. J. (1985). "Energetics of Reproduction". In: Fish Energetics, pp. 231–254. DOI: 10.1007/
- 978-94-011-7918-8_9.

Supplementary Information

4.1 Unit Conversions

4.1.1 Functional Response $(f(\cdot))$

$$kg \cdot s^{-1} \cdot 24 \cdot 60 \cdot 60 = kg \cdot d - 1$$
 (S1)

4.1.2 Metabolic Cost (B_m)

Conversion factor for joules to kg wet mass from Weathers et al. (2012).

$$J \cdot s^{-1} \cdot 24 \cdot 60 \cdot 60 = J \cdot d - 1$$
 (S2)
$$J \cdot d - 1 \cdot 2.5 \times 10^{-4} = kg \cdot d - 1$$

4.2 Growth Curves

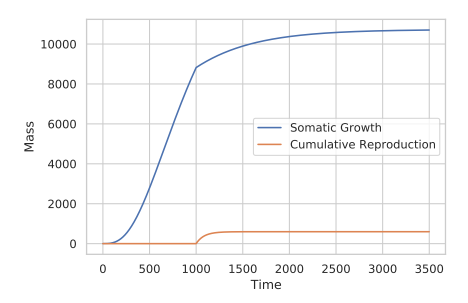


Figure S1: Example of the growth curve and cumulative reproduction expected from a traditional OGM model. Maturation occurs at 1000 days, after which growth is less steep until reaching asymptotic mass.

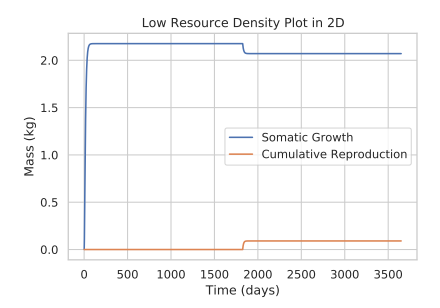


Figure S2: The growth over a fish which consumes in 2D. Maturation occurs at 5 years (1825 days). The fish was allowed to shrink by 5% at the onset of reproduction.

4.3 Sensitivity Analysis

4.3.1 Maturation Time

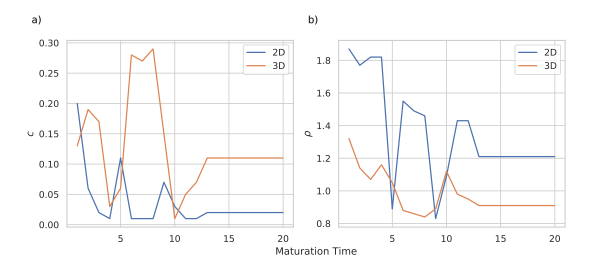


Figure S3: Effect of maturation time on c and ρ where $\mu = 1$ and resource density is high (100 kg/m^D, where D is the dimension).

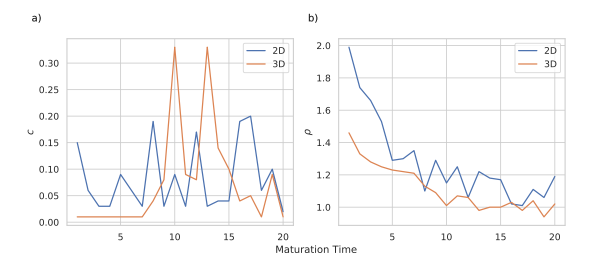


Figure S4: Effect of maturation time on c and ρ where $\mu = 0.75$ and resource density is high (100 kg/m^D, where D is the dimension).

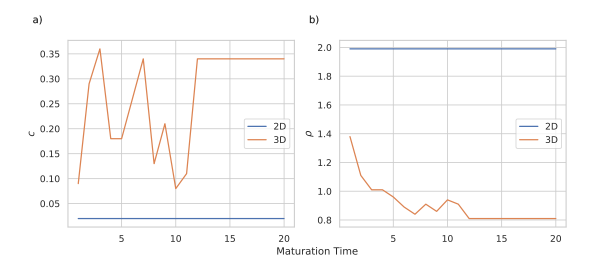


Figure S5: Effect of maturation time on c and ρ where $\mu = 1$ and resource density is low (0.11 kg/m^D, where D is the dimension).

