- <sup>1</sup> Chapter 1
- <sup>2</sup> The Pella-Tomlinson Model

3 Abstract

into two dimensions.

Stock assessments often assume a two-parameter functional form (e.g., Beverton-Holt or Ricker) for the expected recruitment produced by a given level of spawning output. Mangel et al. (2013) and others have shown that biological reference points such as  $\frac{F^*}{M}$  and  $\frac{B^*}{\bar{B}(0)}$ are largely determined by a single parameter (steepness) when using two-parameter relationships. These functions introduce strong correlations between reference points that are pre-determined by the functional form, rather than a biological characteristic of the stock. 9 Mangel et al. note that use of a three-parameter stock-recruitment relationship allows for 10 independent estimation of these reference points. This research seeks to understand the 11 nature of biases in reference points resulting from fitting a two-parameter functional form when the true relationship follows a three-parameter stock-recruitment relationship. This 13 work demonstrates the useful limits of misspecified two-parameter models, and suggests the mechanisms of model failure which arise from mapping a three-dimensional parameter space 15

# 17 Introduction

The most fundamental model in modern fisheries management is the surplus-production model. These models focus on modeling population growth via nonlinear parametric ordi-19 nary differential equations (ODE). Key management quantities called reference points (RPs) are commonly derived from the ODE equilibrium equations and depend upon the parameter-21 ization of biomass production. Two-parameter forms of the production function have been 22 shown to limit the theoretical domain of RPs (Mangel et al., 2013). The limited RP-space of 23 two parameter models are a major source of model misspecification for RPs and thus induce bias in RP estimation. The behavior of RP estimation bias is not well understood and as 25 a result often underappreciated. A metamodeling approach is developed here to describe RP biases and explore mechanisms of model failure under the most common two parameter 27 models. 28 Data for a typical surplus-production model comes in the form of an index of abundance 29

Data for a typical surplus-production model comes in the form of an index of abundance through time which is assumed to be proportional to the reproducing biomass for the population of interest. The index is often observed alongside a variety of other known quantities, but at a minimum, each observed index will be observed in the presence of some known catch for the period. Figure (1.1) shows the classic Namibian Hake dataset exemplifying the form.

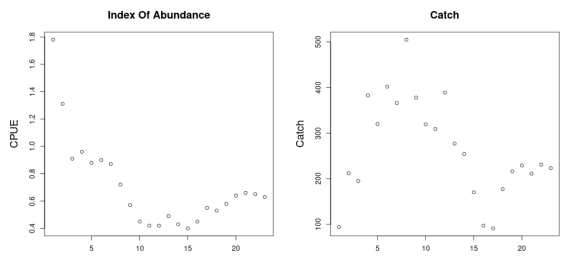


Figure 1.1: *left*: Arl Midex of abundance data, catch per unit Miffort (CPUE), for Namibian Hake from 1965 to 1987 (Hilborn & Mangel, 1997). *right*: The associated catch data for Namibian Hake over the same time period.

Indices are assumed to have multiplicative log-normal errors, and thus the following observation model arises naturally,

$$I_t = qB_t e^{\epsilon} \quad \epsilon \sim N(0, \sigma^2). \tag{1.1}$$

Above q is often referred to as the "catchability parameter"; it serves as the proportionality constant mapping between the observed index of abundance and biomass.  $\sigma^2$  models residual variation. Biologically speaking q and  $\sigma^2$  are often treated as nuisance parameters with the "biological parameters" entering the model through a process model on biomass.

Biomass is assumed to evolve as an ODE; in this case I focus on the following form

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$$\frac{dB}{dt} = P(B(t); \boldsymbol{\theta}) - Z(t)B(t). \tag{1.2}$$

the population, P(B), and various sources of biomass removal, Z, from the population. Firstly, the population grows through a production function, P(B). Production in this setting is defined as the net biomass increase due to all reproduction and maturation processes. The production function is assumed to be a parametric (generally non-linear) function relating the current biomass of the population to an aggregate production of biomass.

Here biomass is assumed to change in time by two processes, net production of biomass into

Secondly, the population decreases as biomass is removed by various sources that are assumed to remove biomass linearly with biomass. Above, Z(t), is an aggregate rate of removal. When the fishing rate, F(t), is the only source of removal Z(t) = F(t), however often models will also included other linear terms in Z(t). Commonly the rate of "natural mortality", M, is also included as an additional term so that Z(t) = M + F(t).

From a management perspective a major goal of modeling is to accurately infer a quantity known as maximum sustainable yield (MSY). One could maximize simple yield at a particular moment in time (and only for that moment) by fishing all available biomass in that moment. This strategy is penny-wise but pound-foolish (not to mention ecologically devastating) since it doesn't leave biomass in the population to reproduce in the future. We seek to fish in a way that allows (or even encourages) future productivity in the population. This is accomplished by maximizing the equilibrium level of catch over time. Equilibrium yield is considered by

replacing the steady state biomass  $(\bar{B})$  in the assumed form for catch, so that  $\bar{Y} = F\bar{B}(F)$ , where  $\bar{}$  indicates a value at steady state. MSY is found by maximizing  $\bar{Y}(F)$  with respect to 58 F, and  $F^*$  is the fishing rate at MSY. Going forward let \* decorate any value derived under 59 the condition of MSY. 60 Fisheries are very often managed based upon reference points which serve as simplified 61 heuristic measures of population behavior. The mathematical form of RPs depends upon 62 the model assumptions through the production function. While a number of different RPs 63 exist which describe the population in different (but related) ways, the most common RPs 64 revolve around the concept of MSY (or robust ways of measuring MSY (Hilborn, 2010; Punt 65 et al., 2016)). Here the focus is primarily on the RPs  $\frac{B^*}{\bar{B}(0)}$  and  $F^*$  ( $\frac{F^*}{M}$  when appropriate) for their pervasive use in modern fisheries (Punt & Cope, 2019). 67  $F^*$  is the afore mentioned fishing rate which results in MSY.  $\frac{B^*}{\overline{B}(0)}$  is the depletion of the 68 stock at MSY. That is to say  $\frac{B^*}{\overline{B}(0)}$  describes the fraction of the unfished population biomass 69 that will remain in the equilibrium at MSY. In general  $F^* \in \mathbb{R}^+$  and  $\frac{B^*}{B(0)} \in (0,1)$ , however 70 under the under the assumption of a two parameter production function production models 71 will be structurally unable to capture the full theoretical range of RPs. 72 Many of the most commonly used production functions depend only on two parameters. 73 For example, the Schaefer model depends only on the biological parameters r and K, and 74 limits RP inference so that under the Schaefer model  $\left(F^*, \frac{B^*}{B(0)}\right) \in \left(\mathbb{R}^+, \frac{1}{2}\right)$ . The two pa-75 rameter Fox model (Fox Jr., 1970) limits  $\left(F^*, \frac{B^*}{\overline{B}(0)}\right) \in \left(\mathbb{R}^+, \frac{1}{e}\right)$ . Similarly the two parameter Cushing (Cushing, 1971), Beverton-Holt (Beverton & Holt, 1957, BH) and Ricker (Ricker, 77 1954) production functions do not model the full theoretical space of RPs (Mangel et al., 78 2013; Yeakel & Mangel, 2015). 79 The bias-variance trade-off (Ramasubramanian & Singh, 2017) makes it clear that the 80 addition of a third parameter in the production function will necessarily reduce estimation 81 bias. However the utility of this bias reduction is still under debate because the particular 82 mechanisms and behavior (direction and magnitude) of these biases for key management 83 quantities are not fully understood or described. Lee et al. (2012) provides some evidence 84 that estimation of productivity parameters are dependent on biomass contrast as well as model specification. Conn et al. (2010) comes to similar conclusions via calibration modeling techniques. These studies indicate important factors that contribute to inferential failure.

However they do not offer mechanisms of model failure, nor do their experimental designs allow for the control of different types of model misspecification.

In this study I consider the behavior of inference when index data are simulated from three parameter PT and Schnute production models, but the simulated data are fit using intentionally misspecified two parameter logistic or BH production models. The work begins with a derivation of RPs under the three parameter models. A method is then presented for generating simulation designs based on the parametric form of RPs which serves as a control on the nature of simulated model misspecification. Finally a Gaussian Process (GP) metamodel (Gramacy, 2020) is constructed for exploration and analysis of RP biases.

A key insight of this approach is that bias is considered broadly across RP-space to uncover patterns and correlations between RPs. The GP metamodel is explicit about tradeoffs between RPs so as to inform the full utility of reducing bias, as well as to suggest mechanisms for understanding what causes bias. Further, the effect of contrast on estimation is considered together with model misspecification.

# <sup>102</sup> 2 Methods

# pprox 2 .1 Pella-Tomlinson Model

The three parameter Pella-Tomlinson (PT) family has a convenient form that includes, among others (Fox Jr., 1970; Rankin & Lemos, 2015), the logistic production function as a special case. PT production function is parameterized so that  $\boldsymbol{\theta} = [r, K, \gamma]$  and the family takes the following form,

$$P_p(B; [r, K, \gamma]) = \frac{rB}{\gamma - 1} \left( 1 - \left( \frac{B}{K} \right)^{(\gamma - 1)} \right). \tag{1.3}$$

 $\gamma$  is a parameter which breaks PT out of the 104 restrictive symmetry of the logistic curve. In gen-105 eral  $\gamma \in (1, \infty)$ , with the logistic model appear-106 ing in the special case of  $\gamma = 2$ , and the Fox model appearing as a limiting case as  $\gamma \to 1$ . The 108 parameter r controls the maximum reproductive 109 rate of the population in the absence of compe-110 tition for resources (i.e. the slope of production 111 function at the origin). K is the so called "carrying capacity" of the population. In this con-113 text the carrying capacity can be formally stated 114 as steady state biomass in the absence of fishing 115 (i.e. B(0) = K). In Figure (3.1) PT recruitment 116 is shown for a range of parameter values so as to demonstrate the various recruitment shapes that 118 can be achieved by PT recruitment. 119

While the form of the PT curve produces some limitations (Fletcher, 1978), importantly

the introduction of a third parameter allows enough flexibility to fully describe the space of reference points used in management. To see this, the reference points are analytically derived for the PT model below.

#### 125 PT Reference Points

With B(t) representing biomass at time t, under PT production, the dynamics of biomass are defined by the following ODE,

$$\frac{dB}{dt} = \frac{rB}{\gamma - 1} \left( 1 - \left( \frac{B}{K} \right)^{\gamma - 1} \right) - FB. \tag{1.4}$$

An expression for the equilibrium biomass is attained by setting Eq (1.4) equal to zero, and rearranging the resulting equation to solve for B. Thinking of the result as a function

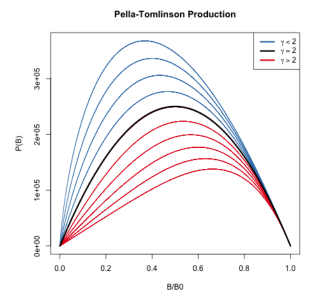


Figure 1.2: The Pella-Tomlinson production function plotted across a variety of parameter values. The special cases of Logistic production is shown in black, and the left-leaning and right-leaning regimes are shown in blue and red respectively.

of F gives,

$$\bar{B}(F) = K \left( 1 - \frac{F(\gamma - 1)}{r} \right)^{\frac{1}{(\gamma - 1)}}.$$
(1.5)

At this point it is convenient to notice that  $\bar{B}(0) = K$ . The expression for  $B^*$  is given by evaluating Eq (1.5) at  $F^*$ . To get an expression for  $F^*$ , the equilibrium yield is maximized with respect to F,

$$F^* = \operatorname*{argmax}_F F\bar{B}(F). \tag{1.6}$$

In the case of PT production this maximization can be done analytically, by differentiating the equilibrium yield with respect to F as follows,

$$\frac{d\bar{Y}}{dF} = \bar{B}(F) + F\frac{d\bar{B}}{dF} \tag{1.7}$$

$$\frac{d\bar{B}}{dF} = -\frac{K}{r} \left( 1 - \frac{F(\gamma - 1)}{r} \right)^{\frac{1}{\gamma - 1} - 1}.$$
(1.8)

Setting Eq (1.7) equal to 0, substituting  $\bar{B}(F)$  and  $\frac{d\bar{B}}{dF}$  by Equations (1.5) and (1.8) respectively, and solving for F produces the following expression for the fishing rate required to produce MSY,

$$F^* = \frac{r}{\gamma} \tag{1.9}$$

Plugging the above expression for  $F^*$  back into Eq (1.5) gives the following expression for biomass at MSY,

$$B^* = K \left(\frac{1}{\gamma}\right)^{\frac{1}{\gamma - 1}}. (1.10)$$

The above derived expressions for  $\bar{B}(0)$ ,  $B^*$ , and  $F^*$  can then be used to build a specific analytical form for the biological reference points in terms of only productivity parameters.

$$F^* = \frac{r}{\gamma} \qquad \frac{B^*}{\bar{B}(0)} = \left(\frac{1}{\gamma}\right)^{\frac{1}{\gamma - 1}} \tag{1.11}$$

#### 131 Simulation

Generating simulated indices of abundance from the PT model requires inverting the relationship between  $\left(F^*, \frac{B^*}{B(0)}\right)$ , and  $(r, \gamma)$ . It is not generally possible to analytically invert this relationship for many three parameter production functions (Punt & Cope, 2019; J. T. Schnute & Richards, 1998). Most three parameter production functions lead to RPs that require expensive numerical methods to invert; more over the numerical inversion procedure can often be unstable. That said, for the case of PT this relationship is analytically invertible, and leads to the following relationship

$$r = \gamma F^* \qquad \qquad \gamma = \frac{W\left(\frac{B^*}{\overline{B}(0)}\log\left(\frac{B^*}{\overline{B}(0)}\right)\right)}{\log\left(\frac{B^*}{\overline{B}(0)}\right)}. \tag{1.12}$$

Above W is the Lambert product logarithm function. More details about this derivation, and the Lambert product logarithm, are given in Appendix (6).

Using Eq. (1.12) to obtain production parameters, a PT production model can be fully defined for any combination of the RPs  $F^*$  and  $\frac{B^*}{\overline{B}(0)}$ . Since K does not enter the RP calculation its value is fixed arbitrarily at 10000.

Indices of abundance are simulated from the three parameter PT production model broadly over the space of  $F^*$  and  $\frac{B^*}{\overline{B}(0)}$  via a space filling design as described in Section (0.2). A small amount of residual variation,  $\sigma = 0.01$ , is added to the simulated index, and these data are then fit with a Schaefer model, at various degrees of misspecification, so as to observe the effect of productivity model misspecification upon RP inference.

#### 142 PT Design

Letting  $\mathcal{F}$  and  $\mathcal{B}$  be regular grids, of size n=100, on  $F^* \in (0.1, 0.7)$  and  $\frac{B^*}{B_0} \in (0.2, 0.6)$ respectively, a LHS design of size 100 is collected among the cells produced by  $\mathcal{F} \times \mathcal{B}$ .

Each of the sampled LHS design locations represent a unique PT model with the sampled RP values. Since the relationship mapping RPs analytically to productivity parameters can be found for the PT model, LHS designs the PT model are computed directly in RP space and Eq. (1.12) is used to map the sampled RP design locations to PT productivity parameters.

### 2.2 Gaussian Process Metamodel

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At its core, a metamodel is simply a model of some mapping of inputs to outputs (the 151 mapping itself is typically defined by a computer model). By modeling the mapping with a 152 statistical model (that explicitly defines the relevant features of the mapping) a metamodel 153 defines a specific ontology for the mapping. By simulating examples of the mapping, the 154 inferential infrastructure of the statistical model is used to empirically learn an effective 155 emulation of the mapping within the ontology defined by the statistical model. The pre-156 dictive infrastructure of the statistical model is then useful as an approximate abstraction 157 of the system itself to better understand the system through further data collection, cheap 158 approximation of the mapping, and/or study of the mapping itself. 159

In this setting, the aim of metamodeling is to study how well RPs are inferred when typical 160 two parameter models of productivity (Logistic and BH) are misspecified for populations 161 that are actually driven by more complicated dynamics. The simulation design, X, provides 162 a sample of different population dynamics that are driven by three parameter production 163 functions broadly in RP space. By simulating index of abundance data from the three 164 parameter model, and fitting those data with the two parameter production model, we 165 observe particular instances of how well RPs are inferred at the given misspecification of the 166 two parameter model relative to the true three parameter production model. By gathering 167 all of the simulated instances of how RPs are inferred (under the two parameter model), 168 we form a set of example mappings to train a metamodel which represents the mapping 169 of true RPs (under the three parameter model) to estimates of RPs under the misspecified 170 two parameter production model. The metamodel is essentially a surrogate for inference under the misspecified two parameter production model that controls for the specific degree 172 of model misspecification. 173

A flexible GP model is assumed for the structure of the metamodel to describe the mapping of RPs under misspecified two parameter models of productivity. A GP is a stochastic process generalizing the multivariate normal distribution to an infinite dimensional analog. GP models are often specified primarily through the choice of a covariance (or correlation) function which defines the relationship between locations in the input space. Typically corre-

lation functions are specified so that points closely related in space result in correlated effects in the model. In this setting the inputs to the GP metamodel are the space of reference points 180 which define the simulated three parameter production models. 181

While index of abundance data are generated from three parameter models, at each design location of the simulation, fitting the restricted two parameter model results in a maximum likelihood estimate (MLE; and associated estimation uncertainty) of each of the productivity parameters (i.e. Schaefer: [log(r), log(K)], BH:  $[log(\alpha), log(\beta)]$ ). To simplify the specification of the metamodel, let y be a vector collecting the fitted MLEs for one of the productivity parameters, and let  $\omega$  be a vector of estimates of the estimator variances (via the inverted Fisher information) at each y. Each of the fitted productivity parameter estimates are then modeled using independent instances of the following GP metamodel.

$$\mathbf{y} = \beta_0 + \mathbf{X}\boldsymbol{\beta} + \mathbf{v} + \boldsymbol{\epsilon}$$

$$\mathbf{v} \sim N_n(\mathbf{0}, \tau^2 \mathbf{R}_{\ell})$$

$$\boldsymbol{\epsilon} \sim N_n(\mathbf{0}, \boldsymbol{\omega}' \mathbf{I})$$
(1.13)

X is the  $n \times 2$  LHS design matrix of RPs for each simulated three parameter data 182 generating model as described in Section (0.2).  $\epsilon$  models independent normally distributed 183 error, which provides an ideal mechanism for propagating uncertainty from inference in the 184 simulation step into the metamodel. By matching each  $y_i$  with an observed  $\omega_i$  variance term, 185  $\epsilon$  serves to down weight the influence of each  $y_i$  in proportion to the inferred production model 186 sampling distribution uncertainty. This has the effect of smoothing the GP model in a way similar to the nugget effect (Gramacy & Lee, 2012), although the application here models 188 this effect heterogeneously. 189

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The term,  $\boldsymbol{v}$ , contains spatially correlated GP effects. The correlation matrix,  $\boldsymbol{R}_{\ell}$  describes how RPs close together in the simulation design are more correlated than those that are far away. This spatial effect is modeled with a squared exponential correlation function,

$$R(\boldsymbol{x}, \tilde{\boldsymbol{x}}) = \exp\left(\sum_{i=1}^{2} \frac{-(x_i - \tilde{x}_i)^2}{2\ell_j^2}\right). \tag{1.14}$$

R has an anisotropic separable form which allows for differing length scales,  $\ell_1$  and  $\ell_2$ , in the different RP axes. The flexibility to model correlations separately in the different RP axes is key due to the differences in the extent of the RP domains marginally. The metamodel parameters  $\beta_0$ ,  $\beta$ ,  $\tau^2$ ,  $\ell_1$  and  $\ell_2$  are fit via MLE against the observations  $\mathbf{y}$ ,  $\mathbf{X}$ , and  $\boldsymbol{\omega}$  from simulation fits.

