# Metamodeling for Bias Estimation of Fisheries

# Reference Points Under Two Parameter Production

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5 Abstract

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  $\frac{B^*}{B(0)}$  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. Mangel et al. note that use of a three-parameter stock-recruitment relationship allows for independent estimation of these reference points. This research seeks to understand the 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 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 into two dimensions.

# 19 1 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-21 nary differential equations (ODE). Key management quantities called reference points (RPs) are commonly derived from the ODE equilibrium equations and depend upon the parameter-23 ization of biomass production. Two-parameter forms of the production function have been 24 shown to limit the theoretical domain of RPs (Mangel et al., 2013). The limited RP-space 25 of two-parameter models makes these models vulnerable to model misspecification with re-26 spect to RPs, and thus the limiting stucture of two-parameter models may in and of itself 27 induce bias in RP estimation using these models. The behavior of RP estimation is not well 28 understood and as a result patterns of bias in RP estimation may easily go unnoticed. A 29 metamodeling approach is developed here to describe RP biases and explore mechanisms of 30 model failure under the most common two-parameter models. 31

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 modelled population that is vulnerable to fishing. The index is often observed alongside a variety of other known quantities, but at a minimum, each index will be observed in the presence of some known catch for the period. Figure (1) shows the classic Namibian Hake dataset (Punt, 1988; Hilborn & Mangel, 1997; Mangel, 2006) exemplifying the form.

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

$$I_t = qB_t e^{\epsilon} \quad \epsilon \sim N(0, \sigma^2). \tag{1}$$

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.

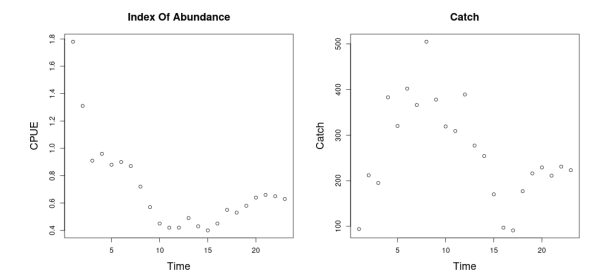


Figure 1: *left*: An index of abundance data, catch per unit effort (CPUE), for Namibian Hake from 1965 to 1987. *right*: The associated catch data for Namibian Hake over the same time period.

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

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$$\frac{dB}{dt} = P(B(t); \boldsymbol{\theta}) - Z(t)B(t). \tag{2}$$

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

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 some 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 57 it doesn't leave biomass in the population to reproduce in the future. We seek to fish in a way 58 that allows (or even encourages) future productivity in the population. This is accomplished 59 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  $\bar{Y} = F\bar{B}(F)$ , 61 where  $\bar{p}$  indicates a value at steady state. MSY is found by maximizing  $\bar{Y}(F)$  with respect to 62 F, and  $F^*$  is the fishing rate at MSY. Going forward let \* decorate any value derived under 63 the condition of MSY. 64 Fisheries are very often managed based upon reference points which serve as simplified 65 heuristic measures of population behavior. The mathematical form of RPs depends upon the model assumptions through the production function. While a number of different RPs 67 exist which describe the population in different (but related) ways, the most common RPs 68 revolve around the concept of MSY (or robust ways of measuring MSY (Hilborn, 2010; Punt 69 et al., 2016)). Here the focus is primarily on the RPs  $\frac{B^*}{\bar{B}(0)}$  and  $F^*$  ( $\frac{F^*}{M}$  when appropriate) for 70 their pervasive use in modern fisheries (Punt & Cope, 2019). 71  $F^*$  is the afore mentioned fishing rate which results in MSY.  $\frac{B^*}{\overline{B}(0)}$  is the depletion of the 72 stock at MSY. That is to say  $\frac{B^*}{\bar{B}(0)}$  describes the fraction of the unfished population biomass 73

 $F^*$  is the afore mentioned fishing rate which results in MSY.  $\frac{B^*}{\overline{B}(0)}$  is the depletion of the stock at MSY. That is to say  $\frac{B^*}{\overline{B}(0)}$  describes the fraction of the unfished population biomass that will remain in the equilibrium at MSY. In general  $F^* \in \mathbb{R}^+$  and  $\frac{B^*}{\overline{B}(0)} \in (0,1)$ , however under the assumption of two-parameter production, models will be structurally unable to capture the full theoretical range of RPs.

Many of the most commonly used production functions depend only on two-parameters. For example, the Schaefer model depends only on the biological parameters r and K, and limits RP inference so that under the Schaefer model  $\left(F^*, \frac{B^*}{B(0)}\right) \in (\mathbb{R}^+, \frac{1}{2})$ . The two-parameter Fox model (Fox Jr., 1970) limits  $\left(F^*, \frac{B^*}{B(0)}\right) \in (\mathbb{R}^+, \frac{1}{e})$ . Similarly the two-parameter Cushing (Cushing, 1971), Beverton-Holt (Beverton & Holt, 1957, BH) and Ricker (Ricker, 1954) production functions do not model the full theoretical space of RPs (Mangel et al., 2013; Yeakel & Mangel, 2015).

The bias-variance trade-off (Ramasubramanian & Singh, 2017) makes it clear that the addition of a third parameter in the production function will necessarily reduce estimation bias.

However the utility of this bias reduction is still under debate because the particular mechanisms and behavior (direction and magnitude) of these biases for key management quantities are not fully understood or described. Lee et al. (2012) provides some evidence that estimation of productivity parameters are dependent on changes in biomass trend through time (i.e. 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 trade-offs 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>106</sup> 2 Methods

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## 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,

$$P_p(B; [r, K, \gamma]) = \frac{rB}{\gamma - 1} \left( 1 - \left( \frac{B}{K} \right)^{(\gamma - 1)} \right). \tag{3}$$

 $\gamma$  is a parameter which breaks PT out of the 108 restrictive symmetry of the logistic curve. In gen-109 eral  $\gamma \in (1, \infty)$ , with the logistic model appear-110 ing in the special case of  $\gamma = 2$ , and the Fox model appearing as a limiting case as  $\gamma \to 1$ . The 112 parameter r controls the maximum per-capita 113 growth rate of the population in the absence of 114 competition for resources (i.e. the slope of pro-115 duction function at the origin). K is the so called "carrying capacity" of the population. In this 117 context the carrying capacity can be formally 118 stated as steady state biomass in the absence of 119 fishing (i.e. B(0) = K). In Figure (2) PT produc-120 tion is shown for a range of parameter values so as 121 to demonstrate the various productivity shapes 122 that can be achieved under PT. 123

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.