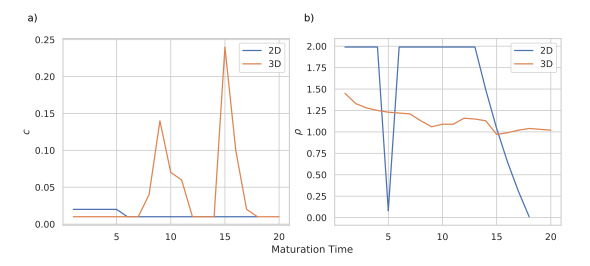


Figure S6: Effect of maturation time on c and ρ where $\mu = 0.75$ and resource density is low(0.11 kg/m^D, where D is the dimension).

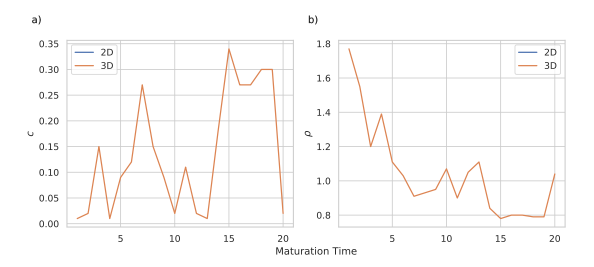


Figure S7: Effect of maturation time on c and ρ where $\mu = 1$ and resource density is very low (0.01 kg/m^D, where D is the dimension). At this resource density reproduction can only occur in 3D.

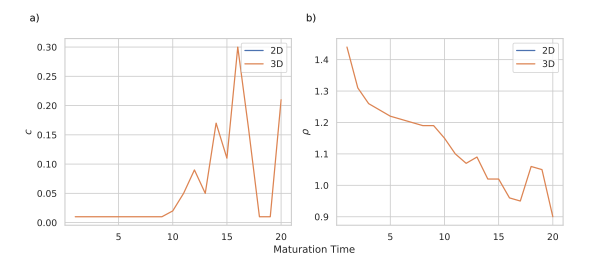


Figure S8: Effect of maturation time on c and ρ where $\mu = 0.75$ and resource density is very low (0.01 kg/m^D, where D is the dimension). At this resource density reproduction can only occur in 3D.

4.3.2 Metabolic Exponent (μ)

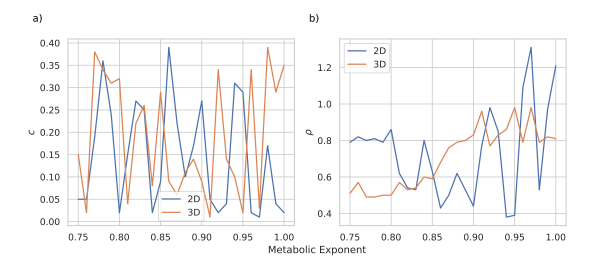


Figure S9: Effect of metabolic on c and ρ where resource density is high (100 kg/m^D, where D is the dimension)

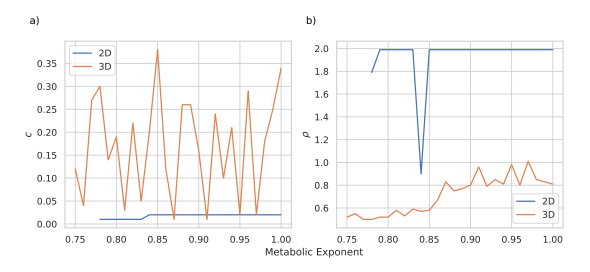


Figure S10: Effect of metabolic on c and ρ where resource density is low (0.11 kg/m^D, where D is the dimension)