Fitting the metamodel allows for a full predictive description of inference under the misspecified restricted models. Predictive estimates are obtained via kriging (Cressie, 2015)

 $\hat{y}(\mathbf{x})$  is the predicted value of the modeled productivity parameter MLE under the two

$$\hat{y}(\mathbf{x}) = \beta_0 + \mathbf{x}\boldsymbol{\beta} + \mathbf{r}(\mathbf{x})' R_{\ell}^{-1} \Big( \mathbf{y} - (\beta_0 + \boldsymbol{X}\boldsymbol{\beta}) \Big)$$
(1.15)

parameter production model, when the index of abundance is generated from the three 198 parameter production model at RP location  $\mathbf{x}$ .  $\mathbf{r}(\mathbf{x})$  is a vector-valued function of correlation 199 function evaluations for the predictive location x against all observations in X (i.e.  $\mathbf{r}(\mathbf{x}) =$ 200  $\boldsymbol{R}(\mathbf{x}, \boldsymbol{x}_i) \ \forall \ \boldsymbol{x}_i \in \boldsymbol{X}).$ 201 While metamodeling occurs on the inferred productivity parameters of the restricted 202 production model, the metamodel can also be used to build estimates of major biological 203 RPs. For the BH model the relevant transformations for relating productivity parameters 204 with RPs are given in Eqs. (2.5, 2.8) with  $\gamma$  fixed to -1; for the Schaefer model  $\hat{B}^* = \frac{\hat{K}}{2}$  and  $\hat{F}^* = \frac{\hat{r}}{2}$ . Applying the metamodel predictive surfaces on the scale of RP estimates allows for 206 the quantification of estimation bias that is induced by fitting a misspecified two parameter 207

production model to indices of abundance generated under three parameter productivity.

# 209 2 .3 Catch

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It is known that contrast in the observed index and catch time series can effect inference on the productivity parameters (Hilborn & Walters, 1992). In this setting contrast refers to changes in the long term trends of index data. Figure (1.3, right) demonstrates an example of biomass that includes contrast induced by catch. It is not well understood how contrast may factor into inferential failure induced by model misspecification. Thus catch is parameterized so as to allow for a spectrum of possible contrast simulation settings.

Catch is parameterized so that F(t) can be controlled with respect to  $F^*$ . Recall that 216 catch is assumed to be proportional to biomass, so that C(t) = F(t)B(t). To control F(t)217 with respect to  $F^*$ , C(t) is specified by defining the quantity  $\frac{F(t)}{F^*}$  as the relative fishing rate. 218 B(t) is defined by the solution of the ODE, and  $F^*$  is defined by the biological parameters 219 of the model. By defining  $\frac{F(t)}{F^*}$ , catch can then be written as  $C(t) = F^*\left(\frac{F(t)}{F^*}\right)B(t)$ . 220 Intuitively  $\frac{F(t)}{F^*}$  describes the fraction of  $F^*$  that F(t) is specified to for the current B(t). 221 When  $\frac{F(t)}{F^*} = 1$ , F(t) will be held at  $F^*$ , and the solution of the ODE brings B(t) into 222 equilibrium at  $B^*$ . When  $\frac{F(t)}{F^*}$  is held constant in time biomass comes to equilibrium as an 223 exponential decay from K approaching  $B^*$ . When  $\frac{F(t)}{F^*} < 1$ , F(t) is lower than  $F^*$  and B(t) is 224 pushed toward  $\bar{B} > B^*$ . Contrarily, when  $\frac{F(t)}{F^*} > 1$ , F(t) is higher than  $F^*$  and B(t) is pushed 225 toward  $\bar{B} < B^*$ ; the precise values of  $\bar{B}$  can be calculated from the steady state biomass 226 equations provided above and depend upon the specific form of the production function. 227

For the simulations presented here, a family of fishing behaviors are considered where the fishing rate accelerates as technology and fishing techniques improve rapidly until management practices are applied, which ultimately brings fishing into equilibrium at  $F^*$ . This is parameterized as three distinct phases, over a total of 45 units of time, with each phase lasting 15 time units. The specific form is given below.

$$\frac{F(t)}{F^*} = ae^{bt}\mathbf{1}_{0 \le t < 15} + (d - ct)\mathbf{1}_{15 \le t < 30} + \mathbf{1}_{30 \le t \le 45}$$
(1.16)

The first term of Eq(1.16) is an exponential increase in fishing, the second term is a linear decline in relative fishing as initial management practices are applied, and the third term,  $\mathbf{1}_{30 \le t \le 45}$ , simply holds the fishing rate at  $F^*$  there after. These three phases are controlled by the four parameters a, b, c, and d. By enforcing that the interface of the phases meet at  $\chi_{max}$  and 1 respectively the relative fishing series is reduced to a two parameter family.

$$a = e^{\log(\chi_{max}) - 15b} \qquad b = \frac{1}{t - 15} \log\left(\frac{\chi_{min}}{\chi_{max}}\right) \tag{1.17}$$

$$c = \frac{\chi_{max} - 1}{15 - 1} \qquad d = 15c + \chi_{max} \tag{1.18}$$

By further specifying  $\chi_{max}=1.6^{\chi}$  and  $\chi_{min}=0.4^{\chi}$  the two parameters  $\chi_{max}$ , and  $\chi_{min}$ 

can be reduced to the single parameter  $\chi$ . The tuning parameter  $\chi$  then singularly controls contrast that appears in time series data.

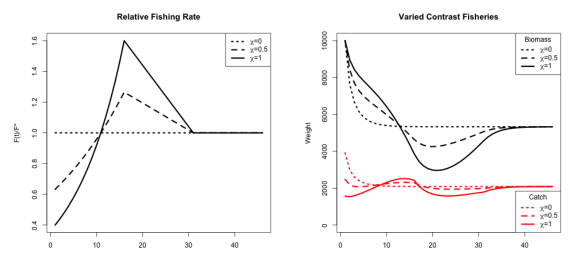


Figure 1.3: (left) Relative fishing with low, medium, and high contrast. (right) Population biomass and catch at each associated level of contrast.

When  $\chi=0$ , the relative fishing rate is a constant at 1 to create a low contrast simulation environment. As  $\chi$  increases Eq (1.16) induces more and more contrast in the observed index and catch time series until  $\chi=1$  which produces a high contrast simulation environment. Figure (1.3) demonstrates a spectrum of contrast simulation environments as well as the time series data they induce in the solution of the production model ODE.

# 2.4 Two Parameter Production Model Inference

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The simulated mapping results from fitting an intentionally misspecified two parameter production model to index of abundance data that are generated from a more complex three parameter model of productivity. Thus, let  $I_t$  be an index of abundance simulated from the three parameter PT or Schnute production models at time  $t \in \{1, 2, 3, ..., T\}$ . However the fitted model is specified to be intentionally misspecified so that the fitted model is driven by a two parameter Schaefer, or BH production model respectively.

The observation model for the fitted model is log-normal such that,

$$I_t|q, \sigma^2, \boldsymbol{\theta} \sim LN(qB_t(\boldsymbol{\theta}), \sigma^2).$$
 (1.19)

 $B_t(\boldsymbol{\theta})$  is defined by the solution of the ODEs defined by the Schaefer, or BH models. For

the Schaefer model  $\theta = [r, K]$ , and for the BH model  $\theta = [\alpha, \beta]$ . From the perspective of the fitted model, the observed  $I_t$  are assumed independent conditional on q,  $\sigma^2$ , r, K and the two parameter ODE model for biomass. Thus the log likelihood can be written as

$$\log \mathcal{L}(q, \sigma^2, \boldsymbol{\theta}; I) = -\frac{T}{2} \log(\sigma^2) - \frac{1}{2\sigma^2} \sum_{t} \log\left(\frac{I_t}{qB_t(\boldsymbol{\theta})}\right)^2.$$
 (1.20)

In this setting, q is fixed at the true value of 0.0005 to focus on the inferential effects of model misspecification on biological parameters.  $\sigma^2$  and  $\theta$  are reparameterized to the log scale and fit via MLE. Reparameterizing the parameters to the log scale improves the reliability of optimization, in addition to facilitating the use of Hessian information for estimating MLE standard errors.

Given that the biological parameters enter the likelihood via a nonlinear ODE, and further the parameters themselves are related to each other nonlinearly, the likelihood function can often be difficult to optimize. A hybrid optimization scheme is used to maximize the log likelihood to ensure that a global MLE solution is found. The R package GA (Scrucca, 2013, 2017) is used to run a genetic algorithm to explore parameter space globally. Optimization periodically jumps into the L-BFGS-B local optimizer to refine optima within a local mode. The scheme functions by searching globally, with the genetic algorithm, across many initial values for starting the local gradient-based optimizer. The genetic algorithm serves to iteratively improve hot starts for the local gradient-based optimizer. Additionally, optimization is only considered to be converged when the optimum results in an invertible Hessian at the

### 2.5 Continuous model formulation

found MLE.

An important (and often overlooked) implementation detail is the solution to the ODE which defines the progression of biomass through time. As a statistical model it is of paramount importance that this ODE not only have a solution, but also that the solution be unique. Of primary concern, uniqueness of the ODE solution is necessary for well conditioned inference.

If the form of  $\frac{dB}{dt}$  is at least Lipschitz continuous, then the Cauchy-Lipschitz-Picard theorem provides local existence and uniqueness of B(t). Recall from Eq(1.2) that  $\frac{dB}{dt}$  is

separated into a term for biomass production, P(B), and a term for removals, Z(t)B(t). For determining Lipschitz continuity of  $\frac{dB}{dt}$ , the smallest Lipschitz constant of  $\frac{dB}{dt}$  will be the sum of the constants for each of the terms P(B) and Z(t)B(t) separately. Typically any choice of P(B) will be continuously differentiable, which implies Lipschitz continuity. At a minimum Z(t) typically contains fishing mortality as a function of time Z(t) to model catch in time as Z(t) typically Z(t) may or may not contain Z(t) but typically Z(t) is modeled as stationary in time and does not pose a continuity issue, unlike some potential assumptions for Z(t).

In practice C(t) is determined by a series of observed, assumed known, catches. Catch 273 observations are typically observed on a quarterly basis, but in practice may not be complete 274 for every quarter (or year) of the modeled period. It is overwhelmingly common to discretize 275 the ODE in time via Euler's method with integration step sizes to match the observation 276 frequency of the modeled data. This is often computationally convenient when the underlying 277 species dynamics are resonably well behaved, however when the dynamics model is used as a 278 statistical model, with the goal of inferring the behavior of the underlying species dynamics, 279 the regularity of the dynamics are not guaranteed. An implicit assumption of continuity 280 of catch in time provides the necessary regularity for the statistical model. Furthermore 281 a continuous handling of the dynamics provides improved accruacy in evaluating the ODE, 282 particually when inferring productivity parameters which largely control the regularity of 283 the dynamics. 284

While there are many ways to handle catch continuity, here I assume that catches accrue linearly between observed catches. This assumption defines the catch function as a piecewise linear function of time, with the smallest Lipschitz constant for the catch term defined by the steepest time segment of the catch function. This assumption represents one of the simplest ways of handling catch, while retaining Lipschitz continuity overall. Furthermore linearly interpolated catch is adequately parsimonious for the typical handling of catches.

#### 291 Integration and Stiffness

As previously mentioned, the overwhelming majority of implementations of stock assessment models discretized the ODE using Euler's method with the integration step sized fixed so as to match the observation frequency. In this setting we explore model parameterizations that explore the full extent of biologically relevant reference points. This exercise produces some combinations of parameters that result in numerically stiff ODEs.

The concept of stiffness in ODEs is hard to precisely characterize. Wanner and Hairer (1996, p.2) describe stiffness in the following pragmatic sense, "Stiff equations are problems for which explicit methods don't work". It is hard to make this definition more mathematically precise, but this a consistent issue for models of fast growing species in the low contrast simulation. Euler's method, as often implemented, is particularly poorly suited for these stiff regions of parameter space. In these stiff regions it is necessary to integrate the ODE with an implicit integration method.

Several of the most common implicit methods were tried including the Livermore Solver for ODEs (Isode), and the Variable Coefficient ODE Solver (vode) as implemented in the deSolve package of R (Soetaert et al., 2010). The difference between implicit solvers is negligible, while explicit methods result in wildly varying solutions to the ODE in stiff regions of parameter space. Results shown here are computed using the Isode integration since it runs relatively quickly and has a relatively smaller footprint in system memory.

# 310 Results

# $_{311}$ 3.1 PT/Schaefer

## 12 An MSY-Optimal Catch History

When F(t) is held constant at  $F^*$ , as it is in the "low contrast" simulation setting, B(t) comes to equilibrium as an exponential decay from K to  $B^*$ . Understanding model misspecification bias is simplified in this setting due to the relative simplicity that this induces in B(t). However this simplicity is known to poorly inform estimates of r, and thus  $F^*$ , due to the limited range of the production function that is observed (Hilborn & Walters, 1992).

Figure (1.4) shows four of the most mis-318 specified example production function fits as 319 compared to the true data generating PT 320 production functions. The rug plots below 321 each set of curves show how the observed 322 biomasses decay exponentially from K to  $B^*$ 323 in each case. In particular, notice how ob-324 servations only exist where the PT biomass 325 is greater than  $B^*$ . Due to the leaning of 326 the true PT curves, and the symmetry of 327 the logistic parabola, the logistic curve only 328 observes information about its slope at the 329 origin from data observed on the right por-330 tion of the PT curves. The top two panels of 331 Figure (1.4) shows PT data generated such 332 that  $\frac{B^*}{\overline{B}(0)} > 0.5$ ; in these cases PT is steeper 333 to the right of  $B^*$  than it is on the left, and so 334 the the logistic curve over-estimates r, and 335 consequently also over-estimates  $F^*$ . The 336

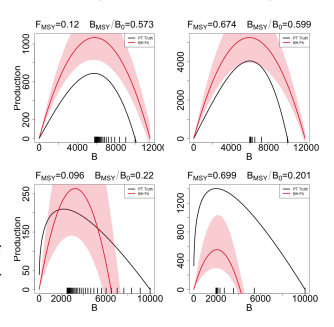


Figure 1.4: A comparison of the true PT production function (in black) and the estimated logistic curve (in red) with 95% CI shown. The examples shown represent the four corners of maximum model misspecification in the simulated RP-space. Observed biomasses are plotted in the rug plots below the curves.

bottom two panels of Figure (1.4) show PT data generated with  $\frac{B^*}{B(0)} < 0.5$  and where the vice versa phenomena occurs. PT is shallower to the right of  $B^*$  than it is on the left

and so the logistic parabola estimate tends to under estimate  $F^*$ .

### 340 Metamodeled Trends

Each point in the space of the RPs  $F^*$  and  $\frac{B^*}{B(0)}$  uniquely identifies a complete PT model 341 with different combinations of parameters values. Recall that when  $\gamma = 2$  for the PT model, 342 the PT curve becomes a parabola and is equivalent to the logistic curve of the Schaefer model. Since the logistic curve is symmetric about  $B^*$ , the Schaefer model must fix the 344 value of  $\frac{B^*}{\bar{B}(0)}$  at the constant 0.5 for any value of  $F^*$ . So the line through RP space defined 345 by  $\frac{B^*}{B(0)} = 0.5 \ \forall F^*$ , defines the subset of RP space where  $\gamma = 2$  and where the PT model 346 is equivalent to the Schaefer model. For brevity this subset of RP where  $\frac{B^*}{\bar{B}(0)} = 0.5$  will be referred to as the "Schaefer set". Thus simulated data that are generated along the Schaefer 348 set will be the only data that are not misspecified relative to the Schaefer model; as PT data 349 are simulated farther and farther away from this line at  $\frac{B^*}{\overline{B}(0)} = 0.5$  model misspecification of 350 the Schaefer model becomes worse and worse. 351

While Figure (1.4) demonstrates a real trend in simulation results, individual simulation 352 runs will at best show jittery trends due to the stochastic nature of statistical inference. The 353 GP process metamodel accounts for this stochasticity to focus analysis on the signal in the 354 simulation results. Recall that metamodeling occurs on the scale of the inferred productivity 355 parameters of the restricted production model, by transforming metamodel predictions via 356 Eq. (1.11), metamodeled predictions are obtained for Schaefer RPs. By further subtracting 357 the true data generating PT RPs from the predicted Schaefer RPs at each point in RP space 358 a pattern of inferential RP bias, induced by model misspecification of the Schaefer model, 359 can be seen. 360

Figure (1.5) shows the pattern of biases the Schaefer model creates when fit to PT data generated at each point of RP space. An equivalent way to think of Figure (1.5) is that since the Schaefer model must estimate RPs in the Schaefer set, the metamodel arrows indicate the mapping that is created by inferring RPs under a misspecified Schaefer model fit to PT data generated at each point over the pictured region.