#### 129 2.1.1 PT Reference Points

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

$$\frac{dB}{dt} = \frac{rB}{\gamma - 1} \left( 1 - \left( \frac{B}{K} \right)^{\gamma - 1} \right) - FB. \tag{4}$$

An expression for the equilibrium biomass is attained by setting Eq (4) equal to zero, and rearranging the resulting equation to solve for B. Thinking of the result as a function

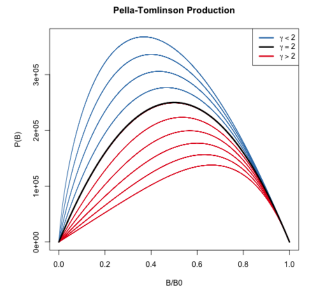


Figure 2: The Pella-Tomlinson production function plotted across a variety of parameter values. The special cases of Logistic production is shown in black, and the left-leaning and right-leaning regimes are shown in blue and red respectively.

of F gives,

$$\bar{B}(F) = K \left( 1 - \frac{F(\gamma - 1)}{r} \right)^{\frac{1}{(\gamma - 1)}}.$$
 (5)

At this point it is convenient to notice that  $\bar{B}(0) = K$ . The expression for  $B^*$  is given by evaluating Eq (5) at  $F^*$ . To get an expression for  $F^*$ , the equilibrium yield is maximized with respect to F,

$$F^* = \operatorname*{argmax}_F F\bar{B}(F). \tag{6}$$

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

$$\frac{d\bar{Y}}{dF} = \bar{B}(F) + F\frac{d\bar{B}}{dF} \tag{7}$$

$$\frac{d\bar{B}}{dF} = -\frac{K}{r} \left( 1 - \frac{F(\gamma - 1)}{r} \right)^{\frac{1}{\gamma - 1} - 1}.$$
 (8)

Setting Eq (7) equal to 0, substituting  $\bar{B}(F)$  and  $\frac{d\bar{B}}{dF}$  by Equations (5) and (8) respectively, and solving for F produces the following expression for the fishing rate required to produce MSY,

$$F^* = \frac{r}{\gamma} \tag{9}$$

Plugging the above expression for  $F^*$  back into Eq (5) gives the following expression for biomass at MSY,

$$B^* = K \left(\frac{1}{\gamma}\right)^{\frac{1}{\gamma - 1}}. (10)$$

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.

$$F^* = \frac{r}{\gamma} \qquad \frac{B^*}{\bar{B}(0)} = \left(\frac{1}{\gamma}\right)^{\frac{1}{\gamma - 1}} \tag{11}$$

#### 35 2.1.2 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

$$r = \gamma F^* \qquad \qquad \gamma = \frac{W\left(\frac{B^*}{\overline{B}(0)}\log\left(\frac{B^*}{\overline{B}(0)}\right)\right)}{\log\left(\frac{B^*}{\overline{B}(0)}\right)}. \tag{12}$$

Above W is the Lambert product logarithm function. More details about this derivation, and the Lambert product logarithm, are given in Appendix (5).

Using Eq. (12) to obtain production parameters, a PT production model can be fully defined for any combination of the RPs  $F^*$  and  $\frac{B^*}{B(0)}$ . 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 (2.3). 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.

#### $_{\scriptscriptstyle 146}$ 2.2 Schnute 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  $\gamma$ ,

$$P_s(B; [\alpha, \beta, \gamma]) = \alpha B (1 - \beta \gamma B)^{\frac{1}{\gamma}}.$$
 (13)

The BH and Logistic production func-147 tions arise when  $\gamma$  is fixed to -1 or 1 respec-148 tively. The Ricker model is a limiting case 149 as  $\gamma \to 0$ . For  $\gamma < -1$  a family of strictly in-150 creasing Cushing-like curves arise, culminating in linear production as  $\gamma \to -\infty$ . These 152 special cases form natural regimes of simi-153 larly behaving production functions as seen 154 in Figure (3). 155

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

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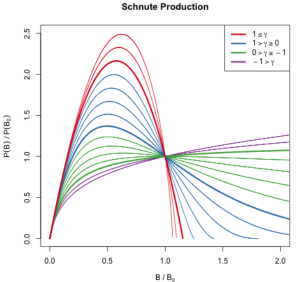


Figure 3: The Schnute production function plotted across a variety of parameter values. Regimes of similarly behaving curves are grouped by color.

wide variety of possible productivity behaviors, they present an ideal simulation environment for inquiry of the reliability of inference under the BH assumption.

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

$$\frac{dB}{dt} = P_s(B;\theta) - (M+F)B. \tag{14}$$

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 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. (14) 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

$$\bar{B}(F) = \frac{1}{\gamma \beta} \left( 1 - \left( \frac{M+F}{\alpha} \right)^{\gamma} \right). \tag{15}$$

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

$$B_0 = \frac{1}{\gamma \beta} \left( 1 - \left( \frac{M}{\alpha} \right)^{\gamma} \right) \tag{16}$$

$$\frac{B^*}{B_0} = \frac{1 - \left(\frac{M + F^*}{\alpha}\right)^{\gamma}}{1 - \left(\frac{M}{\alpha}\right)^{\gamma}}.$$
 (17)

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

$$\frac{d\bar{Y}}{dF} = \bar{B}(F) + F\frac{d\bar{B}}{dF} \tag{18}$$

$$\frac{d\bar{B}}{dF} = -\frac{1}{\beta} \left( \frac{\left( \frac{M+F}{\alpha} \right)^{\gamma}}{F+M} \right). \tag{19}$$

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,

$$0 = \frac{1}{\gamma} - \left(\frac{1}{\gamma} + \frac{F^*}{F^* + M}\right) \left(\frac{F^* + M}{\alpha}\right)^{\gamma}.$$
 (20)

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  $\gamma$ .

Since RPs are left without a closed form expression, computing RPs from productivity parameters amounts to numerically solving the system formed by collecting the expressions (20), (16), and (17).