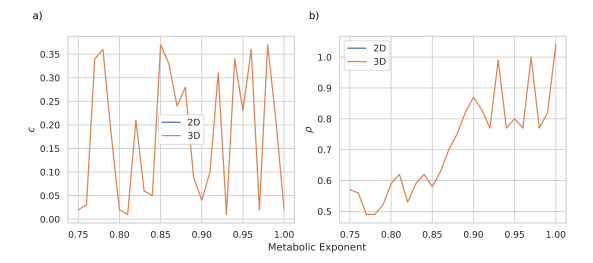


Figure S11: Effect of metabolic on c and ρ where resource density is very low (0.01 kg/m^D, where D is the dimension). At this resource density reproduction can only occur in 3D.

4.3.3 Resource Density

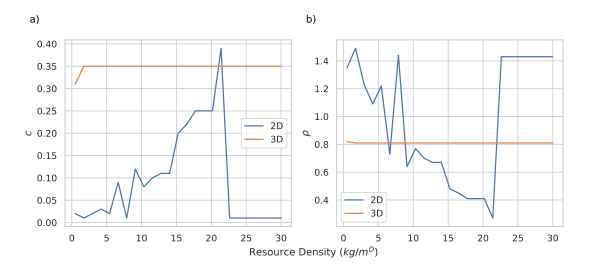


Figure S12: Effect of resource density on c and ρ where $\mu = 1$. Over larger values for resource density. 3D quickly saturates at this density, thus is a nearly straight horizontal line. See Fig. 4 for detail at lower resource density. Units are kg/m^D , where D is the dimension.

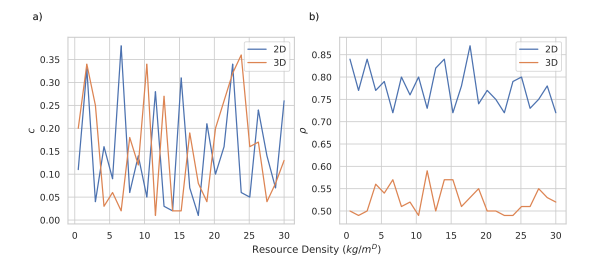


Figure S13: Effect of resource density on c and ρ where $\mu=0.75$. Over larger values for resource density. There is a lot of numeric instability across resource densities, but the trend appears to be somewhat stable around ~ 0.8 in 2D and ~ 0.53 in 3D See Fig. 4 for detail at lower resource density. Units are kg/m^D , where D is the dimension.

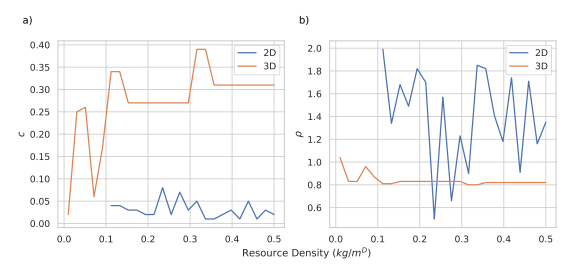


Figure S14: Effect of resource density on c and ρ where $\mu=1$. Demonstrates the expected trend that under limiting resources the higher scaling of 3D search rate allows for steeper reproductive scaling (Table 1). As resources increase and supply shifts more towards being defined by the inverse of handling time, steeper scaling in 2D allows for higher ρ values. Units are kg/m^D, where D is the dimension.

4.3.4 Proportion of Shrinking Allowed

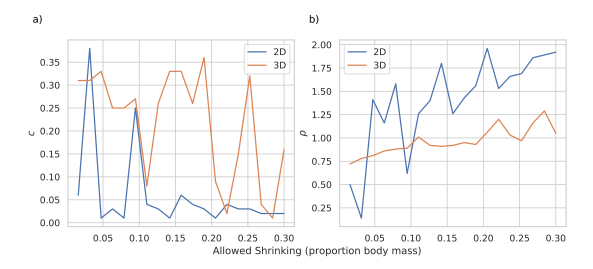


Figure S15: Effect of proportion of shrinking allowed on c and ρ where $\mu = 1$ and resource density is high (100 kg/m^D, where D is the dimension).