Since  $\frac{B^*}{B_0}$  must be 0.5 under the Schaefer model, biases in the  $\frac{B^*}{B_0}$  direction must simply map vertically onto the Schaefer set. Due to this simplified RP geometry under the Schaefer

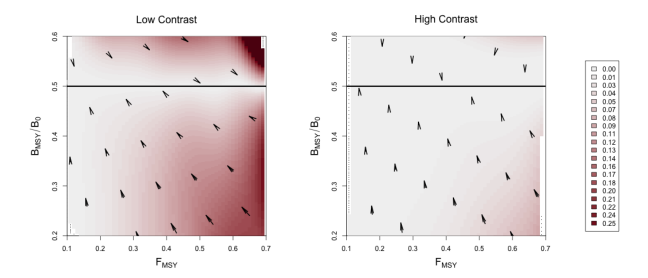


Figure 1.5: Joint bias direction for  $(F^*, \frac{B^*}{B_0})$  estimates under the misspecified Schaefer Model. The intensity of color represents the excess bias relative to the shortest possible mapping. Results in the low contrast setting are shown left, and the high contrast setting is shown right.

model, the degree of bias in  $\frac{B^*}{B_0}$  estimation is defined solely by the degree of model misspecification irrespective of  $F^*$ . Furthermore, the closest possible point along the Schaefer set that Schaefer model inference could map RPs would be the perfectly vertical mapping. This pattern only contains the strictly necessary bias present in  $\frac{B^*}{B_0}$ , and zero bias in  $F^*$ . Any deviation from this minimal bias pattern is necessarily due to added bias in  $F^*$ .

The two simulation settings shown in Figure (1.5) are identical except for the amount of contrast present in the simulated index. The left panel of Figure (1.5) shows RP biases in the low contrast setting, while the right panel shows the high contrast setting. Notice that in the low contrast setting the RP bias pattern is far from the minimum distance mapping, however when contrast is added the mapping becomes much closer to a minimal bias mapping. In the low contrast setting the observed bias is consistent with the pattern and mechanism described in Figure (1.4), where  $F^*$  is underestimated for data generated below the Schaefer line and overestimated above the Schaefer set. In the high contrast simulation the mapping is nearly minimal distance with the exception of PT data generated with simultaneously low  $\frac{B^*}{B_0}$  and high  $F^*$ .

Figure (3.1) demonstrates how bias in  $F^*$  estimation decreases as contrast is added to

#### Bias in Estimated Schaefer FMSY

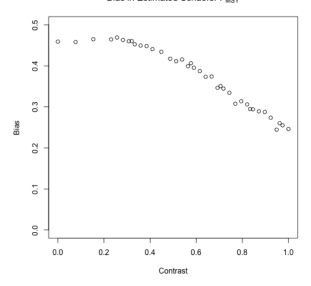


Figure 1.6: Bias in  $F^*$  under the Schaefer model when PT data are generated with increasing contrast so that  $F^*$  and  $\frac{B^*}{B_0}$  are fixed at 0.699 and 0.201 respectively.

PT data as generated in the low  $\frac{B^*}{B_0}$  and high  $F^*$  regime. By including additional contrast  $F^*$  bias is decreased, however parameterizing contrast so as to fully extinguish  $F^*$  bias may require a more complex model of fishing.

# <sup>387</sup> 4 Discussion

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Results presented here generally agree with what is known about estimating growth rate 388 parameters (Lee et al., 2012; Conn et al., 2010; Magnusson & Hilborn, 2007). These study's 389 appreciate the role of contrast for estimating growth rates, however they struggle to make 390 generally extensible conclusions since they focus only on a handful of stocks that fall short 391 of forming a random sample of the greater population of possible stock behaviors. The LHS 392 design methods presented here are designed specifically to simulate a representative sample 393 of stocks broadly across the space of possible RPs. Furthermore, the simulation design, taken together with the GP metamodel of productivity parmater estimates, allows this study to 395 control the degree of model misspecification and generalize conclusions about the behavior 396 of productivity estimation within the production model setting presented. 397

In the presence of contrast,  $F^*$  estimation can enjoy very low bias even for a wide range of poorly specified models; conversely in the absence of contrast  $F^*$  estimation can suffer very large bias even for slightly misspecified models. This pattern is particularly true for inference

under the Schaefer model where the geometry of the restricted RP set isolates estimation failure of  $F^*$  from  $\frac{B^*}{B(0)}$ . While contrast has a similar impact on  $F^*$  estimation under the BH model, the geometry of the BH RP set correlates estimation bias of  $F^*$  and  $\frac{B^*}{B(0)}$ . The GP metamodeling approach reveals a more general pattern that highly informative data sets (high contrast) produces a nearly minimal distance mapping of RPs onto the constrained RP set.

In all cases when model misspecification is removed, even with weakly informative data,
RP estimation is unbiased and well estimated. Thus contrast alone is not the only factor
leading to inferential failure. Model misspecification is a necessary but not sufficient condition for inducing RP estimation bias. The particular RP bias present depends on the RP
geometry of the fitted model and how that geometry is misspecified relative to the data. The
RP mapping is then oriented to the RP geometry of the fitted model.

While the relative fishing rate parameterized in Section (2.3) captures a usefully broad 413 spectrum of relevant fishing behaviors, it is still limiting in the amount of information that it can induce. Improved methods for quantifying contrast in fisheries data, and/or methods 415 of discovering more informative fishing behavior, could improve this analysis. In the absence 416 of a maximally informative dataset simulation methods will not fully describe how inference 417 fails, but the methods presented here tell the most complete picture yet, with explicit control 418 of the degree model misspecification, contrast, and a simulation design that allows for uniform 419 representative data generation across biologically meaningful stocks. The results presented 420 here suggest the conjecture that under a maximally informative dataset, RP inference with 421 a two parameter production function will be biased in the direction a shortest distance map 422 from the true RPs onto restricted set of RPs under the two parameter model. 423

Given the potential for model misspecification of RPs, a minimal distance mapping of RPs represents a best-case scenario where the total bias of RPs, when measured jointly, is minimized. That said, without recognizing the geometry of how two parameter models of productivity limit RP space this may lead to unintuitive implications in RP estimation. For example, due to the shape of the BH RP set a minimal distance mapping ensures that if there is bias in one of  $\frac{B^*}{B_0}$  or  $F^*$ , there will necessarily be bias in the other RP. However under the Schaefer model, since the RP set is a constant in  $\frac{B^*}{B_0}$ , bias in  $F^*$  is not adulterated in the

same way by bias in  $\frac{B^*}{B_0}$  estimation. While models with constant RPs, such as the logistic model  $\frac{B^*}{B_0} = \frac{1}{2}$  or the Fox model  $\frac{B^*}{B_0} = \frac{1}{e}$ , are extremely limited, they can be valuable tools for developing intuition precisely because they isolate RP estimation in their free RPs from the correlated RP biases present in models like the BH or Ricker model.

When one considers the implications of RP bias, overestimation of RPs carries the severe 435 implication of management recommendations potentially leading to overfishing, while un-436 derestimation of RP leads to overly conservative management. In this sense, when the true 437 model is not known, the geometry of the BH set together with the metamodeled bias trends 438 makes the BH model a naturally conservative estimator of RPs for most stocks. For most 439 non-BH populations the BH model is likely to make conservative errors in its estimates of  $F^*$  and  $\frac{B^*}{B_0}$ . The one notable exception to the conservatism of the BH model stands for data 441 generated in the Cushing-like regime of Schnute RPs. In this regime the BH model tends 442 to be fairly unbiased overall, however the bias that is present for these populations tends to 443 be overestimation in both RPs, leading to much more severe management consequences for 444 those populations. 445

The RP bias trends of the Schaefer model demonstrate much less conservatism than the BH overall. For any population with  $\frac{B^*}{B_0} < 0.5$ ,  $\frac{B^*}{B_0}$  will be overestimated. When the population comes from the regime where  $\frac{B^*}{B_0} > 0.5$ ,  $\frac{B^*}{B_0}$  will be under estimated, but  $F^*$  is likely to be overestimated depending on the degree of contrast present in the data. So while the Schaefer model is an intuitive model, it tends to lead to much less conservative RP estimation.

While it is important to recognize these limitations of two parameter models of produc-452 tivity, we should not solely accept conservativism as a rational of choosing a BH model of 453 productivity. Increasing the flexibility of the production function by moving toward three 454 parameter models would release the underlying structural limitations (Mangel et al., 2013) 455 that cause these RP biases in the first place. Punt and Cope (2019) considers a suite of pos-456 sible three parameter curves which could be used instead of current two parameter curves. 457 For all of their benefits, three parameter production functions have their own complicating 458 factors, and the structure present in the Schnute model explored here makes it an intuitive 459 bridge model for developing three parameter models going forward.

• show a schnute fit to data? (Yeakel & Mangel, 2015) Prior

- summary of  $\sigma$  over RP space comparing between models (PT, Schnute, Schnute DD) to show areas of model breakdown.
- miss-identifying signal for noise.
- It happens more as the dynamics get more complex.
- point to the full age structed models.
- show the constrained BH space over a grid of  $M, \kappa, \omega, W_{\infty}$
- Show that the constrained spaces vary only slightly as compared with the consequences of misspecifing the functional form.
- estimating these other quantities (while they can create quite different Biomass series)
  can only do so much to improve (expand) RP inference as compared with correctly
  modeling P.
- mapping distance as a function of contrast at (3.5, 0.5)
- for LHS grid locations show  $\frac{B^*}{B_0}$  and  $F^*$  biases for grids in  $M \in (0, 0.5)$  For sure in High Contrast, maybe also in Low??.

# 476 5 Appendix: Inverting $\frac{B^*}{\bar{B}(0)}$ and $\gamma$ for the PT Model

For brevity let  $\zeta = \frac{B^*}{\bar{B}(0)}$ .

$$\zeta = \left(\frac{1}{\gamma}\right)^{\frac{1}{\gamma - 1}}$$

$$\zeta = \gamma \zeta^{\gamma}$$

$$\zeta = \gamma e^{\gamma \log(\zeta)}$$

$$\zeta \log(\zeta) = \gamma \log(\zeta) e^{\gamma \log(\zeta)}$$

The Lambert product logarithm, W, is defined as the inverse function of  $z = xe^x$  such that x = W(z). Applying this definition allows for the isolation of  $\gamma$ .

$$\gamma \log(\zeta) = W(\zeta \log(\zeta))$$

$$\gamma = \frac{W(\zeta \log(\zeta))}{\log(\zeta)}$$
(1.21)

The Lambert product logarithm is a multivalued function with a branch point at  $-\frac{1}{e}$ . The principal branch,  $W_0(z)$ , is defined on  $z \in \left(-\frac{1}{e}, \infty\right)$ , and the lower branch,  $W_{-1}(z)$ , is defined on  $z \in \left(-\frac{1}{e}, 0\right)$ . Taken individually, each respective branch is analytic, but cannot be expressed in terms of elementary functions.

When  $\zeta \in \left(0, \frac{1}{e}\right)$  the solution of interest in Eq. (1.12) comes from  $W_0$ . When  $\zeta \to \frac{1}{e}$ , the Fox Model emerges as  $\gamma \to 1$ . When  $\zeta \in \left(\frac{1}{e}, 1\right)$  the solution of interest comes from  $W_{-1}$ . For the use case presented here, Eq. (1.12) is to be interpreted as,

$$\gamma = \begin{cases}
\frac{W_0(\zeta \log(\zeta))}{\log(\zeta)} & \zeta \in \left(0, \frac{1}{e}\right) \\
\frac{W_{-1}(\zeta \log(\zeta))}{\log(\zeta)} & \zeta \in \left(\frac{1}{e}, 1\right)
\end{cases}$$
(1.22)

Prager 2002, Figure(2).

https://math.stackexchange.com/questions/3004835/is-the-lambert-w-function-analyticif-not-everywhere-then-on-what-set-is-it-ana https://researchportal.bath.ac.uk/en/publications/algebraicproperties-of-the-lambert-w-function-from-a-result-of-r https://cs.uwaterloo.ca/research/tr/1993/03/W.pdf

489 Chapter 2

The Schnute Model

#### 0.1Schnute Model

The Schnute production function is a three parameter generalization of many of the most common two parameter production functions (Deriso, 1980; J. Schnute, 1985). It can be written in the following form, with parameters  $\alpha$ ,  $\beta$ , and  $\gamma$ ,

$$P_s(B; [\alpha, \beta, \gamma]) = \alpha B(1 - \beta \gamma B)^{\frac{1}{\gamma}}.$$
 (2.1)

The BH and Logistic production func-492 tions arise when  $\gamma$  is fixed to -1 or 1 respec-493 tively. The Ricker model is a limiting case 494 as  $\gamma \to 0$ . For  $\gamma < -1$  a family of strictly in-495 creasing Cushing-like curves arise, culminat-496 ing in linear production as  $\gamma \to -\infty$ . These 497 special cases form natural regimes of simi-498 larly behaving production functions as seen 499 in Figure (2.1). 500

The behavior of RP inference under the BH model is of particular interest due to the overwhelming popularity of the BH assumption in fisheries models. Since Schnute production models can represent a quantifiably wide variety of possible productivity behav-

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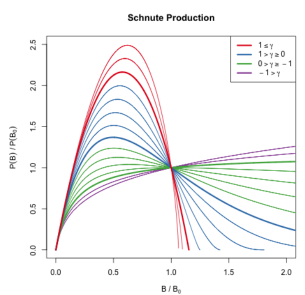


Figure 2.1: The Schnute production function plotted across a variety of parameter values. Regimes of similarly behaving curves are grouped by color.

iors, they present an ideal simulation environment for inquiry of the reliability of inference under the BH assumption. 508

Under Schnute production, biomass dynamics evolve according to the following ODE,

$$\frac{dB}{dt} = P_s(B;\theta) - (M+F)B. \tag{2.2}$$

This equation largely takes the same form as previously described, except that  $P_s$  is the Schnute production function and natural mortality, M, is modeled explicitly here. Natural 510

mortality models the instantaneous rate of mortality from all causes outside of fishing. Explicitly modeling natural mortality is not only a typical assumption of fisheries models, but is also key to the making RPs well defined over the relevant domain of  $\gamma$ .

The derivation of RPs under Eq. (2.2) follows a similar logic as under the PT model. An expression for equilibrium biomass is attained by setting  $\frac{dB}{dt} = 0$  and rearranging the resulting expression to solve for B

$$\bar{B}(F) = \frac{1}{\gamma \beta} \left( 1 - \left( \frac{M+F}{\alpha} \right)^{\gamma} \right). \tag{2.3}$$

The above expression quickly yields  $B_0$ ,  $B^*$  by evaluation at F = 0 and  $F^*$  respectively,

$$B_0 = \frac{1}{\gamma \beta} \left( 1 - \left( \frac{M}{\alpha} \right)^{\gamma} \right) \tag{2.4}$$

$$\frac{B^*}{B_0} = \frac{1 - \left(\frac{M + F^*}{\alpha}\right)^{\gamma}}{1 - \left(\frac{M}{\alpha}\right)^{\gamma}}.$$
 (2.5)

Attaining an expression for  $F^*$  requires maximization of equilibrium yield,  $\bar{Y} = F\bar{B}(F)$ , with respect to F. Analytically maximizing proceeds by differentiating  $\bar{Y}$  to produce

$$\frac{d\bar{Y}}{dF} = \bar{B}(F) + F\frac{d\bar{B}}{dF} \tag{2.6}$$

$$\frac{d\bar{B}}{dF} = -\frac{1}{\beta} \left( \frac{\left( \frac{M+F}{\alpha} \right)^{\gamma}}{F+M} \right). \tag{2.7}$$

Setting  $\frac{d\bar{Y}}{dF} = 0$ , filling in the expressions for  $\bar{B}(F)$  and  $\frac{d\bar{B}}{dF}$ , then rearranging to solve for  $F^*$  is less yielding here than it was in the case of the PT model. This procedure falls short of providing an analytical solution for  $F^*$  directly in terms of  $\theta$ , but rather shows that  $F^*$  must respect the following expression,

$$0 = \frac{1}{\gamma} - \left(\frac{1}{\gamma} + \frac{F^*}{F^* + M}\right) \left(\frac{F^* + M}{\alpha}\right)^{\gamma}. \tag{2.8}$$

The lack of an analytical solution here is understood. J. T. Schnute and Richards (1998, pg. 519) specifically points out that  $F^*$  cannot be expressed analytically in terms of productivity parameters, but rather gives a partial analytical expression for the inverse relationship.

Although parameterized slightly differently, J. T. Schnute and Richards (1998) derives expressions for  $\alpha$  and  $\beta$  as a function of RPs and  $\gamma$ . 518

Since RPs are left without a closed form expression, computing RPs from productivity 519 parameters amounts to numerically solving the system formed by collecting the expressions 520 (2.8), (2.4), and (2.5).

#### Simulation 522

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For the purposed of simulation, it is not necessary to completely know the precise relation-523 ships mapping RPs  $\mapsto \theta$  or  $\theta \mapsto$  RPs. Simulation only requires enough knowledge of these 524 mappings to gather a list of  $(\alpha, \beta, \gamma)$  tuples, for data generation under the Schnute model, 525 and the corresponding RPs in some reasonable space-filling design over RP space. 526

Similarly to J. T. Schnute and Richards (1998), expressions (2.8) and (2.4) are solved for  $\alpha$  and  $\beta$  respectively. This leads to the partial mapping  $(F^*, B_0) \mapsto (\alpha(\cdot, \gamma), \beta(\cdot, \cdot, \gamma))$  in terms of RPs and  $\gamma$ . By further working with Eq. (2.5), to identify  $\gamma$ , the following system is obtained,

$$\alpha = (M + F^*) \left( 1 + \frac{\gamma F^*}{M + F^*} \right)^{1/\gamma}$$

$$\beta = \frac{1}{\gamma B_0} \left( 1 - \left( \frac{M}{\alpha} \right)^{\gamma} \right)$$

$$\frac{B^*}{B_0} = \frac{1 - \left( \frac{M + F^*}{\alpha} \right)^{\gamma}}{1 - \left( \frac{M}{\alpha} \right)^{\gamma}}.$$
(2.9)

For a population experiencing natural mortality M, by fixing  $F^*$ ,  $B_0$ , and  $\frac{B^*}{B_0}$  the above system can fully specify  $\alpha$  and  $\beta$  for a given  $\gamma$ . Notice for a given  $\gamma$  a cascade of closed 528 form solutions for  $\alpha$  and  $\beta$  can be obtained. First  $\alpha(\gamma)$  can be computed, and then  $\beta(\alpha(\gamma), \gamma)$  can be computed. If  $\alpha(\gamma)$  is filled back into the expression for  $\frac{B^*}{B_0}$ , the system collapses into a single onerous expression for  $\frac{B^*}{B_0}(\alpha(\gamma), \gamma)$ . For brevity, define the function 531  $\zeta(\gamma) = \frac{B^*}{B_0} (\alpha(\gamma), \gamma, F^*, M)$  based on Eq. (2.5). 532 Inverting  $\zeta(\gamma)$  for  $\gamma$ , and computing the cascade of  $\alpha(\gamma)$ , and then  $\beta(\alpha(\gamma), \gamma)$ , fully defines 533 the Schnute model for a given  $(\frac{F^*}{M}, \frac{B^*}{B_0})$ . However inverting  $\zeta$  accurately is extremely difficult. Inverting  $\zeta$  analytically is not feasible, and numerical methods for inverting  $\zeta$  are unstable and can be computationally expensive. Rather than numerically invert precise values of  $\zeta(\gamma)$ ,  $\gamma$  is sampled so that the overall simulation design is space filling as described in Section (0
538 .2).