#### 176 2.2.1 Simulation

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

Similarly to J. T. Schnute and Richards (1998), expressions (20) and (16) 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. (17), to identify  $\gamma$ , the following system is obtained,

For a population experiencing natural mortality M, by fixing  $F^*$ ,  $B_0$ , and  $\frac{B^*}{B_0}$  the above

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

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

$$\frac{B^*}{B_0} = \frac{1 - \left( \frac{M + F^*}{\alpha} \right)^{\gamma}}{1 - \left( \frac{M}{\alpha} \right)^{\gamma}}.$$
(21)

system can fully specify  $\alpha$  and  $\beta$  for a given  $\gamma$ . Notice for a given  $\gamma$  a cascade of closed 182 form solutions for  $\alpha$  and  $\beta$  can be obtained. First  $\alpha(\gamma)$  can be computed, and then 183  $\beta(\alpha(\gamma), \gamma)$  can be computed. If  $\alpha(\gamma)$  is filled back into the expression for  $\frac{B^*}{B_0}$ , the system 184 collapses into a single onerous expression for  $\frac{B^*}{B_0}(\alpha(\gamma), \gamma)$ . For brevity, define the function 185  $\zeta(\gamma) = \frac{B^*}{B_0} (\alpha(\gamma), \gamma, F^*, M)$  based on Eq. (17). 186 Inverting  $\zeta(\gamma)$  for  $\gamma$ , and computing the cascade of  $\alpha(\gamma)$ , and then  $\beta(\alpha(\gamma), \gamma)$ , fully 187 defines the Schnute model for a given  $(\frac{F^*}{M}, \frac{B^*}{B_0})$ . However inverting  $\zeta$  accurately is extremely 188 difficult. Inverting  $\zeta$  analytically is not feasible, and numerical methods for inverting  $\zeta$  are 189 unstable and can be computationally expensive. Rather than numerically invert precise 190 values of  $\zeta(\gamma)$ ,  $\gamma$  is sampled so that the overall simulation design is space filling as described 191 in Section (2.3.2). 192 Each design location defines a complete Schnute production model with the given RP 193 values. Indices of abundance are simulated from the Schnute model at each design location, 194 a small amount of residual variation,  $\sigma = 0.01$ , is added to the simulated index, and the data 195 are then fit with a misspecified BH production model. The design at large captures various 196 degrees of model misspecification relative to the BH model, so as to observe the effect of 197 productivity model misspecification upon RP inference. 198

#### 2.3 Latin Hypercube Sampling 199

The goal of space filling design in this setting is to extend the notion of the random sample 200 (and its desirable parameter estimation properties) across the simulated RP domain so as 201 to represent the simulated space as well as possible (Gramacy, 2020). The simple random 202 sample is the classical approach to unbiased parameter estimation, however simple random-203 ness is patchy, often sampling some regions of design space quite densely, while leaving other 204 regions of design space empty. Space filling designs aim to preserve (or enhance) parameter 205 estimation properties across the simulated domain (Devon Lin & Tang, 2015; Stein, 1987), 206 while constraining samples to be spaced in some notion of spread over the entire space. 207 Latin hypercube sampling (McKay et al., 2000, LHS) is among the most foundational of 208 space filling designs used in computer experiments. 209

A LHS of size n, in the 2 dimensional space defined by RPs, distributes samples so as to spread points across a design region in a broadly representative way. A LHS design extends the notion of a univariate random uniform sample across multiple dimensions so that each margin of the design space enjoys a uniform distribution.

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LHS designs achieve this notion of uni-218 formity by first partitioning each dimension 219 of the design space into regular grids of size 220 n. By intersecting the grids of each dimen-221 sion, cells are produced that evenly partition 222 the design space. In two dimensions  $n^2$  cells 223 are produced, from which a total of n samples are taken. Crucially only one sample is 225

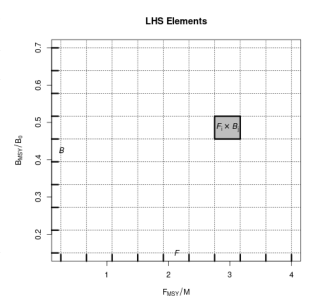


Figure 4: LHS grids. Intersecting  $\mathcal{F}$  and  $\mathcal{B}$ produces  $n^2$  cells; a particular cell  $\mathcal{F}_i \times \mathcal{B}_j$  is shown in grey. Maybe just show points.

taken from a given element of each grid in each dimension so as to reduce clumping of the 226 n samples across the design space. 227

#### 228 2.3.1 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 the PT model are computed directly in RP space and Eq. (12) is used to map the sampled RP design locations to PT productivity parameters.

#### 236 2.3.2 Schnute Design

Due to the lack of an analytical relationship mapping RPs  $\mapsto \theta$ , analogous to the PT model's Eq. (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 (2.2.1), allows an approximate LHS to be obtained by a careful navigation of the system of equations seen in Eq. (21).

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 approximate 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 5: 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,

$$\gamma \sim \zeta_{min}\delta(\gamma_{min}) + t(\mu, \sigma, \nu)\mathbf{1}_{\gamma > \gamma_{min}}.$$
 (22)

Above, t is the density of the three-247 parameter location-scale family Student's t 248 distribution with location  $\mu$ , scale  $\sigma$ , and 249 degrees of freedom  $\nu$ .  $\mathbf{1}_{\gamma>\gamma_{min}}$  is an indica-250 tor function that serves to truncate the Stu-251 dent's t distribution at the lower bound  $\gamma_{min}$ . 252  $\delta(\gamma_{min})$  is the Dirac delta function evaluated 253 at  $\gamma_{min}$ , which is scaled by the known value 254  $\zeta_{min}$ ; this places probability mass  $\zeta_{min}$  at 255 the point  $\gamma_{min}$ . Since sampling from a Stu-256 dent's t distribution is readily doable, sam-257 pling from a truncated Student's t mixture 258 only requires slight modification. 259

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

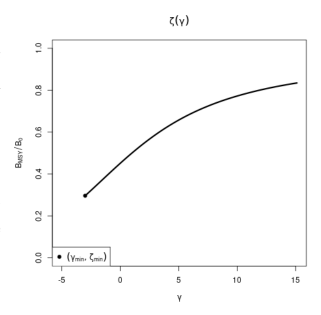


Figure 6:  $\zeta(\gamma)$  Plotted for  $F^* = 0.1$  and M = 0.2. The point  $(\gamma_{min}, \zeta_{min})$  shows the lowest biologically meaningful value of  $\gamma$ ; below which productivity is negative.

known from the dynamics of the Schnute model at a given RP, full specification of Eq. (22) 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  $\zeta(\gamma)$ .

$$[\hat{\mu}, \hat{\sigma}, \hat{\nu}] = \underset{[\mu, \sigma, \nu]}{\arg\min} \int_{\Gamma} \left( T(\gamma; \mu, \sigma, \nu) - \zeta(\gamma) \right)^2 d\gamma \tag{23}$$