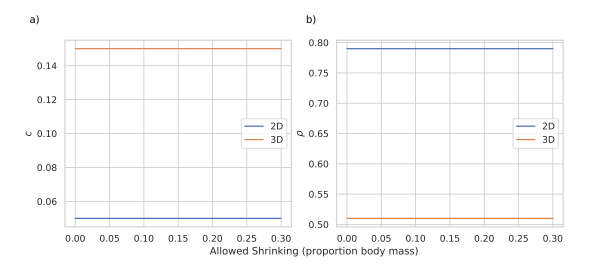


Figure S16: Effect of proportion of shrinking allowed on c and ρ where $\mu = 0.75$ and resource density is high (100 kg/m^D, where D is the dimension).

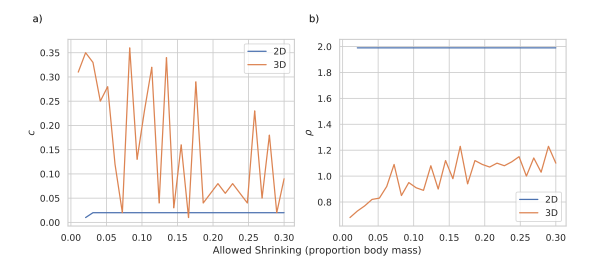


Figure S17: Effect of proportion of shrinking allowed on c and ρ where $\mu = 1$ and resource density is low (0.11 kg/m^D, where D is the dimension)

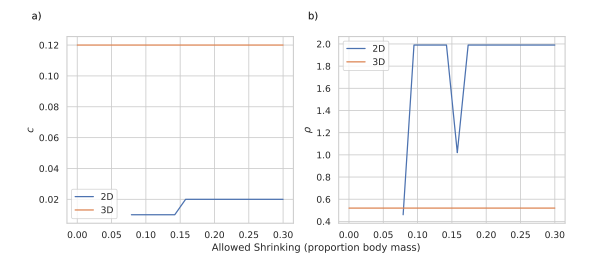


Figure S18: Effect of proportion of shrinking allowed on c and ρ where $\mu = 0.75$ and resource density is low (0.11 kg/m^D, where D is the dimension)

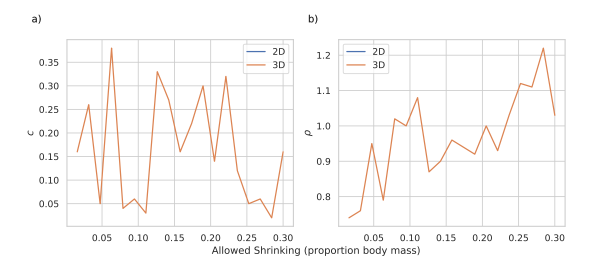


Figure S19: Effect of proportion of shrinking allowed on c and ρ where $\mu = 1$ and resource density is very low (0.01 kg/m^D, where D is the dimension). The resource density only allows for reproduction to occur on 3D.

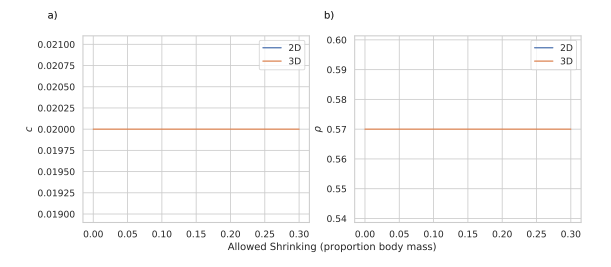


Figure S20: Effect of proportion of shrinking allowed on c and ρ where $\mu = 0.75$ and resource density is very low (0.01 kg/m^D, where D is the dimension). The resource density only allows for reproduction to occur on 3D.

References

Weathers, Kathleen C, David L Strayer, and Gene E Likens (2012). Fundamentals of ecosystem science. Academic Press.