Each design location defines a complete Schnute production model with the given RP values. Indices of abundance are simulated from the Schnute model at each design location, a small amount of residual variation,  $\sigma = 0.01$ , is added to the simulated index, and the data are then fit with a misspecified BH production model. The design at large captures various degrees of model misspecification relative to the BH model, so as to observe the effect of productivity model misspecification upon RP inference.

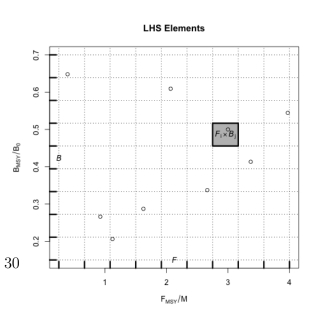
# 545 0.2 Latin Hypercube Sampling

The goal of space filling design in this setting is to extend the notion of the random sample 546 (and its desirable parameter estimation properties) across the simulated RP domain so as 547 to represent the simulated space as well as possible (Gramacy, 2020). The simple random 548 sample is the classical approach to unbiased parameter estimation, however simple random-549 ness is patchy, often sampling some regions of design space quite densely, while leaving other 550 regions of design space empty. Space filling designs aim to preserve (or enhance) parameter 551 estimation properties across the simulated domain (Devon Lin & Tang, 2015; Stein, 1987), 552 while constraining samples to be spaced in some notion of spread over the entire space. 553 Latin hypercube sampling (McKay et al., 2000, LHS) is among the most foundational of space filling designs used in computer experiments. 555

A LHS of size n, in the 2 dimensional 556 space defined by RPs, distributes samples so 557 as to spread points across a design region in 558 a broadly representative way. A LHS design 559 extends the notion of a univariate random 560 uniform sample across multiple dimensions 561 so that each margin of the design space en-562 joys a uniform distribution. 563

LHS designs achieve this notion of uni-

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formity by first partitioning each dimension

of the design space into regular grids of size

n. By intersecting the grids of each dimen-

sion, cells are produced that evenly partition

the design space. In two dimensions  $n^2$  cells

 $_{570}$  are produced, from which a total of n sam-

ples are taken. Crucially only one sample is

taken from a given element of each grid in each dimension so as to reduce clumping of the

n samples across the design space.

# 574 Schnute Design

Due to the lack of an analytical relationship mapping RPs  $\mapsto \theta$ , analogous to the PT model's

<sup>576</sup> Eq. (1.12), producing a LHS design over Schnute RPs requires a more tactful approach. The

structured relationship between the RPs and productivity parameters, described in Section

578 (0.1), allows an approximate LHS to be obtained by a careful navigation of the system of

equations seen in Eq. (2.9).

Under the Schnute model, let  $\mathcal{F}$  and  $\mathcal{B}$ 

represent regular grids on  $\frac{F^*}{M} \in (0.25, 4)$  and

 $\frac{B^*}{B_0} \in (0.15, 0.7)$  respectively which can serve

as the scaffolding for computing an approx-

imate LHS.

Since it is not practical to invert  $\zeta(\gamma)$ , a uniform sample in  $\frac{B^*}{B_0}$  can be obtained by modeling  $\gamma$  as a random variable, with realization  $\gamma^*$ , and thinking of  $\zeta(\gamma)$  as its cumulative distribution function (CDF). The aim is to model  $\gamma$  as an easily sampled random

Given  $B_0$ , M, and  $F^*$ :

- 1) Draw  $\gamma^* \sim \gamma | F^*, M$ .
- 2) Compute  $\frac{B^*}{B_0} = \zeta(\gamma^*)$
- 3) Compute  $\alpha^* = \alpha(\gamma^*, F^*, M)$
- 4) Compute  $\beta^* = \beta(\alpha^*, \gamma^*, M, B_0)$

Figure 2.3: An outline of the sampling procedure for  $\gamma$  given  $B_0$ , M, and  $F^*$ .

variable with a CDF that closely approximates  $\zeta$ , so that  $\zeta(\gamma^*) \sim U(\zeta_{min}, 1)$  as closely as possible. There may be many good models for the distribution of  $\gamma$ , but in this setting the

following distribution is very effective,

$$\gamma \sim \zeta_{min}\delta(\gamma_{min}) + t(\mu, \sigma, \nu)\mathbf{1}_{\gamma > \gamma_{min}}.$$
 (2.10)

Above, t is the density of the three pa-585 rameter location-scale family Student's t dis-586 tribution with location  $\mu$ , scale  $\sigma$ , and de-587 grees of freedom  $\nu$ .  $\mathbf{1}_{\gamma > \gamma_{min}}$  is an indica-588 tor function that serves to truncate the Stu-589 dent's t distribution at the lower bound  $\gamma_{min}$ . 590  $\delta(\gamma_{min})$  is the Dirac delta function evaluated 591 at  $\gamma_{min}$ , which is scaled by the known value 592  $\zeta_{min}$ ; this places probability mass  $\zeta_{min}$  at 593 the point  $\gamma_{min}$ . Since sampling from a Student's t distribution is readily doable, sam-595 pling from a truncated Student's t mixture 596 only requires slight modification. 597

Let T be the CDF of the modeled distribution of  $\gamma$ . Since the point  $(\gamma_{min}, \zeta_{min})$  is



Figure 2.4:  $\zeta(\gamma)$  Plotted for  $F^* = 0.1$  and M = 0.2. The point  $(\gamma_{min}, \zeta_{min})$  shows the lowest biologically meaningful value of  $\gamma$ ; below which productivity is negative.

known from the dynamics of the Schnute model at a given RP, full specification of Eq. (2.10) only requires determining the values for  $\mu$ ,  $\sigma$ , and  $\nu$  which make T best approximate  $\zeta(\gamma)$ . Thus, the values of  $\mu$ ,  $\sigma$ , and  $\nu$  are chosen by minimizing the  $L^2$  distance between  $T(\gamma)$  and  $\zeta(\gamma)$ .

$$[\hat{\mu}, \hat{\sigma}, \hat{\nu}] = \underset{[\mu, \sigma, \nu]}{\operatorname{arg \, min}} \int_{\Gamma} \left( T(\gamma; \mu, \sigma, \nu) - \zeta(\gamma) \right)^2 d\gamma \tag{2.11}$$

```
The distribution T(\gamma|\hat{\mu}, \hat{\sigma}, \hat{\nu}) is fit for use
                                                                     Algorithm 1 LHS of size n on rectangle R.
598
                                                                       1: procedure LHS_n(R)
     in generating \gamma^* random variates at a spe-
599
                                                                       2:
                                                                               Define n-grids \mathcal{F}, \mathcal{B} \in R
     cific F^* and M. This approximation releases
600
                                                                               for each grid element i do
     the need to invert \zeta w.r.t \gamma by using sam-
                                                                       3:
601
                                                                                    Draw \frac{F^*}{M} \sim Unif(\mathcal{F}_i)
     ples of \gamma^* values to generate approximatly
                                                                       4:
602
                                                                                    Compute [\hat{\mu}, \hat{\sigma}, \hat{\nu}] given F^* \& M
                                                                       5:
     uniform samples of \zeta(\gamma^*). By sampling ap-
603
                                                                                    while \mathcal{B}_j not sampled do
                                                                       6:
     proximatly uniform \zeta(\gamma^*) random variates in
604
                                                                                         Draw \gamma^* \sim T(\gamma | \hat{\mu}, \hat{\sigma}, \hat{\nu})
                                                                       7:
     this way, and making use of the structure in
605
                                                                                         Compute \zeta^* = \zeta(\gamma^*)
                                                                       8:
     Eq. (2.9), an approximate LHS sample can
606
                                                                                         Compute j such that \zeta^* \in \mathcal{B}_i
                                                                       9:
     be collected via Algorithm (1).
607
          \frac{F^*}{M} is drawn uniformly from \mathcal{F}_i. Con-
                                                                                    end while
                                                                     10:
608
                                                                                    Compute \alpha^* = \alpha(\gamma^*, F^*, M)
     ditioning on the sample of F^*, and M,
                                                                      11:
609
                                                                                    Compute \beta^* = \beta(\alpha^*, \gamma^*, M, B_0)
     T(\gamma|\hat{\mu},\hat{\sigma},\hat{\nu}) is fit and \gamma^* is sampled. \zeta^* is
                                                                     12:
610
                                                                                    Save (\frac{F^*}{M}, \zeta^*) \Leftrightarrow (\alpha^*, \beta^*, \gamma^*) in \mathcal{F}_i \times \mathcal{B}_j
                                                                      13:
     then computed and placed into the appropri-
611
                                                                               end for
                                                                      14:
     ate grid element \mathcal{B}_{j}. Given \gamma^{*}, the cascade
612
                                                                     15: end procedure
     \alpha(\gamma^*), and \beta(\alpha(\gamma^*), \gamma^*), can be computed.
613
     The algorithm continues until all of the de-
614
```

### 16 Design Refinement

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Since the behavior of RP inference, under misspecified models, will vary in yet-unknown ways, the exact sampling design density may be hard to know a priori. Several factors, including the particular level of observation uncertainty, high variance (i.e. hard to resolve) features of the response surface, or simply "gappy" instantiations of the initial LHS design may necessitate adaptive design refinement, to accurately describe RP biases. Given the temperamental relationship between RPs and productivity parameters in the Schnute model, a recursive refinement algorithm that makes use of the previously described LHS routine, is developed.

sign elements,  $(\frac{F^*}{M}, \zeta^*) \Leftrightarrow (\alpha^*, \beta^*, \gamma^*)$ , have been computed for all  $i \in [1, ..., n]$ .

While LHS ensures uniformity in the design margins, and a certain degree of spread, it

is widely recognized that particular LHS instantiations may leave substantive gaps in the simulation design. To correct this, LHS is often paired with design elements of maximin 627 design (Morris & Mitchell, 1995; Devon Lin & Tang, 2015). Maximin designs sample the 628 design space by maximizing the minimum distance between sampled points. This has the 629 advantage of definitionally filling holes in the design, however because no points are ever drawn outside of the design domain, samples tend to clump around edges (particularly 631 corners) of the design domain. Since LHS ensures uniformity in the margins and maximin 632 designs enjoys a certain sense of optimality in how they define and fill gaps (Johnson et al., 633 1990), the methods are quite complimentary when combined. 634

Making use of this complimentary relationship, holes in the existing LHS design of RPs are identified based on maximin design principles. New design points are collected based on areas of the RP design space which maximizes the minimum distance between all pairs of points in the current design, based on the following distance function

$$d(\boldsymbol{x}, \boldsymbol{x'}) = \sqrt{(\boldsymbol{x} - \boldsymbol{x'})^T \boldsymbol{D}^{-1}(\boldsymbol{x} - \boldsymbol{x'})}$$

$$\boldsymbol{D} = \operatorname{diag} \left[ \left( \max(\mathcal{F}) - \min(\mathcal{F}) \right)^2, \left( \max(\mathcal{B}) - \min(\mathcal{B}) \right)^2 \right].$$
(2.12)

Above, d is a scaled distance function that defines the distance between points in the differing scales of  $\frac{B^*}{B_0}$  and  $\frac{F^*}{M}$ .  $\mathbf{D}$  is a diagonal matrix that measures the squared size of the domain in each axis of so as to normalize distances to a common scale.

If  $X_n$  is the initial design, computed on  $R_{full}$ , let  $x_a$  be the augmenting point which maximizes the minimum distance between all of the existing design points,

$$x_a = \underset{x'}{\operatorname{argmax}} \min\{d(x_i, x') : i = 1, ..., n\}.$$
 (2.13)

The point  $x_a$  is used as an anchor for augmenting  $X_n$ . An additional  $LHS_{n'}$  (via Algorithm (1)) is collected, adding n' design points, centered around  $x_a$ , to the overall design. The augmenting region,  $R_{(x_a,d_a)}$ , for collecting  $LHS_{n'}$  is defined based on the square centered at  $x_a$  with side length  $2d_a$ , where  $d_a = \min\{d(x_i, x_a) : i = 1, ..., n\}$ , in the space defined by the metric d.

Due to the tendency of maximin sampling to cluster augmenting points on the edges of the design space,  $R_{(x_a,d_a)}$  is truncated by the outer most limits of  $R_{full}$  so as to focus design augmentation within the specified domain of the simulation. Furthermore, since the design space has a nonlinear constraint at low values of  $\frac{B^*}{B_0}$ , the calculation of  $x_a$  is further truncated based on a convex hull defined by the existing samples in the overall design.

Design refinement then proceeds as follows. An initial design is computed,  $X_n = LHS_n(R_{full})$ , based on an overall simulated region of RPs  $R_{full}$ . The maximin augmenting point,  $x_a$ , is computed at a maximin distance of  $d_a$  from the existing samples. An augmenting design  $X_{n'} = LHS_{n'}(R_{(x_a,d_a)})$  is collected and added to  $X_n$ . Design refinement carries on recursively collecting augmenting designs in this way until the maximin distance falls below the desired level.

#### 654 0.3 Gaussian Process Metamodel

At its core, a metamodel is simply a model of some mapping of inputs to outputs (the 655 mapping itself is typically defined by a computer model). By modeling the mapping with a 656 statistical model (that explicitly defines the relevant features of the mapping) a metamodel 657 defines a specific ontology for the mapping. By simulating examples of the mapping, the 658 inferential infrastructure of the statistical model is used to empirically learn an effective 659 emulation of the mapping within the ontology defined by the statistical model. The pre-660 dictive infrastructure of the statistical model is then useful as an approximate abstraction 661 of the system itself to better understand the system through further data collection, cheap 662 approximation of the mapping, and/or study of the mapping itself. 663

In this setting, the aim of metamodeling is to study how well RPs are inferred when typical two parameter models of productivity (Logistic and BH) are misspecified for populations that are actually driven by more complicated dynamics. The simulation design, X, provides a sample of different population dynamics that are driven by three parameter production functions broadly in RP space. By simulating index of abundance data from the three parameter model, and fitting those data with the two parameter production model, we observe particular instances of how well RPs are inferred at the given misspecification of the two parameter model relative to the true three parameter production model. By gathering

all of the simulated instances of how RPs are inferred (under the two parameter model),
we form a set of example mappings to train a metamodel which represents the mapping
of true RPs (under the three parameter model) to estimates of RPs under the misspecified
two parameter production model. The metamodel is essentially a surrogate for inference
under the misspecified two parameter production model that controls for the specific degree
of model misspecification.

A flexible GP model is assumed for the structure of the metamodel to describe the map-678 ping of RPs under misspecified two parameter models of productivity. A GP is a stochastic 679 process generalizing the multivariate normal distribution to an infinite dimensional analog. 680 GP models are often specified primarily through the choice of a covariance (or correlation) 681 function which defines the relationship between locations in the input space. Typically corre-682 lation functions are specified so that points closely related in space result in correlated effects 683 in the model. In this setting the inputs to the GP metamodel are the space of reference points 684 which define the simulated three parameter production models. 685

While index of abundance data are generated from three parameter models, at each design location of the simulation, fitting the restricted two parameter model results in a maximum likelihood estimate (MLE; and associated estimation uncertainty) of each of the productivity parameters (i.e. Schaefer:[log(r), log(K)], BH:[ $log(\alpha)$ ,  $log(\beta)$ ]). To simplify the specification of the metamodel, let  $\mathbf{y}$  be a vector collecting the fitted MLEs for one of the productivity parameters, and let  $\boldsymbol{\omega}$  be a vector of estimates of the estimator variances (via the inverted Fisher information) at each  $\mathbf{y}$ . Each of the fitted productivity parameter estimates are then modeled using independent instances of the following GP metamodel.

$$\mathbf{y} = \beta_0 + \mathbf{X}\boldsymbol{\beta} + \mathbf{v} + \boldsymbol{\epsilon}$$

$$\mathbf{v} \sim N_n(\mathbf{0}, \tau^2 \mathbf{R}_{\ell})$$

$$\boldsymbol{\epsilon} \sim N_n(\mathbf{0}, \boldsymbol{\omega}' \mathbf{I})$$
(2.14)

X is the  $n \ge 2$  LHS design matrix of RPs for each simulated three parameter data generating model as described in Section (0.2).  $\epsilon$  models independent normally distributed error, which provides an ideal mechanism for propagating uncertainty from inference in the

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simulation step into the metamodel. By matching each  $y_i$  with an observed  $\omega_i$  variance term,  $\epsilon$  serves to down weight the influence of each  $y_i$  in proportion to the inferred production model 690 sampling distribution uncertainty. This has the effect of smoothing the GP model in a way similar to the nugget effect (Gramacy & Lee, 2012), although the application here models 692 this effect heterogeneously.