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The distribution T(\gamma|\hat{\mu}, \hat{\sigma}, \hat{\nu}) is fit for use
                                                                     Algorithm 1 LHS of size n on rectangle R.
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                                                                       1: procedure LHS_n(R)
     in generating \gamma^* random variates at a spe-
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                                                                       2:
                                                                               Define n-grids \mathcal{F}, \mathcal{B} \in R
     cific F^* and M. This approximation releases
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                                                                               for each grid element i do
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     the need to invert \zeta w.r.t \gamma by using sam-
                                                                                    Draw \frac{F^*}{M} \sim Unif(\mathcal{F}_i)
     ples of \gamma^* values to generate approximatly
                                                                       4:
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                                                                                    Compute [\hat{\mu}, \hat{\sigma}, \hat{\nu}] given F^* \& M
                                                                       5:
     uniform samples of \zeta(\gamma^*). By sampling ap-
265
                                                                                    while \mathcal{B}_j not sampled do
                                                                       6:
     proximatly uniform \zeta(\gamma^*) random variates in
266
                                                                                        Draw \gamma^* \sim T(\gamma | \hat{\mu}, \hat{\sigma}, \hat{\nu})
                                                                       7:
     this way, and making use of the structure in
267
                                                                                        Compute \zeta^* = \zeta(\gamma^*)
                                                                      8:
     Eq. (21), an approximate LHS sample can
268
                                                                                         Compute j such that \zeta^* \in \mathcal{B}_i
                                                                      9:
     be collected via Algorithm (1).
269
          \frac{F^*}{M} is drawn uniformly from \mathcal{F}_i. Con-
                                                                                    end while
                                                                     10:
270
                                                                                    Compute \alpha^* = \alpha(\gamma^*, F^*, M)
     ditioning on the sample of F^*, and M,
                                                                     11:
271
                                                                                    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:
272
                                                                                    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-
273
                                                                               end for
     ate grid element \mathcal{B}_j. Given \gamma^*, the cascade
                                                                     14:
274
                                                                     15: end procedure
     \alpha(\gamma^*), and \beta(\alpha(\gamma^*), \gamma^*), can be computed.
275
     The algorithm continues until all of the de-
276
```

#### 278 2.3.3 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 289 design (Morris & Mitchell, 1995; Devon Lin & Tang, 2015). Maximin designs sample the 290 design space by maximizing the minimum distance between sampled points. This has the 291 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 293 corners) of the design domain. Since LHS ensures uniformity in the margins and maximin 294 designs enjoys a certain sense of optimality in how they define and fill gaps (Johnson et al., 295 1990), the methods are quite complimentary when combined. 296

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

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

$$\boldsymbol{D} = \operatorname{diag} \left[ \left( \max(\mathcal{F}) - \min(\mathcal{F}) \right)^2, \left( \max(\mathcal{B}) - \min(\mathcal{B}) \right)^2 \right].$$
(24)

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,

$$\boldsymbol{x_a} = \underset{\boldsymbol{x'}}{\operatorname{argmax}} \min\{d(\boldsymbol{x_i}, \boldsymbol{x'}) : i = 1, ..., n\}.$$
 (25)

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.

## 316 2.4 Gaussian Process Metamodel

At its core, a metamodel is simply a model of some mapping of inputs to outputs (the 317 mapping itself is typically defined by a computer model). By modeling the mapping with a statistical model (that explicitly defines the relevant features of the mapping) a metamodel 319 defines a specific ontology for the mapping. By simulating examples of the mapping, the 320 inferential infrastructure of the statistical model is used to empirically learn an effective 321 emulation of the mapping within the ontology defined by the statistical model. The pre-322 dictive infrastructure of the statistical model is then useful as an approximate abstraction 323 of the system itself to better understand the system through further data collection, cheap 324 approximation of the mapping, and/or study of the mapping itself. 325

In this setting, the aim of metamodeling is to study how well RPs are inferred when 326 typical two-parameter models of productivity (Logistic and BH) are misspecified for pop-327 ulations that are actually driven by more complicated dynamics. The simulation design, 328 X, provides a sample of different population dynamics that are driven by three-parameter 329 production functions broadly in RP space. By simulating index of abundance data from the 330 three parameter model, and fitting those data with the two-parameter production model, we 331 observe particular instances of how well RPs are inferred at the given misspecification of the 332 two-parameter model relative to the true three-parameter production model. By gathering 333

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 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 correlation 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
which define the simulated three-parameter production models.

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.