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The term,  $oldsymbol{v}$ , contains spatially correlated GP effects. The correlation matrix,  $oldsymbol{R}_{\ell}$  describes how RPs close together in the simulation design are more correlated than those that are far away. This spatial effect is modeled with a squared exponential correlation function,

$$R(\boldsymbol{x}, \tilde{\boldsymbol{x}}) = \exp\left(\sum_{i=1}^{2} \frac{-(x_i - \tilde{x}_i)^2}{2\ell_j^2}\right). \tag{2.15}$$

R has an anisotropic separable form which allows for differing length scales,  $\ell_1$  and  $\ell_2$ , 694 in the different RP axes. The flexibility to model correlations separately in the different 695 RP axes is key due to the differences in the extent of the RP domains marginally. The 696 metamodel parameters  $\beta_0$ ,  $\boldsymbol{\beta}$ ,  $\tau^2$ ,  $\ell_1$  and  $\ell_2$  are fit via MLE against the observations  $\mathbf{y}$ ,  $\boldsymbol{X}$ , 697 and  $\omega$  from simulation fits. 698

Fitting the metamodel allows for a full predictive description of inference under the misspecified restricted models. Predictive estimates are obtained via kriging (Cressie, 2015)

 $\hat{y}(\mathbf{x})$  is the predicted value of the modeled productivity parameter MLE under the two

$$\hat{y}(\mathbf{x}) = \beta_0 + \mathbf{x}\boldsymbol{\beta} + \mathbf{r}(\mathbf{x})' \mathbf{R}_{\ell}^{-1} \Big( \mathbf{y} - (\beta_0 + \mathbf{X}\boldsymbol{\beta}) \Big)$$
(2.16)

parameter production model, when the index of abundance is generated from the three 702 parameter production model at RP location  $\mathbf{x}$ .  $\mathbf{r}(\mathbf{x})$  is a vector-valued function of correlation 703 function evaluations for the predictive location  $\mathbf{x}$  against all observations in  $\mathbf{X}$  (i.e.  $\mathbf{r}(\mathbf{x}) =$ 704  $\boldsymbol{R}(\mathbf{x}, \boldsymbol{x}_i) \ \forall \ \boldsymbol{x}_i \in \boldsymbol{X}).$ 705 While metamodeling occurs on the inferred productivity parameters of the restricted 706 production model, the metamodel can also be used to build estimates of major biological 707 RPs. For the BH model the relevant transformations for relating productivity parameters 708 with RPs are given in Eqs. (2.5, 2.8) with  $\gamma$  fixed to -1; for the Schaefer model  $\hat{B}^* = \frac{\hat{K}}{2}$  and 709

 $\hat{F}^* = \hat{\frac{\hat{r}}{2}}$ . Applying the metamodel predictive surfaces on the scale of RP estimates allows for the quantification of estimation bias that is induced by fitting a misspecified two parameter production model to indices of abundance generated under three parameter productivity.

# 713 1 Results

## 4 1.1 Schnute/BH

#### 715 Design

Algorithm (1) enforces uniform marginals in  $\frac{F^*}{M}$ 716 directly, as well as the adherence of the overall 717 design to latin squares. Figure (2.5) shows a uni-718 form Q-Q plot for sampled  $\zeta$ , using Algorithm (1), against theoretical uniform quantiles. 720 evidence by the excellent coherence to the the-721 oretical uniform quantiles, the approximation in 722 Section (0.2) for sampling  $\gamma$  (and therefore  $\zeta(\gamma)$ ), 723 is very effective. Furthermore since numerical in-724 version of  $\zeta(\gamma)$  is costly and unreliable, the rel-725 ative speed and accuracy that this approximate 726

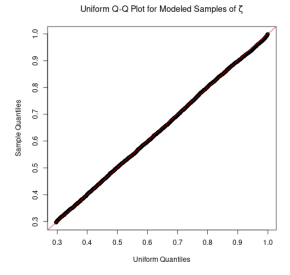


Figure 2.5: Uniform Q-Q plot for  $\zeta$  plotted for  $F^* = 0.1$  and M = 0.2.

LHS sampling method provides is pivotal for the rest of the work presented here.

Similarly to the PT model, the three pa-728 rameter Schnute model is uniquely identified 729 by each point in the space of  $\frac{F^*}{M}$  and  $\frac{B^*}{B_0}$  RPs. 730 As seen in Figure (2.6), Schnute production 731 has different behaviors in different ranges of 732 RPs space, which are entirely defined by the 733 value of  $\gamma$  (shown in Figure (2.1)). When  $\gamma \geq 1$  the Schnute model produces a family 735 of Logistic-like curves that are increasingly 736 right leaning as  $\gamma$  increases. For  $1 > \gamma \ge 0$ , 737 Schnute production takes a family of left 738 leaning Ricker-like curves that all, at least, 739 approach the x-axis. For  $0 > \gamma > -1$  there 740 are a family of BH-like curves that do not 741

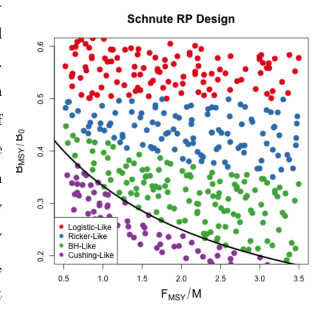


Figure 2.6: A Schnute RP design. Colors indicate different regimes of Schnute production.

The black curve shows the BH set.

742 approach the x-axis but still have decreas-

ing productivity for large biomass stocks. When  $\gamma$  is exactly -1 Schnute reduces to BH production which has asymptoting production for large biomass. Finally when  $-1 > \gamma$  Schnute produces a family of increasing Cushing-like curves that do not asymptote, and produces linear production as  $\gamma \to -\infty$ .

Modeling index data that are simulated broadly over the theoretical space of RPs with

misspecified BH production greatly limits the range of possible RPs that can be inferred. Under BH production the full theoretical space of RPs are limited to the curve  $\frac{B^*}{B_0} = \frac{1}{F^*/M+2}$ .

Define the "BH set" as the set of RPs defined by this limited space, i.e. the curve  $\left\{\left(\frac{F^*}{M}, \frac{B^*}{B_0}\right) \middle| \frac{B^*}{B_0} = \frac{1}{F^*/M+2}\right\}$ . as seen in the black curve in Figure (2.6). The farther away from this set that Schnute data are simulated, the worse the BH model is misspecified for

#### 754 Metamodeled Trends

those data.

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Unlike the Schaefer model, the BH set is not a constant in  $\frac{B^*}{B_0}$ . Under the BH model, bias in  $\frac{B^*}{B_0}$  is no longer entirely defined by the degree of model misspecification, but rather the structure of BH RPs allows bias in both  $\frac{B^*}{B_0}$  and  $\frac{F^*}{M}$  to interact as a function of contrast in the data.

High Contrast Figure (2.7) shows metamodeled RP bias surfaces for inference under the BH model in the high contrast setting. The (left) and (bottom) panels focus only on the  $\frac{B^*}{B(0)}$  and  $\frac{F^*}{M}$  components of bias respectively. In these panels bias is shown as relative bias, similar to a percent error calculation. Where RP represents the true value of the three parameter RP, and  $\widehat{RP}$  refers to the metamodel estimate.

Figure (2.7, top-right) combines the components of bias to show the overall mapping of RPs under BH inference in the high contrast simulation setting. Unlike high contrast RP inference under the Schaefer model, the BH model does shows bias in both RPs here. Despite the bias in  $\frac{B^*}{B(0)}$  and  $\frac{F^*}{M}$  these results are similar to that of the Schaefer model in that the overall mapping of RPs is very nearly a minimal distance mapping onto the constrained set of RPs. The primary difference between Schaefer model and BH RP inference is the geometry of their limited RP spaces. Unlike the Schaefer model the BH set encourages bias

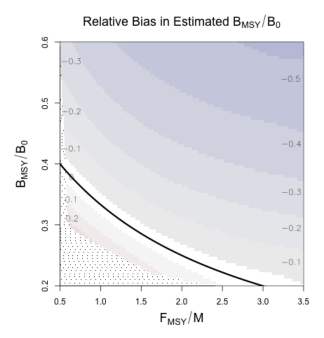
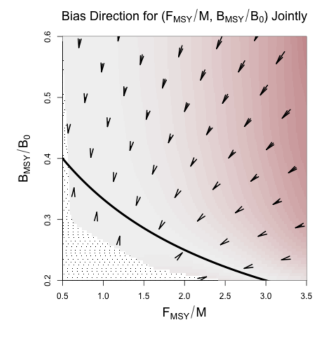
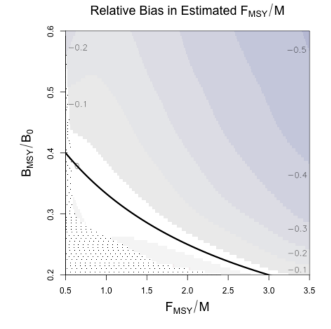


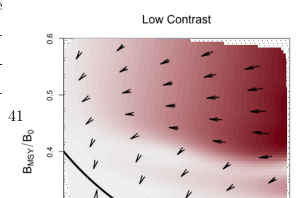
Figure 2.7: Heatplots showing the bias in RP estimation induced by model misspecification of the BH model in the high contrast simulation setting. In all cases the restricted RP-space of the BH set is shown as the black curve. (left) Relative bias in  $\frac{B^*}{B(0)}$ . (top-right) Bias in RP-space shown directionally. Arrows point from the location where data is generated, toward the location in the BH set where MLE projects estimated RPs. The intensity of color represents the excess bias relative to the shortest possible mapping. (bottom) Relative bias in  $F^*$ .





in both RPs for misspecified models even in very well informed setting.

Low Contrast Figure (2.8) shows the mapping of RPs in the low contrast simulation setting. Figures (2.8) and (2.7, top-right) share a common scale for the inten-



sity of color to facilitate comparison. In Figure (2.8) notice that the mildly misspecified 777 area around the BH set produces mappings 778 onto the BH set which resemble the minimal 779 distance mapping seen in the high contrast setting. The primary difference in this low 781 contrast setting, is the break point around 782  $\frac{B^*}{B(0)} = 0.4$  above which  $\frac{F^*}{M}$  is sharply under-783 estimated. 784

The region of RPs where the BH model manages to recover the minimal distance mapping may be considered a "safe regime"

of data types that are reasonably well modeled by a BH model. By comparison of Figure (2.8), with Figure (2.6), this safe regime of the BH model occurs for data generated for Cushing-like or BH-like production. While bias of the RPs can still become concerningly large, this region can be considered safe in the sense that even for low contrast data RP estimation under the the BH model recovers the minimal distance mapping.

Outside of this safe regime, RP estima-793 tion breaks from the minimal distance map-794 ping at the interface between BH-Like and 795 Ricker-Like regimes of the Schnute model 796 (again see Figure (2.6)). The Ricker model 797 lies along this regime interface, and repre-798 sents the first model to approach the x-axis for large biomasses as  $\gamma$  increases. 800 markedly unBH-like productivity in the low 801 information simulation setting breaks MLE 802 inference from the minimal distance map-803 ping and instead maps RPs to extremely low 804 values of  $F^*$ ; consequently  $\frac{B^*}{\overline{B}(0)}$  is estimated

#### **Estimated Yield Curves For Poorly Specified BH**

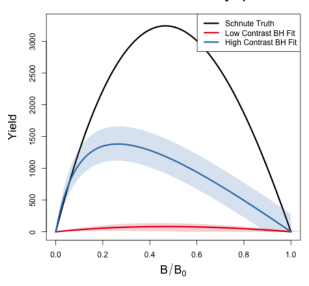


Figure 2.9: Yield curves for data generated with  $\frac{F^*}{M} = 3.48$  and  $\frac{B^*}{\overline{B}(0)} = 0.48$ .

near the limiting value under the BH (i.e.

 $\lim_{F^*\to 0} \frac{1}{F^*/M+2} = 0.5$ ). Similarly the set of Ricker RPs (as well as the Schaeffer set) include this trivial limiting point in common  $(\frac{F^*}{M} = 0, \frac{B^*}{B(0)} = 0.5)$ .

Interestingly, in the high contrast setting this trivial mapping for highly misspecified BH 809 models is not present. This suggests that, under a misspecified BH model, the presence of 810 adequate information in the data to produce reasonable estimates of  $\frac{F^*}{M}$ , drives  $\frac{B^*}{B(0)}$  below 0.5 811 in accordance with  $\frac{B^*}{\bar{B}(0)} = \frac{1}{F^*/M+2}$ , even when the true  $\frac{B^*}{\bar{B}(0)} > 0.5$ . This phenomena balances 812 RP estimation within the constrained BH set as mediated by the information content of the 813 data and the degree of model misspecification. When the information content in the data 814 is too small to drive a compromised RP estimate, inference completely disregards accurate 815 estimation of  $F^*$  in order to better estimate  $\frac{B^*}{\overline{B}(0)}$  by exploiting the common limiting behavior 816 of the BH set and that of Ricker-like and Logistic-like models. 817

# <sup>818</sup> 2 Discussion

Results presented here generally agree with what is known about estimating growth rate 819 parameters (Lee et al., 2012; Conn et al., 2010; Magnusson & Hilborn, 2007). These study's 820 appreciate the role of contrast for estimating growth rates, however they struggle to make 821 generally extensible conclusions since they focus only on a handful of stocks that fall short 822 of forming a random sample of the greater population of possible stock behaviors. The LHS 823 design methods presented here are designed specifically to simulate a representative sample 824 of stocks broadly across the space of possible RPs. Furthermore, the simulation design, taken 825 together with the GP metamodel of productivity parmater estimates, allows this study to 826 control the degree of model misspecification and generalize conclusions about the behavior 827 of productivity estimation within the production model setting presented. 828

In the presence of contrast,  $F^*$  estimation can enjoy very low bias even for a wide range of poorly specified models; conversely in the absence of contrast  $F^*$  estimation can suffer very large bias even for slightly misspecified models. This pattern is particularly true for inference under the Schaefer model where the geometry of the restricted RP set isolates estimation failure of  $F^*$  from  $\frac{B^*}{B(0)}$ . While contrast has a similar impact on  $F^*$  estimation under the BH model, the geometry of the BH RP set correlates estimation bias of  $F^*$  and  $\frac{B^*}{B(0)}$ . The

GP metamodeling approach reveals a more general pattern that highly informative data sets (high contrast) produces a nearly minimal distance mapping of RPs onto the constrained RP set.

In all cases when model misspecification is removed, even with weakly informative data,
RP estimation is unbiased and well estimated. Thus contrast alone is not the only factor
leading to inferential failure. Model misspecification is a necessary but not sufficient condition for inducing RP estimation bias. The particular RP bias present depends on the RP
geometry of the fitted model and how that geometry is misspecified relative to the data. The
RP mapping is then oriented to the RP geometry of the fitted model.

While the relative fishing rate parameterized in Section (2.3) captures a usefully broad 844 spectrum of relevant fishing behaviors, it is still limiting in the amount of information that 845 it can induce. Improved methods for quantifying contrast in fisheries data, and/or methods 846 of discovering more informative fishing behavior, could improve this analysis. In the absence 847 of a maximally informative dataset simulation methods will not fully describe how inference 848 fails, but the methods presented here tell the most complete picture yet, with explicit control 849 of the degree model misspecification, contrast, and a simulation design that allows for uniform 850 representative data generation across biologically meaningful stocks. The results presented 851 here suggest the conjecture that under a maximally informative dataset, RP inference with 852 a two parameter production function will be biased in the direction a shortest distance map 853 from the true RPs onto restricted set of RPs under the two parameter model. 854

Given the potential for model misspecification of RPs, a minimal distance mapping of 855 RPs represents a best-case scenario where the total bias of RPs, when measured jointly, is 856 minimized. That said, without recognizing the geometry of how two parameter models of 857 productivity limit RP space this may lead to unintuitive implications in RP estimation. For 858 example, due to the shape of the BH RP set a minimal distance mapping ensures that if 859 there is bias in one of  $\frac{B^*}{B_0}$  or  $F^*$ , there will necessarily be bias in the other RP. However under 860 the Schaefer model, since the RP set is a constant in  $\frac{B^*}{B_0}$ , bias in  $F^*$  is not adulterated in the 861 same way by bias in  $\frac{B^*}{B_0}$  estimation. While models with constant RPs, such as the logistic 862 model  $\frac{B^*}{B_0} = \frac{1}{2}$  or the Fox model  $\frac{B^*}{B_0} = \frac{1}{e}$ , are extremely limited, they can be valuable tools for developing intuition precisely because they isolate RP estimation in their free RPs from

the correlated RP biases present in models like the BH or Ricker model.

When one considers the implications of RP bias, overestimation of RPs carries the severe 866 implication of management recommendations potentially leading to overfishing, while un-867 derestimation of RP leads to overly conservative management. In this sense, when the true 868 model is not known, the geometry of the BH set together with the metamodeled bias trends makes the BH model a naturally conservative estimator of RPs for most stocks. For most 870 non-BH populations the BH model is likely to make conservative errors in its estimates of 871  $F^*$  and  $\frac{B^*}{B_0}$ . The one notable exception to the conservatism of the BH model stands for data 872 generated in the Cushing-like regime of Schnute RPs. In this regime the BH model tends 873 to be fairly unbiased overall, however the bias that is present for these populations tends to 874 be overestimation in both RPs, leading to much more severe management consequences for 875 those populations. 876

The RP bias trends of the Schaefer model demonstrate much less conservatism than the BH overall. For any population with  $\frac{B^*}{B_0} < 0.5$ ,  $\frac{B^*}{B_0}$  will be overestimated. When the population comes from the regime where  $\frac{B^*}{B_0} > 0.5$ ,  $\frac{B^*}{B_0}$  will be under estimated, but  $F^*$  is likely to be overestimated depending on the degree of contrast present in the data. So while the Schaefer model is an intuitive model, it tends to lead to much less conservative RP estimation.

While it is important to recognize these limitations of two parameter models of produc-883 tivity, we should not solely accept conservativism as a rational of choosing a BH model of 884 productivity. Increasing the flexibility of the production function by moving toward three 885 parameter models would release the underlying structural limitations (Mangel et al., 2013) 886 that cause these RP biases in the first place. Punt and Cope (2019) considers a suite of pos-887 sible three parameter curves which could be used instead of current two parameter curves. 888 For all of their benefits, three parameter production functions have their own complicating 889 factors, and the structure present in the Schnute model explored here makes it an intuitive 890 bridge model for developing three parameter models going forward. 891

• show a schnute fit to data? (Yeakel & Mangel, 2015) Prior

892

- summary of  $\sigma$  over RP space comparing between models (PT, Schnute, Schnute DD) to show areas of model breakdown.
- miss-identifying signal for noise.
- It happens more as the dynamics get more complex.
- point to the full age structed models.
- show the constrained BH space over a grid of  $M, \kappa, \omega, W_{\infty}$
- Show that the constrained spaces vary only slightly as compared with the consequences of misspecifing the functional form.
- estimating these other quantities (while they can create quite different Biomass series)
  can only do so much to improve (expand) RP inference as compared with correctly
  modeling P.
- mapping distance as a function of contrast at (3.5, 0.5)
- for LHS grid locations show  $\frac{B^*}{B_0}$  and  $F^*$  biases for grids in  $M \in (0, 0.5)$  For sure in High Contrast, maybe also in Low??.