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

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

$$\boldsymbol{\epsilon} \sim N_n(\mathbf{0}, \boldsymbol{\omega}' \mathbf{I})$$
(26)

X is the  $n \ge 2$  LHS design matrix of RPs for each simulated three-parameter data generating model as described in Section (2.3.3).  $\epsilon$  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, serves to down weight the influence of each  $y_i$  in proportion to the inferred production model 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 this effect heterogeneously.

The term, v, contains spatially correlated GP effects. The correlation matrix,  $R_{\ell}$  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,

$$R(\boldsymbol{x}, \tilde{\boldsymbol{x}}) = \exp\left(\sum_{i=1}^{2} \frac{-(x_i - \tilde{x}_i)^2}{2\ell_j^2}\right).$$
 (27)

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)

$$\hat{y}(\mathbf{x}) = \beta_0 + \mathbf{x}\boldsymbol{\beta} + \mathbf{r}(\mathbf{x})' \mathbf{R}_{\ell}^{-1} \Big( \mathbf{y} - (\beta_0 + \mathbf{X}\boldsymbol{\beta}) \Big)$$
(28)

 $\hat{y}(\mathbf{x})$  is the predicted value of the modeled productivity parameter MLE under the twoparameter production model, when the index of abundance is generated from the threeparameter production model at RP location  $\mathbf{x}$ .  $\mathbf{r}(\mathbf{x})$  is a vector-valued function of correlation function evaluations for the predictive location  $\mathbf{x}$  against all observations in  $\mathbf{X}$  (i.e.  $\mathbf{r}(\mathbf{x}) =$  $\mathbf{R}(\mathbf{x}, \mathbf{x}_i) \ \forall \ \mathbf{x}_i \in \mathbf{X}$ ). While metamodeling occurs on the inferred productivity parameters of the restricted

production model, the metamodel can also be used to build estimates of major biological RPs. For the BH model the relevant transformations for relating productivity parameters with RPs are given in Eqs. (17, 20) 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 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.

It is known that contrast in the observed index and catch time series can effect inference

## 375 2.5 Catch

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on the productivity parameters (Hilborn & Walters, 1992). In this setting contrast refers to 377 changes in the long term trends of index data. Figure (7, right) demonstrates an example of 378 biomass that includes contrast induced by catch. It is not well understood how contrast may 379 factor into inferential failure induced by model misspecification. Thus catch is parameterized 380 so as to allow for a spectrum of possible contrast simulation settings. 381 Catch is parameterized so that F(t) can be controlled with respect to  $F^*$ . Recall that 382 catch is assumed to be proportional to biomass, so that C(t) = F(t)B(t). To control F(t)383 with respect to  $F^*$ , C(t) is specified by defining the quantity  $\frac{F(t)}{F^*}$  as the relative fishing rate. 384 B(t) is defined by the solution of the ODE, and  $F^*$  is defined by the biological parameters of the model. By defining  $\frac{F(t)}{F^*}$ , catch can then be written as  $C(t) = F^*\left(\frac{F(t)}{F^*}\right)B(t)$ . 386 Intuitively  $\frac{F(t)}{F^*}$  describes the fraction of  $F^*$  that F(t) is specified to for the current B(t). 387 When  $\frac{F(t)}{F^*} = 1$ , F(t) will be held at  $F^*$ , and the solution of the ODE brings B(t) into 388 equilibrium at  $B^*$ . When  $\frac{F(t)}{F^*}$  is held constant in time biomass comes to equilibrium as an 389 exponential decay from K approaching  $B^*$ . When  $\frac{F(t)}{F^*} < 1$ , F(t) is lower than  $F^*$  and B(t) is pushed toward  $\bar{B} > B^*$ . Contrarily, when  $\frac{F(t)}{F^*} > 1$ , F(t) is higher than  $F^*$  and B(t) is pushed 391

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.

toward  $\bar{B} < B^*$ ; the precise values of  $\bar{B}$  can be calculated from the steady state biomass

equations provided above and depend upon the specific form of the production function.

$$\frac{F(t)}{F^*} = ae^{bt} \mathbf{1}_{0 \le t < 15} + (d - ct) \mathbf{1}_{15 \le t < 30} + \mathbf{1}_{30 \le t \le 45}$$
 (29)

The first term of Eq(29) 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.

$$a = e^{\log(\chi_{max}) - 15b} \qquad b = \frac{1}{t - 15} \log\left(\frac{\chi_{min}}{\chi_{max}}\right) \tag{30}$$

$$c = \frac{\chi_{max} - 1}{15 - 1} \qquad d = 15c + \chi_{max} \tag{31}$$

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.

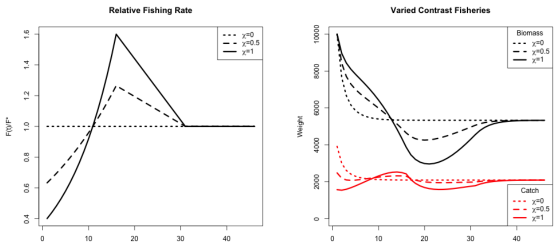


Figure 7: (left) Relative fishing with low, medium, and high confrast. (right) Population biomass and catch at each associated level of contrast.

When  $\chi = 0$ , the relative fishing rate is a constant at 1 to create a low contrast simulation environment. As  $\chi$  increases Eq (29) induces more and more contrast in the observed index and catch time series until  $\chi = 1$  which produces a high contrast simulation environment. Figure (7) demonstrates a spectrum of contrast simulation environments as well as the time series data they induce in the solution of the production model ODE.

#### 2.6 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,

$$I_t|q, \sigma^2, \boldsymbol{\theta} \sim LN(qB_t(\boldsymbol{\theta}), \sigma^2).$$
 (32)

 $B_t(\boldsymbol{\theta})$  is defined by the solution of the ODEs defined by the Schaefer, or BH models. For the Schaefer model  $\boldsymbol{\theta} = [r, K]$ , and for the BH model  $\boldsymbol{\theta} = [\alpha, \beta]$ . 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

In this setting, q is fixed at the true value of 0.0005 to focus on the inferential effects

$$\log \mathcal{L}(q, \sigma^2, \boldsymbol{\theta}; I) = -\frac{T}{2} \log(\sigma^2) - \frac{1}{2\sigma^2} \sum_{t} \log\left(\frac{I_t}{qB_t(\boldsymbol{\theta})}\right)^2.$$
 (33)

of model misspecification on biological parameters.  $\sigma^2$  and  $\boldsymbol{\theta}$  are reparameterized to the log scale and fit via MLE. Reparameterizing the parameters to the log scale improves the 411 reliability of optimization, in addition to facilitating the use of Hessian information for 412 estimating MLE standard errors. 413 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 416 likelihood to ensure that a global MLE solution is found. The R package GA (Scrucca, 2013, 417 2017) is used to run a genetic algorithm to explore parameter space globally. Optimization 418 periodically jumps into the L-BFGS-B local optimizer to refine optima within a local mode. 419 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 found MLE.

An important (and often overlooked) implementation detail is the solution to the ODE which