907 Chapter 3

A Delay Differential Model

#### • Introduction

- piggy back intro off of simpleModel
- 911 problem statement and motivation
- introduce reference point and management decision making
- new dynamics of cohorting.

#### • Methods

- state and decribe model
- Reference Point Derivation
- layout data generation/space filling problem
- how far to get the math for inputting into CAS
- method of CAS.
- describe and plot  $\zeta$ .
- constrained BH space (method for visualizing)
- appendix for RP CAS calculation
- Results
- summary of  $\sigma$  over RP space comparing between models (PT, Schnute, Schnute DD) to show areas of model breakdown.
- miss-identifying signal for noise.
- It happens more as the dynamics get more complex.
- point to the full age structed models.
- Show that the constrained spaces vary only slightly as compared with the consequences of misspecifing the functional form.
- ?Discussion?

- summary of  $\sigma$  over RP space comparing between models (PT, Schnute, Schnute DD) to show areas of model breakdown.
- miss-identifying signal for noise.
- It happens more as the dynamics get more complex.
- point to the full age structed models.
- show the constrained BH space over a grid of  $M, \kappa, \omega, W_{\infty}$
- Show that the constrained spaces vary only slightly as compared with the consequences of misspecifing the functional form.
- estimating these other quantities (while they can create quite different Biomass series)
  can only do so much to improve (expand) RP inference as compared with correctly
  modeling P.

## 943 1 Introduction

- the delay model: J. Schnute (1985) J. Schnute (1987) Fournier and Doonan (1987).
- discrete: Hilborn and Walters (1992, pg. 334)
- Walters (2020)

947

automatic accounting for cohort cycles

# 948 2 Methods

## 49 2 .1 Delay Differential Model

Age structured fisheries models typically assume 950 Von Bertalanffy (1938, VB) gorwth in length 951 with age. To model weight the assumption of 952 VB growth in length is composed with a power law relating length to weight,  $w = al^b$ . Since b954 is usually  $\sim 3$  this composition of assumed func-955 tional forms typically results in a monotonically 956 increasing sigmoidal curve of weight with age. When  $b \leq 1$  weight at age takes a VB-like form 958 with b = 1 resulting in an exact correspondence 959 of simulations VB-growth in length and weight. 960

The delay model slightly abridges these relationships by directly assuming VB growth in weight as follows,

$$w(a) = w_{\infty}(1 - e^{-\kappa(a - a_0)}).$$
 (3.1)

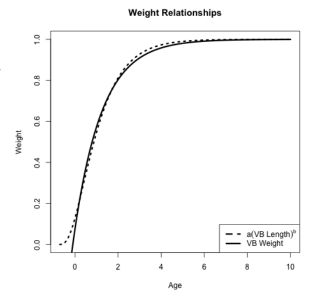


Figure 3.1: The typical composition of allometric weight (b=3) with VB growth in length, as approximated by VB growth in weight directly.

 $\kappa$  is a parameter that controls the instantaneous rate of individual growth (in weight) with age.  $w_{\infty}$  is the maximum weight of individuals in the population, and w(a) is the average

weight of an individual at age a. The parameter  $a_0$  controls the age at which individuals 963 are assummed to have zero weight; by letting  $a_0 < 0$  this allows fish of age zero to have 964 positive weight. Rather than taking a sigmoidally increasing function, VB growth directly in 965 weight results in an monotonically inceasing curve that asymptotes with a strictly decreasing 966 growth rate with age. (only a good approximation for older ages where growth begins to decline) 968

Together with VB growth, the delay model is derived from the assumption that both 969 natural mortality and fishing selectivity are separately proportional to a common heavyside 970 step function with age. That is to say, before a threshold age of selectivity,  $a_s$ , the population 971 is assumed not to experience any mortality whatsoever, but all fish older then  $a_s$  experience 972 the same rate of natural mortaility. Simulaneously all fish older than  $a_s$  are equally vulnerable 973 to fishing (i.e. knife edge selectivity at age  $a_s$ ), although fishing effort may vary from through 974 time. 975

Walters (2020) shows that within these assumptions the following delay differential system of equations exactly models the population dynamics of the total exploitable biomass B(t) and number of indivuduals N(t) through time.

$$\frac{dB}{dt} = w(a_s)R(B;\theta) + \kappa \left[w_{\infty}N - B\right] - (M+F)B \tag{3.2}$$

$$\frac{dB}{dt} = w(a_s)R(B;\theta) + \kappa \left[w_{\infty}N - B\right] - (M+F)B$$

$$\frac{dN}{dt} = R(B;\theta) - (M+F)N$$
(3.2)

This formulation separates the number of individuals in the population from the biomass 976 of the population. The dynamics of N, as seen in Eq (3.3), are very similar to that of the 977 production models previously presented, however the role of the production function is now 978 filled by a "recruitment" function, R(B), which describes the number of new individuals 979 recruiting into the expoitable population as a function of exploitable biomass. In turn, the 980 biomass dynamics are coupled to the numbers dynamics by the assumption of VB growth 981 with growth parameters appearing in Eq (3.2), converting population numbers into biomass 982 and accounting for the growth of biomass with age. 983

Eq (3.2) of the above model expands the notion of biomass production into the processes of recruitment, individual growth, and maturity. The term  $w(a_s)R(B;\theta)$  represents the

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biomass of new recruits; with  $w(a_s)$  representing the weight of individuals at the age of 986 maturity,  $a_s$ , and  $R(B;\theta)$  representing the number of new recruits entering the exploitable 987 population at time t. The negative term, (M+F)B, represents all causes of mortality as 988 it is applied to biomass. Finally, the term  $\kappa \left[ w_{\infty} N - B \right]$  accounts for the net growth of the 989 existing biomass by discounting the limiting maximal individual growth rate by metabolic weight loss proportional to B(t). This term, together with the delay structure in R, provides 991 the major computational savings of the delay differential setting, as compared with full age 992 structured models, by automatically keeping track of changes in the mean size and growth 993 associated with changes in recruitment as cohorts mature into the population. 994

Often a BH functional form is assumed for the stock recruitment relationship, but any adequatly flexible family of functions may model this relationship. For the sake of evaluating the adequacy of assumed BH recruitment the simulation setting below is derived for the delay model under the assumption of the generalized three parameter Schnute recruitment as follows.

$$R(B; [\alpha, \beta, \gamma]') = \alpha B(t - a_s) (1 - \beta \gamma B(t - a_s))^{\frac{1}{\gamma}}$$
(3.4)

The parameters  $\theta' = [\alpha, \beta, \gamma]$  function similarly in this setting as previously described in Section (??). That said, since the delay model explicitly parses out growth in it's dynamics, these parameters only describe the net processes of larval production, and maturation into the population, where as the production model used these parameters to also model the net effects of growth on biomass production. The  $\gamma$  parameter generalizes the family to model varying degrees of decreasing recruitment for large biomasses as  $\gamma$  increases. The Schnute function is exactly equivalent to BH recruitment at the special case when  $\gamma = -1$ , it passes through the Ricker model as  $\gamma \to 0$ , and Logistic recruitment occurs when  $\gamma = 1$ .

Since the delay model assumes knife edge selectivity, at age  $a_s$ , the term  $B(t-a_s)$  appears in R. That is to say fish recruiting into the exploitable population are the result of larval production of biomass  $a_s$  time units in the past. This is because fishing selectivity is only assumed to occur for fish that are at least  $a_s$  time units old and thus fish younger than  $a_s$  are not exploitable. This waiting period requires that new recruits be the result of spawning

biomass  $a_s$  time units in the past. Modeling maturity in this way results in dynamics equations which are a system of delay differential equations as opposed to the simple ODEs that arrise in the production model setting.

- ~ interpretation of recruitment (larval production, recruitment) [growth external] vs. production (larval production, recruitment, growth)
- general structure: Walters (2020) Hilborn and Walters (1992, pg. 334)
- growth: Von Bertalanffy (1938)
- recruitment: J. Schnute (1985); J. T. Schnute and Richards (1998)

#### 1016 2.2 Reference Points

Deriving reference points for the delay model under Schnute recruitment is conceptually similar to the production model setting. The additional nonlinear VB growth assumptions along side Schnute recruitment quickly make the expressions look somewhat unweildy, although analytical solutions can still be derived for most of the same quantities (although complicated by growth parameters).

Starting from Eqs. (3.2) and (3.3), setting both  $\frac{dB}{dt}$  and  $\frac{dN}{dt}$  simultaneously equal to zero, and solving for B and N as a function of fishing, gives the equilibrium biomass and numbers equations.

$$\bar{B}(F) = \frac{1}{\beta \gamma} \left( 1 - \left( \frac{(F+M)(F+M+\kappa)}{\alpha w(a_s)(F+M+\frac{\kappa w_{\infty}}{w(a_s)})} \right)^{\gamma} \right)$$
(3.5)

$$\bar{N}(F) = \frac{\alpha \bar{B}(F)(1 - \beta \gamma \bar{B}(F))^{1/\gamma}}{F + M}$$
(3.6)

Eq. (3.6) is just  $\frac{R(B)}{F+M}$ , and is coupled to  $\bar{B}(F)$  where most of the dynamics appear. Eq. (3.5) resembles Eq (2.3) from the simple production model setting although the growth parameters  $\kappa$ ,  $w_{\infty}$  and  $w(a_s)$ , make slight adjustments to the balance of the maximum rate of recruitment and mortality rate to give an expression for equilibrium biomass that accounts for the factors of individual growth.

Expressions for  $B_0$  and  $B^*$  are attained by evaluating  $\bar{B}(F)$  at F=0 and  $F=F^*$  respectively. Calculation of  $F^*$  typically involves maximization of equilibrium yield,  $\bar{Y}=F\bar{B}(F)$ .

While it was not possible to analytically maximize  $\bar{Y}$ , stable numerical solutions for calculating  $F^*$  were obtained by numerically solving for the roots of the analytical derivative of equilibrium yield with respect to F. Below a greatly simplified expression for  $\frac{d\bar{Y}}{dF}$  is shown; the substitution Z=F+M (total mortality rate) has been made to produce a more compact expression.

$$\frac{d\bar{Y}}{dF} = \frac{1}{\beta\gamma} \left[ 1 - \left( \frac{Z(Z+\kappa)}{\alpha w(a_s)(Z + \frac{\kappa w_\infty}{w(a_s)})} \right)^{\gamma} - \left( \frac{\gamma F}{\alpha w(a_s)} \right) \left( \frac{Z(Z+\kappa)}{\alpha w(a_s)(Z + \frac{\kappa w_\infty}{w(a_s)})} \right)^{\gamma-1} \left( 1 + \frac{\left( \frac{\kappa w_\infty}{w(a_s)} \right) \left( \kappa - \frac{\kappa w_\infty}{w(a_s)} \right)}{(Z + \frac{\kappa w_\infty}{w(a_s)})^2} \right) \right]$$
(3.7)

 $F^*$  is calculated as the numerical root, w.r.t. F, of the above expression. The numerical root is calculated using the base R uniroot function which employs a derivative free search given by Brent (1973).

#### 1037 BH Constraint

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In the simple production model the BH con-1038 strained RPs are fixed to  $\frac{1}{x+2}$ . In the delay 1039 differential modeling setting the constrained 1040 BH RP set is complicated by the growth pa-1041 rameters  $a_s$  and  $\kappa$ . Under BH recruitment 1042 these parameters of the delay model slightly 1043 influence this relationship as seen in Figure 1044 (3.2). That said, the influence of  $a_s$  and  $\kappa$ 1045 on RPs is still largly limited to a confined 1046 region of reference point space which resem-1047 bles the  $\frac{1}{x+2}$  form. In fact the confined region 1048 of RPs is bounded above by  $\frac{1}{x+2}$ . In Figure 1049 (3.2) notice that for values of  $a_s$  and  $\kappa$  that 1050 result in high  $w(a_s)$  (high values of  $\kappa$  and 1051 small values of  $a_s$  seen in red) the BH RP 1052

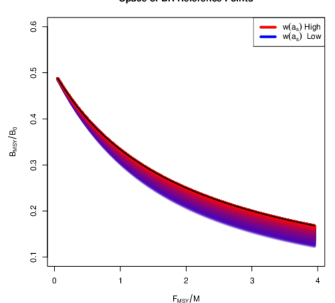


Figure 3.2: The space of BH RPs for the delay model as a function of  $\kappa$  and  $a_s$ . The RP space is plotted for  $80 \times 80$  combinations of  $\kappa \in [0.1, 2]$  and  $a_s \in [0.1, 10]$ . The color drawn is the resulting value of  $w(a_s)$  mapped between blue and red.  $\frac{1}{x+2}$  is plotted in black for reference.

space converges to  $\frac{1}{x+2}$  as derived in the simple production model setting. In opposition to

the simple production model limit, when  $w(a_s)$  is low (as seen in the more blue region of Figure (3.2)), RPs decrease as the influence of growth in the dynamics increases.

## <sup>1056</sup> 2 .3 Delay Differential Integration

The delay model belongs to a class of differential equations known as delay differential 1057 equations (DDE). The delay arrises from the  $B(t-a_s)$  terms found in the recruitment 1058 function. Solving DDEs require special care which depends on the nature of the time delay. 1059 The addition of time-varying delays, many different delays, or very small delays (delays 1060 below the step size of the numerical integrator) results in some of the more challenging 1061 settings for solving DDEs. However with a single stationary model of the age of selectivity, 1062 the delay model in this setting represents one of the most straight forward DDE structures. 1063 The most numerically challenging case presented here arrises in the case of the limiting 1064 production model when  $a_s \to 0$  while  $\kappa \to \infty$ . That said the limiting production model can 1065 be approximated for values of  $a_s \approx 0.1$ , and it was straightforward to ensure that the step 1066 size of the integrator remained reasonably below 0.1. 1067

The DDE presented here is integrated with the initial values fixed at  $B_0$  and  $N_0$  as given 1068 by Eqs. (3.5) and (3.6) with F=0 at any given configuration of  $\theta$  and growth parameters. 1069 The system given in Eqs. (3.2) and (3.3) are then solved numerically using the implicit 1070 Livermore Solver (Isode) as implemented in the dede function of the R package deSolve 1071 (Soetaert et al., 2010). The dede solver provides many methods for integrating DDEs, but 1072 lsode was chosen because it is an implicit method that runs relatively quickly with a relatively 1073 smaller footprint in system memory as compared with other methods. The radau method 1074 was also tried in more computationally challenging settings with good results (albeit running 1075 more slowly that Isode). Ultimately the simulated parameter space did not produce DDEs 1076 that require the more expensive radau integrator to solve accurately. 1077

## 2.4 Simulation Design

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Similarly as previously described in Section (0.1) the relationship between RPs  $\mapsto \theta$  cannot be fully expressed analytically for the Schnute delay model. However, just as in the production model setting, simulation only requires enough knowledge of these mappings to gather

a list of  $(\alpha, \beta, \gamma)$  tuples and the corresponding RPs in some reasonable space-filling design over RP space.

In the delay model a partial mapping for  $(F^*, B_0) \mapsto (\alpha(\cdot, \gamma), \beta(\cdot, \cdot, \gamma))$  can be derived analytically in terms of RPs and  $\gamma$ . The substitution  $Z^* = F^* + M$  is made where  $F^*$  and M appear together to produce a more compact expression.

$$\alpha = \left[ \left( \frac{Z^*(Z^* + \kappa)}{w(a_s)(Z^* + \frac{\kappa w_{\infty}}{w(a_s)})} \right)^{\gamma} + \left( \frac{\gamma F^*}{w(a_s)} \right) \left( \frac{Z^*(Z^* + \kappa)}{w(a_s)(Z^* + \frac{\kappa w_{\infty}}{w(a_s)})} \right)^{\gamma - 1} \left( 1 + \frac{\left( \frac{\kappa w_{\infty}}{w(a_s)} \right) \left( \kappa - \frac{\kappa w_{\infty}}{w(a_s)} \right)}{(Z^* + \frac{\kappa w_{\infty}}{w(a_s)})^2} \right) \right]^{\frac{1}{\gamma}}$$
(3.8)

$$\beta = \frac{1}{\gamma B_0} \left( 1 - \left( \frac{M(M + \kappa)}{\alpha w(a_s)(M + \frac{\kappa w_\infty}{w(a_s)})} \right)^{\gamma} \right)$$
(3.9)

Above Eq. (3.8) results from setting Eq. (3.7) equal to zero and solving for  $\alpha$ , and Eq. (3.9) results from solving the  $\bar{B}(0)$  expression, as derived from Eq. (3.5), for  $\beta$ . The system is completed by further working with the  $\frac{\bar{B}(F^*)}{\bar{B}(0)}$  expression, as seen below, to identify  $\gamma$ .

The system formed by collecting Eqs. (3.8), (3.9), and (3.10) can be navigated similarly

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$$\frac{B^*}{B_0} = \frac{1 - \left(\frac{(F^* + M)(F^* + M + \kappa)}{\alpha w(a_s)(F^* + M + \frac{\kappa w_{\infty}}{w(a_s)})}\right)^{\gamma}}{1 - \left(\frac{M(M + \kappa)}{\alpha w(a_s)(M + \frac{\kappa w_{\infty}}{w(a_s)})}\right)^{\gamma}}$$
(3.10)

to Eq. (2.9) in the Schnute production model setting. For a population experiencing natural 1088 mortality M, VB growth with paramters  $\kappa$  and  $w_{\infty}$ , and age of selectivity  $a_s$  the above 1089 system can fully specify  $\alpha$  and  $\beta$  for a given  $\gamma$ , by fixing  $F^*$ ,  $B_0$ , and  $\frac{B^*}{B_0}$ . For a given  $\gamma$  a 1090 cascade of closed form solutions for  $\alpha$  and  $\beta$  can be obtained, just as in Section (0.1). First 1091  $\alpha(\gamma)$  can be computed, and then  $\beta(\alpha(\gamma), \gamma)$  can be computed. If  $\alpha(\gamma)$  is filled back into the 1092 expression for  $\frac{B^*}{B_0}$ , the system collapses into a single onerous expression for  $\frac{B^*}{B_0}(\alpha(\gamma), \gamma)$ . For 1093 brevity, define the function  $\zeta(\gamma) = \frac{B^*}{B_0}(\alpha(\gamma), \gamma, F^*, M)$  based on Eq. (3.10). 1094 Again rather than inverting  $\zeta(\gamma)$  for  $\gamma$ ,  $\gamma$  is the sampled so that the overall simulation 1095 design is space filling as described in Section (0.2). Given the sampled  $\gamma$ , the cascade of 1096  $\alpha(\gamma)$ , and then  $\beta(\alpha(\gamma), \gamma)$ , can be computed, and the Schnute delay model is fully defined 1097 by a given  $(\frac{F^*}{M}, \frac{B^*}{B_0})$ . While conceputally this framing is similar to the Schnute production 1098

parameters appear explicitly here. Other ways of navigating the RPs  $\mapsto \theta$  system are possible, but for the sake of numerical stability this strategy has proven the most reliably accurate by limiting exposure to numerical error propagation.