#### 2.7 Continuous model formulation

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defines the progression of biomass through time. As a statistical model it is of paramount 427 importance that this ODE not only have a solution, but also that the solution be unique. Of 428 primary concern, uniqueness of the ODE solution is necessary for well conditioned inference. 429 If the form of  $\frac{dB}{dt}$  is at least Lipschitz continuous, then the Cauchy-Lipschitz-Picard 430 theorem provides local existence and uniqueness of B(t). Recall from Eq(2) that  $\frac{dB}{dt}$  is 431 separated into a term for biomass production, P(B), and a term for removals, Z(t)B(t). For 432 determining Lipschitz continuity of  $\frac{dB}{dt}$ , the smallest Lipschitz constant of  $\frac{dB}{dt}$  will be the sum 433 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 435 Z(t) typically contains fishing mortality as a function of time F(t) to model catch in time as 436 C(t) = F(t)B(t). Z(t) may or may not contain M, but typically M is modeled as stationary 437 in time and does not pose a continuity issue, unlike some potential assumptions for C(t). 438 In practice C(t) is determined by a series of observed, assumed known, catches. Catch 439 observations are typically observed on a quarterly basis, but in practice may not be complete 440 for every quarter (or year) of the modeled period. It is overwhelmingly common to discretize 441 the ODE in time via Euler's method with integration step sizes to match the observation 442 frequency of the modeled data. This is often computationally convenient when the underlying 443 species dynamics are resonably well behaved, however when the dynamics model is used as a statistical model, with the goal of inferring the behavior of the underlying species dynamics, 445 the regularity of the dynamics are not guaranteed. An implicit assumption of continuity 446 of catch in time provides the necessary regularity for the statistical model. Furthermore 447 a continuous handling of the dynamics provides improved accruacy in evaluating the ODE, 448 particually when inferring productivity parameters which largely control the regularity of 450 the dynamics.

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.

#### 457 2.7.1 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.

# 476 3 Results

#### 3.1 PT/Schaefer

#### 478 3.1.1 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 (8) shows four of the most mis-484 specified example production function fits as 485 compared to the true data generating PT 486 production functions. The rug plots below 487 each set of curves show how the observed 488 biomasses decay exponentially from K to  $B^*$ 489 in each case. In particular, notice how obser-490 vations only exist where the PT biomass is 491 greater than  $B^*$ . Due to the leaning of the 492 true PT curves, and the symmetry of the 493 logistic parabola, the logistic curve only ob-494 serves information about its slope at the ori-495 gin from data observed on the right portion 496 of the PT curves. The top two panels of Fig-497 ure (8) shows PT data generated such that 498  $\frac{B^*}{B(0)} > 0.5$ ; in these cases PT is steeper to the 490 right of  $B^*$  than it is on the left, and so the 500 the logistic curve over-estimates r, and con-501 sequently also over-estimates  $F^*$ . The bot-502

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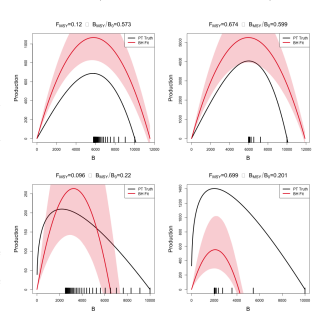


Figure 8: A comparison of the true PT production function (in black) and the estimated logistic curve (in red) with 95% CI shown. The examples shown represent the four corners of maximum model misspecification in the simulated RP-space. Observed biomasses are plotted in the rug plots below the curves.

tom two panels of Figure (8) 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  $F^*$ .

#### 3.1.2 Metamodeled Trends

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Each point in the space of the RPs  $F^*$  and  $\frac{B^*}{B(0)}$  uniquely identifies a complete PT model 507 with different combinations of parameters values. Recall that when  $\gamma = 2$  for the PT model, 508 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 510 value of  $\frac{B^*}{\bar{B}(0)}$  at the constant 0.5 for any value of  $F^*$ . So the line through RP space defined 511 by  $\frac{B^*}{B(0)} = 0.5 \ \forall F^*$ , defines the subset of RP space where  $\gamma = 2$  and where the PT model 512 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 514 set will be the only data that are not misspecified relative to the Schaefer model; as PT data 515 are simulated farther and farther away from this line at  $\frac{B^*}{\overline{B}(0)}=0.5$  model misspecification of 516 the Schaefer model becomes worse and worse. 517

While Figure (8) demonstrates a real trend in simulation results, individual simulation 518 runs will at best show jittery trends due to the stochastic nature of statistical inference. The 519 GP process metamodel accounts for this stochasticity to focus analysis on the signal in the 520 simulation results. Recall that metamodeling occurs on the scale of the inferred productivity 521 parameters of the restricted production model, by transforming metamodel predictions via 522 Eq. (11), metamodeled predictions are obtained for Schaefer RPs. By further subtracting 523 the true data generating PT RPs from the predicted Schaefer RPs at each point in RP space 524 a pattern of inferential RP bias, induced by model misspecification of the Schaefer model, 525 can be seen. 526

Figure (9) 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 (9) 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

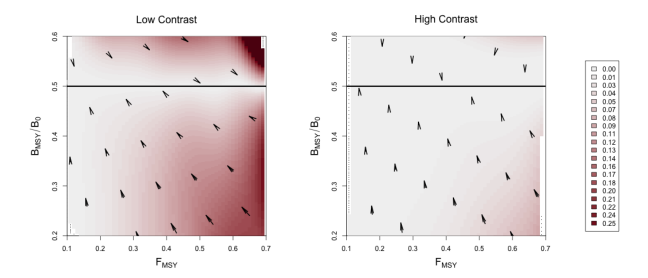


Figure 9: Joint bias direction for  $(F^*, \frac{B^*}{B_0})$  estimates under the misspecified Schaefer Model. The intensity of color represents the excess bias relative to the shortest possible mapping. Results in the low contrast setting are shown left, and the high contrast setting is shown right.

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 (9) are identical except for the amount of contrast present in the simulated index. The left panel of Figure (9) 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 (8), 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 (3.1.2) demonstrates how bias in  $F^*$  estimation decreases as contrast is added to

#### Bias in Estimated Schaefer FMSY

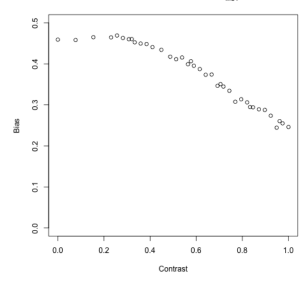


Figure 10: Bias in  $F^*$  under the Schaefer model when PT data are generated with increasing contrast so that  $F^*$  and  $\frac{B^*}{B_0}$  are fixed at 0.699 and 0.201 respectively.

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.