Each design location defines a complete Schnute delay differential model with the given RP values. Indices of abundance are simulated from the Schnute model at each design location, a small amount of residual variation,  $\sigma = 0.01$ , is added to the simulated index, and the data are then fit with a misspecified BH model. The design captures various degrees of model misspecification relative to the BH model, so as to observe the effect of recruitment misspecification upon RP inference.

point to catch, and LHS design, and Metamodel.

#### o 2 .5 Parameter Estimation

• I use B only here

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• quick statement of inference, and reference to previous section

Let  $I_t$ ,  $t \in \{1, 2, 3, ..., T\}$ , be a series of indicies of abundance, proportional to biomass, as simulated from the Schnute Delay model. These data are modelled with the following log-normal observation model that has been intentionally constrained to BH recruitment,

$$I_t \sim LN(qB_t(\boldsymbol{\theta}, \boldsymbol{\phi}), \sigma^2).$$
 (3.11)

 $B_t(\boldsymbol{\theta}, \boldsymbol{\phi})$  is the biomass solution of the BH constrained DDE system. The BH constraint isimplemented by fixing  $\gamma = -1$  so that  $\boldsymbol{\theta}' = [\alpha, \beta, \gamma = -1]$ .  $\boldsymbol{\phi}$  is a vector of growth and maturity parameters,  $\boldsymbol{\phi}' = [\kappa, w_{\infty}, a_0, a_s]$ . The nuisance parameter q models the proportionality constant of the index with process biomass, and  $\sigma^2$  models residual variation of the index.

In this setting,  $\phi$  and q are fixed to focus on the inferential affects of model misspecification on recruitment parameters and RPs. Without an explicite mechanism for the delay model to incorporate age data, under the BH model  $\phi$  is not well informed and would tyically be estimated externally for data limted stocks. Under BH recruitment  $\phi$  can only slightly impact RPs as seen in Figure (3.2).

 $\sigma^2$  and  $\theta$  are reparameterized to the log scale and fit via MLE. Reparameterizing the 1123 parameters to the log scale improves the reliability of optimization, in addition to facili-1124 tating the use of Hessian information for estimating MLE standard errors. Given that the 1125 biological parameters enter the likelihood via a nonlinear differential equation, and further 1126 the parameters themselves are related to each other nonlinearly, the likelihood function can 1127 often be difficult to optimize. A hybrid optimization scheme is used to maximize the log 1128 likelihood to ensure that a global MLE solution is found. The R package GA (Scrucca, 2013, 1129 2017) is used to run a genetic algorithm to explore parameter space globally. Optimization 1130 periodically jumps into the L-BFGS-B local optimizer to refine optima within a local mode. 1131 The scheme functions by searching globally, with the genetic algorithm, across many initial 1132 values for starting the local gradient-based optimizer. The genetic algorithm serves to iter-1133 atively improve hot starts for the local gradient-based optimizer. Additionally, optimization 1134 is only considered to be converged when the optimum results in an invertible Hessian at the 1135 found MLE. 1136

- fixed M = 0.2,  $a_0 = -1$ ,  $w_{\infty} = 1$ 
  - play with  $\kappa$  and age of selectivity  $a_s$

#### 1139 Numbers Indicies

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While not utilized here, age structured models may commonly model indicies as proportional to numbers rather than (or simultaiously to) biomass. When solving the DDE, Eq. (3.3) points out that the full DDE solution will expose a numbers solution simultaneously with a biomass solution that may be used for these purposes. These solutions are often quite similar since the main driver of process behavior comes from the form of R which is shared among N and B. However, it is common on the west coast of the US that indicies derived from commercial fisheries are measured as weights while indicies derived from recreational fisheries are often measured as counts. If a numbers index,  $J_t$ , is observed alongside the previously mentioned biomass index, the following likelihood component is often added as a conditionally independent component of the likelihood,

$$J_t \sim LN(pN_t(\boldsymbol{\theta}, \boldsymbol{\phi}), \tau^2).$$
 (3.12)

 $N_t(\boldsymbol{\theta}, \boldsymbol{\phi})$  is the numbers solution of the DDE system.  $\boldsymbol{\theta}$  and  $\boldsymbol{\phi}$  are the productivity and growth parameters shared in common with the biomass component. p and  $\tau^2$  are then the analogous proportionality constant and residual variation of the numbers index respectively.

## 1143 2.6 GP Metamodel

1145

point to catch, and LHS design, and Metamodel.

## 2.7 Clustering Catastrophic Model Failure

Considering the behavior observed in Section (1.1), where  $\frac{F_{MSY}}{M}$  is dramatically underesti-1146 mated, it is very natural to ask where specifically in RP space we might see this catastrophic 1147 failure of the BH model. The structure of RPs under the BH model suggests several natural 1148 avenues for forming hypotheses to identify this misspecified RP region. The single clearest 1149 feature to identify are cases where  $\frac{F_{MSY}}{M}$  is estimated well below the minimum values sim-1150 ulated. In the follow section this idea is formally stated in a hypothesis testing structure 115 that uses the GP metamodel as a surrogate for the sampling distribution of  $\frac{F_{MSY}}{M}$  under the 1152 misspecified BH model. This allows for a rejection threshold to be derived in terms of the 1153 GP predictive structures that defines a classifier for identifying BH inference break points 1154 broadly over RP space. 1155 For simplicity in outlining the hypothesis testing framework, let  $\theta$  be the metamodeled 1156

For simplicity in outlining the hypothesis testing framework, let  $\theta$  be the metamodeled population parameter, corresponding with  $\frac{F_{MSY}}{M}$ , under the two parameter BH model and let  $\theta$  be the metamodeled population parameter, corresponding with  $\theta$  under the two parameter BH model and let  $\theta$  be the metamodeled population parameter, corresponding with  $\theta$  under the two parameter BH model and let  $\theta$  may be formally represented by the following hypothesis.

$$H_0: \theta \ge \theta_{min} \quad H_a: \theta < \theta_{min}$$
 (3.13)

For evaluating the hypothesis test, the metamodel prediction serves as a descriptor of RP estimation using kriging to provide both a measure of mean behavior as well as propogating estimate uncertainty via the kriging predictive variance. The metamodeled quantity is then predicted by  $N(\hat{y}(\mathbf{x}), \hat{\sigma}^2(\mathbf{x}))$ , where  $\hat{y}(\mathbf{x})$  is as previously described in Eq. (2.16) and  $\hat{\sigma}^2(\mathbf{x})$ 

propagates estimate uncertainty via the kriging predictive variance given by,

$$\hat{\sigma}^2(\mathbf{x}) = \mathbf{R}(\mathbf{x}, \mathbf{x}) - \mathbf{r}(\mathbf{x})' \mathbf{R}_{\ell}^{-1} \mathbf{r}(\mathbf{x}). \tag{3.14}$$

For evaluating the hypotheses given in Eq. (3.13)  $H_0$  is then rejected when  $\hat{y}(\mathbf{x})$  is small in the following sense

$$R = \{ \mathbf{x} : \ \hat{y}(\mathbf{x}) < C \}. \tag{3.15}$$

To calibrate exactly how small the constant C should be the false positive rate,  $\alpha_0$ , is considered. For evaluating R under the null hypothesis, the domain of the metamodel is limited along the axis of  $\theta$  as,  $\mathbf{x}_{min} = [\theta_{min}, \mathbf{x}_{(-\theta)}]$ . Thinking of  $\hat{y}(\mathbf{x})$  as an estimator of  $\theta$ , via kriging, the following test statistic distibution follows naturally under the null,

$$\frac{\hat{y}(\mathbf{x}_{min}) - \theta_{min}}{\hat{\sigma}(\mathbf{x}_{min})} \sim N(0, 1). \tag{3.16}$$

This stratforwardly implies  $\alpha_0 = \Phi\left(\frac{C - \theta_{min}}{\hat{\sigma}(\mathbf{x}_{min})}\right)$ , and rearranging for C gives  $C = \theta_{min} + \Phi^{-1}(\alpha_0)\hat{\sigma}(\mathbf{x}_{min})$ . Replacing C into Eq. (3.15), the rejection region can be made specific as

$$R = \{ \mathbf{x} : \hat{y}(\mathbf{x}) < \theta_{min} + \Phi^{-1}(\alpha_0)\hat{\sigma}(\mathbf{x}_{min}) \}.$$
(3.17)

Other hypotheses may be formed using a similar structure (or added to this structure) to fine tune the rejection threshold, although a point of deminishing returns will quickly set in for describing the catestophic model failure under low contrast BH inference. By simple inspection of Figure (2.8), clearly the under estimation of  $\frac{F_{MSY}}{M}$  is a very powerful classifer for describing BH inference trends.

## 3 Results

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Figure (3.3)shows three hypothetical 1177 individual-growth/maturity curves that span 1178 a wide range of RPs. As seen in Figure (3.2), 1179 the larger values of  $w(a_s)$  correspond to less 1180 dramatic growth with the red curve demon-1181 strating the simle (no growth) production 1182 model limit  $(a_s \to 0 \text{ and } \kappa \to \infty)$ . The cases 1183 with smaller  $w(a_s)$  values (blue and purple 1184 curves) correspond to more dramatic growth 1185 behaviors, with the blue curve where  $a_s = 2$ 1186 and  $\kappa = 0.1$  representing the most dramatic 1187 growth shown here. 1188

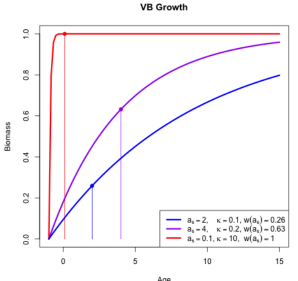


Figure 3.3: Three hypothetical individualgrowth curves, showing  $w(a_s)$  on each curve.

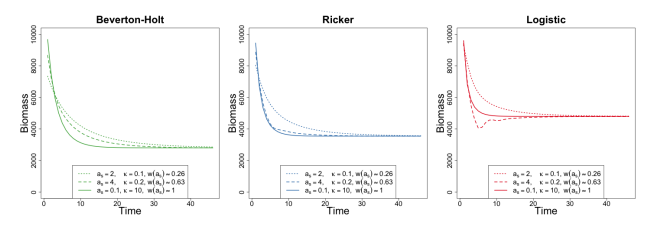


Figure 3.4: Biomass dynamics of BH (left), Ricker (center), and Logistic (right) delay differential models in the low contrast simulation setting. In all cases  $\alpha = 1.2$  and  $\beta$  is chosen so that each model shares the same  $B_{MSY}$  within each given  $\gamma$ .

Figure (3.4) demonstrates a range of biomass dynamics that the Schnute delay model can display under a spectrum of growth behaviors with fishing held consistent at  $F_{MSY}$ . The three special cases of  $\gamma = -1$  (BH),  $\gamma \to 0$  (Ricker), and  $\gamma = 1$  (Logistic) recruitment are shown in each of the above shown growth configurations. Notice under the most dramatic growth ( $a_s = 2$  and  $\kappa = 0.1$ ) setting, biomass of the Logistic model comes into equilibrium at

 $B_{MSY}$  as an oscillating curve. This effect occurs here due to the Logistic model's relatively high  $\frac{B^*}{B_0}$  interacting with the lag in selectivity upon the sudden onset of fishing; this produces a shock that pushes biomass past  $B_{MSY}$  setting up an oscillatory pattern of recruitment. One may also observe these oscillations under the Ricker model by exaggerating the  $a_s$  lag as well as the steepness of the Ricker curve. The BH model may also demonstrate these oscillations, in a heavily lagged setting, by shocking the population past its relatively low  $B_{MSY}$  as a

sudden release in fishing applied to a heavily:

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Figure (3.5) shows the range of RPs that 1201 can be modeled with each of the BH, Ricker, 1202 and Logistic recruitments over the spectrum 1203 of individual-growth/maturitymodels simu-1204 lated here. Notice that the more dramatic 1205 the growth, the further the RP curve lies 1206 from the simple production model, but each 1207 recruitment model reacts differently under 1208 each of the given growth parameters. The 1209 Ricker and BH RP-spaces are qualitatively 1210 similar in shape with more dramatic growth 1211 settings decreasing  $\frac{B_{MSY}}{B_0}$  relative to the sim-1212

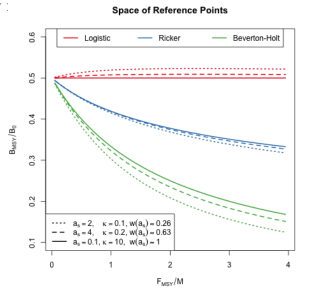


Figure 3.5: Restricted RP-space under each recruitment models, with each growth curve.

ple production model setting. The Logistic model on the other hand increases  $\frac{B_{MSY}}{B_0}$  relative to the simple production model setting as growth parameters become more dramatic. It is also worth noting that the Ricker model's RPs are much less influenced by growth parameters as compared with that of the BH or Logistic model.

## 3.1 Simple Production Model Limit

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Under the delay differential's limiting simple production model ( $a_s = 0.1$  and  $\kappa = 10$ ), the expectation is that RP inference should be identical to that of the model seen in Chapter (2). By way of verifying this equivalence, Figure (3.6) demonstrates a virtually identical pattern of RP biases as previously seen in Figures (2.7) and (2.8) (under both of the high and low contrast settings).

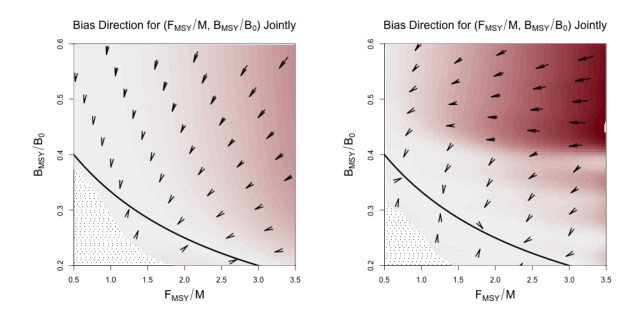


Figure 3.6: RP mapping of BH delay model fit to Schnute delay data under the simple (no growth) production model limit. left: High contrast simulation. Right: Low contrast simulation.

Indeed in the high contrast setting, Figure (3.6, left) shows how the BH model induces 1223 the same pattern of bias as seen in Chapter (2). There is bias in both RPs (in accordance 1224 with the  $\frac{B^*}{\bar{B}(0)} = \frac{1}{F^*/M+2}$  RP-set) so as to produce a nearly minimal distance mapping of 1225 RPs onto the constrained BH set of RPs. Similarly, in the low contrast setting, Figure (3.6, 1226 right) again shows the same two regions pattern of RP inference. Firstly, there is a region 1227 of relatively small model misspecification where the minimal distance mapping is preserved. 1228 Secondly, as model misspecification becomes greater (around the Ricker set)  $\frac{F^*}{M}$  begins to 1229 be sharply underestimated. Above this break point in RP estimation inference appears to 1230 be driven to the trivial RP  $\frac{F^*}{M} = 0$ ,  $\frac{B^*}{\bar{B}(0)} = 0.5$ ) that is shared in common amoung all of the 1231

two-parameter models described here.

These results merely confirm that the theoretical limiting dynamics do indeed replicate expected RP inference patterns as previously observed in Chapter (2).

#### 3.2 Moderate Growth

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Moving past verification of the simple production model, other values of  $a_s$  and  $\kappa$  provide a probe into the effects individual growth dynamics may have on RP inference. Individual growth is a multifaceted phenomena that is not easily reduced to a single number, but for the purposes of this model  $w(a_s)$  serves as a decent proxy for the extent of the model dynamics that are due to individual growth. This follows from the intuition that individuals maturing at a smaller fraction of  $w_{\infty}$  demonstrate the dynamics of growth during an observable (to the model) phase rather than growth occurring prior to selection.

That said,  $w(a_s)$  is not a one-to-one map of  $\kappa$  and  $a_s$ . A level curve of  $w(a_s; \kappa) = c$  is attained by increasing the value of  $a_s$  and decreasing  $\kappa$  corrispondingly, or vice versa. The case where  $a_s = 4$  and  $\kappa = 0.2$  (resulting in  $w(a_s) \approx 0.6$ ) respresents a plausibly biological example of moderate growth. Similar examples of the  $w(a_s) = 0.6$  level curve result in much larger lags (discussed in Section (3.5)) or larger  $\kappa$ 's which quickly tend toward behaviors previously described in the simple production model setting.

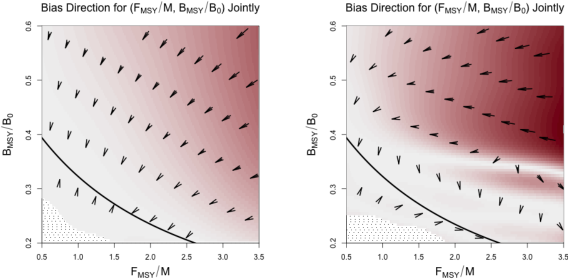


Figure 3.7: RP mapping of BH delay model fit to Schnute delay data under moderate growth ( $a_s = 4$  and  $\kappa = 0.2$ ). Left: High contrast simulation. Right: Low contrast simulation.

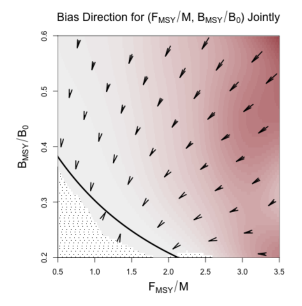
The RP mappings seen in Figure (3.7) show very similar RP mappings to that of the 1249 simple production model, with the biggest differences occurring around the location of the 1250 break point where the low contrast model begins to dramatically underestimate  $\frac{F^*}{M}$ . In the 1251 high contrast simulation setting Figure (3.7; left), the RP mappings again demonstrate a 1252 nearly identical minimal distance mapping of RPs onto the constrained BH RP set. In the 1253 low contrast setting Figure (3.7; right) a very similar two regiem pattern of RP inference is 1254 observed, however the location of the break between these regiems appears at lower values of 1255  $\frac{B^*}{B(0)}$ . In this moderate growth setting the break point occurs around values of  $\frac{B^*}{B(0)}$  just below 1256 0.4 where in the simple production model the break point occurs at  $\frac{B^*}{B(0)}$  just above 0.4. 1257

## 1258 3.3 Emphatic Growth Dynamics

The emphatic growth setting simulated here fixes  $a_s = 2$  and  $\kappa = 0.1$ , to simulate a species that grows quite slowly and yet matures into the reproducing stock at a relatively early age. This combination has the effect of exaggerating the components of the model dynamics which are related to individual growth since individuals recruit at a small size and slowly grow over the extent of the modeled period.

The slow growth of these dynamics oppose the simple production model setting in the sense that they move the constrained RP set a large distance (largest amoung the spectrum of decreasing  $w(a_s)$  populations simulated here) away from the  $\frac{1}{x+2}$  limiting case. It is interesting to note that this is true for all of the two parameter constrained constrained RP sets as seen in Figure (3.5).

Despite the emphaticly growth driven dynamics in this setting, the RP mappings seen 1269 in Figure (3.8) obviously bare a huge resemblance to the previously seen RP mappings. 1270 Again the biggest differences in the RP mappings occur around the location of the break 1271 point where the low contrast model begins to dramatically underestimate  $\frac{F^*}{M}$ . In this low 1272 contrast setting the break point in RP estimation occurs around values of  $\frac{B^*}{\overline{B}(0)}$  well below 1273 0.4 with the behaviour extending as far down as  $\frac{B^*}{B(0)} = 0.3$ . This region shift occurs well 1274 below that of the Ricker set, as initially observed in the production model setting. This 1275 reduced range of acceptible RP inference indicates that under increasingly emphatic growth 1276 the model misspecification issue of the BH model becomes an increasingly brittle assumption 1277



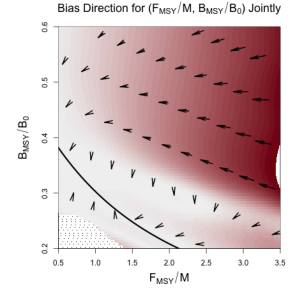


Figure 3.8: RP mapping of BH delay model fit to Schnute delay data under dramatic growth  $(a_s = 2 \text{ and } \kappa = 0.1)$ . Left: High contrast simulation. Right: Low contrast simulation.

1278 with respect of RPs.

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Interestingly this pattern only follows for the low contrast setting. In the high contrast setting inference returns to a pattern resmbleing the minimal distance mapping onto BH RP set. Further pointing to the importance of contrast for informing these models.

## 3.4 Clustering Catastrophic Model Failure

Figure (3.9) shows the rejection thresholds 1283 for the low contrast simulations of each of 1284 the emphatic, moderate, and no growth set-1285 tings. The dark lines represent the rejection 1286 threshold with a false positive rate of about 1287 15%, and the light shaded regions show 1288 how the rejection threshold changes as the 1280 false positive rate rages from 50% to 2.25%. 1290 When applied to the high contrast simula-1291 tions the rejection threshold falls outside of 1292 the simulated RP range as expected by in-1293 spection of the high contrast RP mappings. 1294

Notice in Figure (3.9) that the rejection

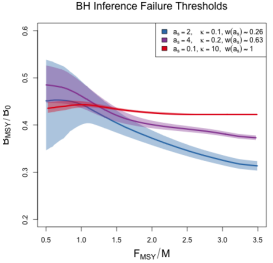


Figure 3.9: BH RP estimation failure threasholds with increasingly emphatic individual growth dynamics.

threshold is subject to two axese of sensativity. Firstly, for each simulated growth the rejection threshold is more sensative for small values of  $\frac{F_{MSY}}{M}$  than for large values. This is a natural result since discerning  $\hat{y}(x)$  below the minimum simulated RP becomes more difficult when the data are truely generated near the minimum simulated  $\frac{F_{MSY}}{M}$ . For large  $\frac{F_{MSY}}{M}$  the minimum distance mapping results in  $\hat{y}(x)$  well above the minimum simulated RP but for small  $\frac{F_{MSY}}{M}$  even the minimum distance mapping may be close to the rejection threshold.

The second axis of sensativity is between individual growth simulations. The no growth setting produces a very clear threshold of model failure, while the failure threshold for emphatic growth is much more varied, especially near the minimum simulated  $\frac{F_{MSY}}{M}$ . This is largely due to the increased RP estimate uncertainty as growth becomes more emphatic in the dynamics.

Model misspecification of the BH model is compounded for the more emphatic growth settings as recruitment can interact with growth dynamics to produce unique behaviors as exemplified in Section (3.5).

## 3.5 Ocillatory Growth Influence

While the above patterns of RP estimation follow for biological regiems of the  $w(a_s; \kappa) = c$  level curve, as  $a_s$  increases an ocillatory regiem also exists within these dynamics. While RP estimation behaves similarly in this ocillatory regiem there are unique features in this setting that are not present in the more biological regiems. Below consider the ocillatory example of a logistic delay model with  $a_s = 10$  fixing fishing at  $F_{MSY}$ .

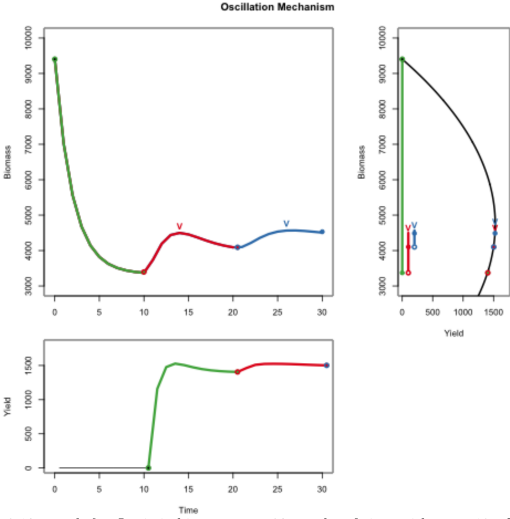


Figure 3.10:  $top\ left$ : Logistic biomass over 30 epochs of time with  $a_s=10$ . Green, red, and blue colors indicate three 10 epoch long windows of biomass. v indicates local biomass ocillation maxima.  $top\ right$ : Yield plotted over the range of biomasses shown. The biomass range of each 10 epoch window is shown in the vertical colored lines.  $bottom\ left$ : Yield plotted through time. Colors correspond to the lagged biomass region that results in the evaluated yield. The black horizontal line demonstrates the pre-model assumption of biomass fixed at  $B_0$ .

Figure (3.10) demonstrates the mechanism of how these oscillatory dynamics form. Oscillatory dynamics appear when fishing pushes biomass past  $B_{MSY}$  within the lagged  $a_s$ window of recruitment. The delay model assumes that biomass is fixed in equilibrium at  $B_0$ ,
for  $t \leq 0$ . Therefore in the green region of the biomass series, 0 < t < 10, the population
recruits at R(B0). Figure (3.10) shows that in this initial period R(B0) results in zero yield
for that period, and biomass falls as a result.

Once t exceeds  $a_s$ , the lagged recruitment refers to the integrated biomass series to evaluate recruitment based on  $R(B_{t-a_s})$ . The red region of the biomass series is the result of yield over the initial green biomasses. Figure (3.10) shows that the yield over the green biomass series first increases, as biomass approaches  $B_{MSY}$  and then decreases as biomass passes  $B_{MSY}$ . This creates the local maximum in the red biomass series.

Furthermore, the blue region of the biomass series is then based on yield over the red biomasses. Notice that since the red biomasses first increase and then decrease, yield increases as the red biomass increases toward  $B_{MSY}$ , and yield subsequently decreases following
the descending leg of the red biomass series. This yield pattern carries the ocillation of the
red biomass region forward into the blue region.

This process of biomass ocillation carries on in this manner nonetheless approaching equilibrium at  $B_{MSY}$ . Equilibrium is reached in an ocillatory manner setoff by the green biomass series crossing over from above  $B_{MSY}$  to below it. The example shown in Figure (3.10) exemplifies the oscillatory phenomena simulated here, but the mechanism that produces these oscillations may occur with other forms of recruitment outside of logistic recruitment whenever fishing cases biomass to cross over  $B_{MSY}$  within the lagged recruitment window.

#### 1339 RP Estimation

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Statistical inference in the oscillatory regiem can be challenging. Depending on the parameters inferred, the likelihood can have multiple local modes which require global optimization techniques to distiguish. Furthermore, parameter estimation is more uncertain in this setting as the likelihood may confuse oscillations with residual noise.

Figure (3.11) shows the BH RP mapping fixing  $w(10;0.1) \approx 0.6$  in the high contrast

simulation setting. This places the dynamics firmly in the ocillatory regiem, but the high

contrast setting provides significant information for inferring recruitment parameters.

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Interestingly in this high contrast set-1348 ting, a very similar two regiem pattern of 1349 RP inference is observed as previously seen 1350 in low contrast settings. That said the 1351 boundary between the regiems in this setting 1352 is much smoother and the location of the 1353 break between these regiems appears around 1354 higher values of  $\frac{B^*}{\bar{B}(0)}$ . 1355

This higher  $\frac{B^*}{\overline{B}(0)}$  break point, hovering around 0.5, is consistent with the mechanism which induces ocillation. Starting the biomass at  $\overline{B}(0)$  in the ocillatory regiem, in-

BMSX/M Pin 2.11 DD (DH 1)

Bias Direction for (F<sub>MSY</sub>/M, B<sub>MSY</sub>/B<sub>0</sub>) Jointly

Figure 3.11: RP mapping of BH delay model fit to high contrast Schnute delay data under ocillatory growth ( $a_s = 10$  and  $\kappa = 0.1$ ).

creased  $\frac{B^*}{B(0)}$  will tend to exasterbate oscillatory behavior by increasing  $B_{MSY}$  so that biomass is more easily pushed past  $B_{MSY}$  within the initial lagged as window of recruitment. This produces more dramatic oscillations in the higher  $\frac{B^*}{B(0)}$  region of RP space.

The fitted BH model does not produce 1363 significant ocillations because under the BH 1364 model  $\frac{B^*}{\overline{B}(0)}$  is constrained below 0.5 with 1365 the majority of the simulation BH  $\frac{B^*}{\overline{B}(0)}$  RPs 1366 falling between 0.4 and 0.2. Therefore, 1367 the fitted BH model will not tend to push 1368 biomass past  $B_{MSY}$  and thus is incapable 1369 of modeling oscillatory biomass series. Fig-1370 ure (3.12) shows a subset of example BH 1371 fits, which demonstrats the limited oscilla-1372 tory capacity of the BH fits. Furthermore, 1373 since the BH model has a limited oscillatory 1374

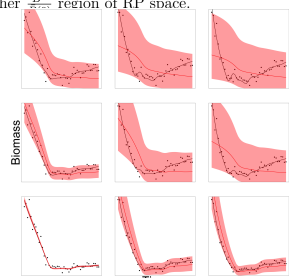


Figure 3.12: Example BH fits (red) to Schnute data (black). Each example plot is arranged to mirror its location in RP space.

capacity in this setting, the BH model tends

to explain the oscillations with artifically high residual variation and artifically low steepness focusing on overly simplistic trends in the data.

#### 1378 Estimating More

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Figure (3.13) shows a subset of example 1379 model fits broadly over RP space. Model 1380 fits are shown both under the two-parameter 1381 BH model as well as under the three pa-1382 rameter Schnute model, each model estimat-1383 ing all of its recruitment parameters as well 1384 as the growth and maturity parameters  $\kappa$ 1385 and  $a_s$ . Notice that the BH model, even 1386 when additionally estimating  $\kappa$  and  $a_s$ , does 1387 not gain the flexibility to properly model 1388 Schnute data. 1389

The lack of oscillatory dynamics produced by the BH model causes the misspecified BH fits in Figure (3.13) to largely estimates  $\kappa$  and  $a_s$  so as to approximate the production model limiting case. The fitted

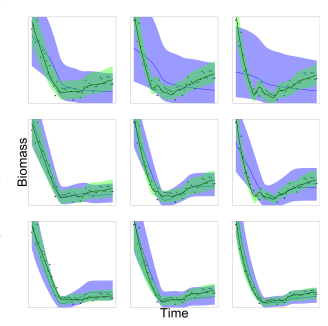


Figure 3.13:  $\kappa$  and  $a_s$  estimation under BH (blue) and Schnute (green) fits to Schnute data (black) arranged to mirror RP space.

Schnute model can produce the oscillatory dynamics and thus the information in the oscillatory data well inform estimates of  $\kappa$  and  $a_s$  under the Schnute model. Furthermore, the Schnute model has no issue learning its  $\gamma$  parameter.

While Statistical inference in the oscillatory region can be challenging in the highly constrained BH model, the Schnute model can easily estimate its extra  $\gamma$  parameter. The flexibility of estimating  $\gamma$  simplifies inference by correctly specifying RPs, and also by opening up the model dynamics to reveal additional information about  $\kappa$  and  $a_s$  in the data.

# 1402 4 Discussion

- break point decreases with growth
- inference becomes more brittle with more dramtic growth.
- interaction between assumed form of growth and stock recruitment.
- low-side steepness bias masks ocillatory/shock patterns induced by growth and maturity parameters
- misspecified BH prevents learning growth
- increasing growth accelerates model misspecification
- statistical evidence of minimum distance mapping within accepible regiem, although
  float idea of PT-like pattern as BH set flattens. (explaining perterbations)

## $_{1412}$ 5 old ideas

- show production model limit (contrast
- $-a_s \rightarrow 0$ : instant maturity
- $-\kappa \to \infty$ : recruit as an adult ()
- describe second order shapes of growth/maturity (and cause)
- weight of recruits => scaling biomass  $(q, \beta, \text{ and } w_{\infty})$
- 1418 —
- describe RP bias
- flat

- summary of  $\sigma$  over RP space comparing between models (PT, Schnute, Schnute DD) to show areas of model breakdown.
- miss-identifying signal for noise.
- It happens more as the dynamics get more complex.
- point to the full age structed models.
- show the constrained BH space over a grid of M,  $\kappa$ ,  $\omega$ ,  $W_{\infty}$
- Show that the constrained spaces vary only slightly as compared with the consequences of misspecifing the functional form.
- estimating these other quantities (while they can create quite different Biomass series)
  can only do so much to improve (expand) RP inference as compared with correctly
  modeling P.
- mapping distance as a function of contrast at (3.5, 0.5)
- for LHS grid locations show  $\frac{B^*}{B_0}$  and  $F^*$  biases for grids in  $M \in (0, 0.5)$  For sure in High Contrast, maybe also in Low??.

# <sup>1435</sup> 6 Appendix: Inverting $\frac{B^*}{\overline{B}(0)}$ and $\gamma$ for the PT Model

For brevity let  $\zeta = \frac{B^*}{\bar{B}(0)}$ .

$$\zeta = \left(\frac{1}{\gamma}\right)^{\frac{1}{\gamma - 1}}$$

$$\zeta = \gamma \zeta^{\gamma}$$

$$\zeta = \gamma e^{\gamma \log(\zeta)}$$

$$\zeta \log(\zeta) = \gamma \log(\zeta) e^{\gamma \log(\zeta)}$$

The Lambert product logarithm, W, is defined as the inverse function of  $z = xe^x$  such that x = W(z). Applying this definition allows for the isolation of  $\gamma$ .

$$\gamma \log(\zeta) = W(\zeta \log(\zeta))$$

$$\gamma = \frac{W(\zeta \log(\zeta))}{\log(\zeta)}$$
(3.18)

The Lambert product logarithm is a multivalued function with a branch point at  $-\frac{1}{e}$ . The principal branch,  $W_0(z)$ , is defined on  $z \in \left(-\frac{1}{e}, \infty\right)$ , and the lower branch,  $W_{-1}(z)$ , is defined on  $z \in \left(-\frac{1}{e}, 0\right)$ . Taken individually, each respective branch is analytic, but cannot be expressed in terms of elementary functions.

When  $\zeta \in \left(0, \frac{1}{e}\right)$  the solution of interest in Eq. (1.12) comes from  $W_0$ . When  $\zeta \to \frac{1}{e}$  the

When  $\zeta \in (0, \frac{1}{e})$  the solution of interest in Eq. (1.12) comes from  $W_0$ . When  $\zeta \to \frac{1}{e}$ , the Fox Model emerges as  $\gamma \to 1$ . When  $\zeta \in (\frac{1}{e}, 1)$  the solution of interest comes from  $W_{-1}$ . For the use case presented here, Eq. (1.12) is to be interpreted as,

$$\gamma = \begin{cases}
\frac{W_0(\zeta \log(\zeta))}{\log(\zeta)} & \zeta \in (0, \frac{1}{e}) \\
\frac{W_{-1}(\zeta \log(\zeta))}{\log(\zeta)} & \zeta \in (\frac{1}{e}, 1)
\end{cases}$$
(3.19)

# 7 Appendix: The Replacement Line for the Delay Model

The replacement line is the rate of productivity which exactly balances biomass loss in the absence of fishing. In the simple production model, productivity must simply balance biomass loss due to M. Thus when  $R(B;\theta) > MB$  there will be some surplus productivity to enable fishing.

In the delay model, productivity is complicated by biomass changing, both with the recruitment of young into the reproducing population, as well as biomass accumulation due to the growth of existing individuals in the population. To derive the replacement line in the case of the delay model the equilibrium equations in the absence of fishing are considered and the R(B) that this implies is then isolated.

$$0 = \frac{dB}{dt} = w(a_s)R(B) + \kappa \left[w_{\infty}N - B\right] - MB \tag{3.20}$$

$$0 = \frac{dN}{dt} = R(B) - MN. \tag{3.21}$$

Eq(3.21) quickly gives  $\bar{N} = R(B)/M$ . Substituting this equilibrium value into Eq(3.20) to rewrite N in terms of B,

$$0 = w(a_s)R(B) + \kappa \left[ w_{\infty} \frac{R(B)}{M} - B \right] - MB. \tag{3.22}$$

Collecting like terms,

$$R(B)\left[w(a_s) + \frac{\kappa w_\infty}{M}\right] = [M + \kappa] B. \tag{3.23}$$

Finally solving for R(B), and simplifying, gives the equation of the replacement line as,

$$R(B) = \left[ \frac{M(M+\kappa)}{w(a_s)M + \kappa w_{\infty}} \right] B. \tag{3.24}$$

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