# 553 3.2 Schnute/BH

#### 554 3.2.1 Design

Algorithm (1) enforces uniform marginals in  $\frac{F^*}{M}$ 555 directly, as well as the adherence of the overall design to latin squares. Figure (11) shows a uni-557 form Q-Q plot for sampled  $\zeta$ , using Algorithm 558 (1), against theoretical uniform quantiles. As ev-559 idence by the excellent coherence to the theoret-560 ical uniform quantiles, the approximation in Sec-561 tion (2.3.2) for sampling  $\gamma$  (and therefore  $\zeta(\gamma)$ ), 562 is very effective. Furthermore since numerical in-563 version of  $\zeta(\gamma)$  is costly and unreliable, the rel-564 ative speed and accuracy that this approximate 565

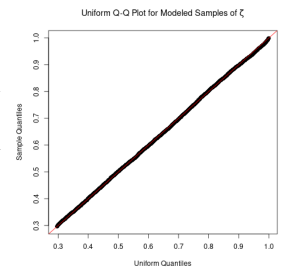


Figure 11: Uniform Q-Q plot for  $\zeta$  plotted for  $F^* = 0.1$  and M = 0.2.

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

Similarly to the PT model, the three-567 parameter Schnute model is uniquely iden-568 tified by each point in the space of  $\frac{F^*}{M}$  and 569  $\frac{B^*}{B_0}$  RPs. As seen in Figure (12), Schnute 570 production has different behaviors in different ranges of RPs space, which are entirely 572 defined by the value of  $\gamma$  (shown in Figure 573 (3)). When  $\gamma \geq 1$  the Schnute model pro-574 duces a family of Logistic-like curves that 575 are increasingly right leaning as  $\gamma$  increases. For  $1 > \gamma \ge 0$ , Schnute production takes 577 a family of left leaning Ricker-like curves 578 that all, at least, approach the x-axis. For 579  $0\,>\,\gamma\,>\,-1$  there are a family of BH-like 580 curves that do not approach the x-axis but 581

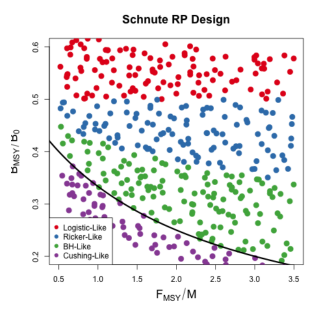


Figure 12: A Schnute RP design. Colors indicate different regimes of Schnute production.

The black curve shows the BH set.

still have decreasing 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 (12). The farther away from this set that Schnute data are simulated, the worse the BH model is misspecified for those data.

#### 3.2.2 Metamodeled Trends

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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

596 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.

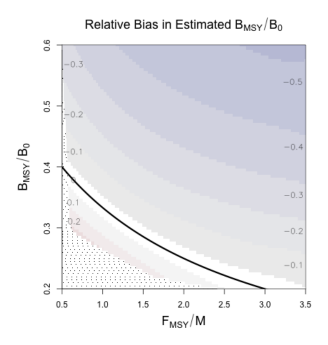
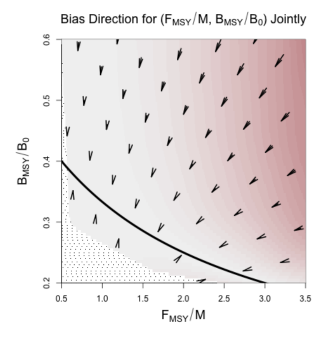
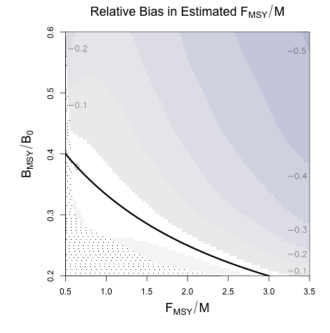


Figure 13: Heatplots showing the bias in RP estimation induced by model misspecification of the BH model in the high contrast simulation setting. In all cases the restricted RP-space of the BH set is shown as the black curve. (left) Relative bias in  $\frac{B^*}{\overline{B}(0)}$ . (top-right) Bias in 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  $F^*$ .





High Contrast Figure (13) 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,

 $\frac{\widehat{RP}-RP}{RP}$ , similar to a percent error calculation. Where RP represents the true value of the 601 three-parameter RP, and  $\widehat{RP}$  refers to the metamodel estimate. 602

Figure (13, top-right) combines the components of bias to show the overall mapping of 603 RPs under BH inference in the high contrast simulation setting. Unlike high contrast RP 604 inference under the Schaefer model, the BH model does shows bias in both RPs here. Despite 605 the bias in  $\frac{B^*}{\overline{B}(0)}$  and  $\frac{F^*}{M}$  these results are similar to that of the Schaefer model in that the 606 overall mapping of RPs is very nearly a minimal distance mapping onto the constrained 607 set of RPs. The primary difference between Schaefer model and BH RP inference is the 608 geometry of their limited RP spaces. Unlike the Schaefer model the BH set encourages bias 609 in both RPs for misspecified models even in very well informed setting. 610

Low Contrast Figure (14) shows the 611 mapping of RPs in the low contrast simulation setting. Figures (14) and (13, top-613 right) share a common scale for the inten-614 sity of color to facilitate comparison. In Fig-615 ure (14) notice that the mildly misspecified 616 area around the BH set produces mappings 617 onto the BH set which resemble the minimal 618 distance mapping seen in the high contrast 619 setting. The primary difference in this low 620 contrast setting, is the break point around 621  $\frac{B^*}{\overline{B}(0)} = 0.4$  above which  $\frac{F^*}{M}$  is sharply under-622 estimated. 623

The region of RPs where the BH model 624 manages to recover the minimal distance 625 mapping may be considered a "safe regime" 626

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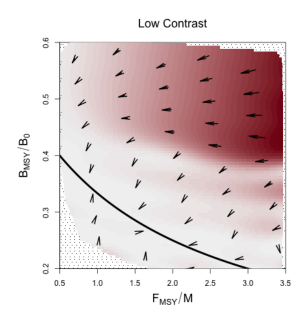


Figure 14: Joint bias direction of RP inference in the low contrast simulation setting. The intensity of color represents the excess bias relative to the shortest possible mapping.

of data types that are reasonably well modeled by a BH model. By comparison of Figure (14), with Figure (12), this safe regime of the BH model occurs for data generated for 628 Cushing-like or BH-like production. While bias of the RPs can still become concerningly 629

large, this region can be considered safe in the sense that even for low contrast data RP 630 estimation under the the BH model recovers the minimal distance mapping. 631

Outside of this safe regime, RP estima-632 tion breaks from the minimal distance map-633 ping at the interface between BH-Like and 634 Ricker-Like regimes of the Schnute model 635 (again see Figure (12)). The Ricker model 636 lies along this regime interface, and repre-637 sents the first model to approach the x-axis 638 for large biomasses as  $\gamma$  increases. markedly unBH-like productivity in the low 640 information simulation setting breaks MLE 641 inference from the minimal distance map-642 ping and instead maps RPs to extremely low 643 values of  $F^*$ ; consequently  $\frac{B^*}{\overline{B}(0)}$  is estimated 644 near the limiting value under the BH (i.e.

645

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

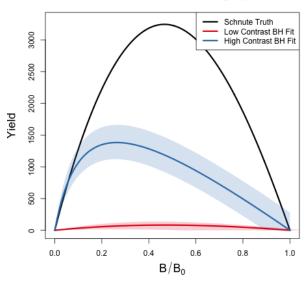


Figure 15: Yield curves for data generated with  $\frac{F^*}{M} = 3.48$  and  $\frac{B^*}{\bar{B}(0)} = 0.48$ .

 $\lim_{F^*\to 0} \frac{1}{F^*/M+2} = 0.5$ ). Similarly the set of Ricker RPs (as well as the Schaeffer set) include 646 this trivial limiting point in common  $(\frac{F^*}{M} = 0, \frac{B^*}{B(0)} = 0.5)$ . 647

Interestingly, in the high contrast setting this trivial mapping for highly misspecified BH 648 models is not present. This suggests that, under a misspecified BH model, the presence of 649 adequate information in the data to produce reasonable estimates of  $\frac{F^*}{M}$ , drives  $\frac{B^*}{\bar{B}(0)}$  below 0.5 650 in accordance with  $\frac{B^*}{\overline{B}(0)} = \frac{1}{F^*/M+2}$ , even when the true  $\frac{B^*}{\overline{B}(0)} > 0.5$ . This phenomena balances 651 RP estimation within the constrained BH set as mediated by the information content of the 652 data and the degree of model misspecification. When the information content in the data 653 is too small to drive a compromised RP estimate, inference completely disregards accurate 654 estimation of  $F^*$  in order to better estimate  $\frac{B^*}{\overline{B}(0)}$  by exploiting the common limiting behavior 655 of the BH set and that of Ricker-like and Logistic-like models. 656

## 4 Discussion

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Results presented here generally agree with what is known about estimating growth rate 658 parameters (Lee et al., 2012; Conn et al., 2010; Magnusson & Hilborn, 2007). These study's 659 appreciate the role of contrast for estimating growth rates, however they struggle to make 660 generally extensible conclusions since they focus only on a handful of stocks that fall short 661 of forming a random sample of the greater population of possible stock behaviors. The LHS 662 design methods presented here are designed specifically to simulate a representative sample 663 of stocks broadly across the space of possible RPs. Furthermore, the simulation design, taken 664 together with the GP metamodel of productivity parmater estimates, allows this study to 665 control the degree of model misspecification and generalize conclusions about the behavior 666 of productivity estimation within the production model setting presented. 667

In the presence of contrast,  $F^*$  estimation can enjoy very low bias even for a wide range of 668 poorly specified models; conversely in the absence of contrast  $F^*$  estimation can suffer very 669 large bias even for slightly misspecified models. This pattern is particularly true for inference 670 under the Schaefer model where the geometry of the restricted RP set isolates estimation 671 failure of  $F^*$  from  $\frac{B^*}{\overline{B}(0)}$ . While contrast has a similar impact on  $F^*$  estimation under the 672 BH model, the geometry of the BH RP set correlates estimation bias of  $F^*$  and  $\frac{B^*}{B(0)}$ . The 673 GP metamodeling approach reveals a more general pattern that highly informative data sets 674 (high contrast) produces a nearly minimal distance mapping of RPs onto the constrained RP set. 676

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 (2.5) captures a usefully broad 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 of discovering more informative fishing behavior, could improve this analysis. In the absence
of a maximally informative dataset simulation methods will not fully describe how inference
fails, but the methods presented here tell the most complete picture yet, with explicit control
of the degree model misspecification, contrast, and a simulation design that allows for uniform
representative data generation across biologically meaningful stocks. The results presented
here suggest the conjecture that under a maximally informative dataset, RP inference with
a two parameter production function will be biased in the direction a shortest distance map
from the true RPs onto restricted set of RPs under the two-parameter model.

Given the potential for model misspecification of RPs, a minimal distance mapping of 694 RPs represents a best-case scenario where the total bias of RPs, when measured jointly, is 695 minimized. That said, without recognizing the geometry of how two-parameter models of 696 productivity limit RP space this may lead to unintuitive implications in RP estimation. For 697 example, due to the shape of the BH RP set a minimal distance mapping ensures that if 698 there is bias in one of  $\frac{B^*}{B_0}$  or  $F^*$ , there will necessarily be bias in the other RP. However under 699 the Schaefer model, since the RP set is a constant in  $\frac{B^*}{B_0}$ , bias in  $F^*$  is not adulterated in the 700 same way by bias in  $\frac{B^*}{B_0}$  estimation. While models with constant RPs, such as the logistic 701 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 702 for developing intuition precisely because they isolate RP estimation in their free RPs from 703 the correlated RP biases present in models like the BH or Ricker model. 704

When one considers the implications of RP bias, overestimation of RPs carries the severe 705 implication of management recommendations potentially leading to overfishing, while un-706 derestimation of RP leads to overly conservative management. In this sense, when the true 707 model is not known, the geometry of the BH set together with the metamodeled bias trends 708 makes the BH model a naturally conservative estimator of RPs for most stocks. For most 709 non-BH populations the BH model is likely to make conservative errors in its estimates of 710  $F^*$  and  $\frac{B^*}{B_0}$ . The one notable exception to the conservatism of the BH model stands for data 711 generated in the Cushing-like regime of Schnute RPs. In this regime the BH model tends 712 to be fairly unbiased overall, however the bias that is present for these populations tends to 713 be overestimation in both RPs, leading to much more severe management consequences for 714 those populations.

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-722 tivity, we should not solely accept conservativism as a rational of choosing a BH model of 723 productivity. Increasing the flexibility of the production function by moving toward three-724 parameter models would release the underlying structural limitations (Mangel et al., 2013) 725 that cause these RP biases in the first place. Punt and Cope (2019) considers a suite of pos-726 sible three-parameter curves which could be used instead of current two-parameter curves. 727 For all of their benefits, three parameter production functions have their own complicating 728 factors, and the structure present in the Schnute model explored here makes it an intuitive 729 bridge model for developing three-parameter models going forward. 730

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

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- summary of  $\sigma$  over RP space comparing between models (PT, Schnute, Schnute DD) to show areas of model breakdown.
  - miss-identifying signal for noise.

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- 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.

# Space of BH Reference Points

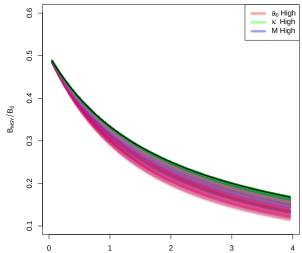


Figure 16: BH RP-space sensitivity to the parmaters M,  $\kappa$ , and  $a_0$ . The black curve shows the BH set in the simple production model setting.

- 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>746</sup> 5 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  $\gamma$ .

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

$$\gamma = \frac{W(\zeta \log(\zeta))}{\log(\zeta)}$$
(34)

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. (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. (12) is to be interpreted as,

$$\gamma = \begin{cases}
\frac{W_0(\zeta \log(\zeta))}{\log(\zeta)} & \zeta \in (0, \frac{1}{e}) \\
\frac{W_{-1}(\zeta \log(\zeta))}{\log(\zeta)} & \zeta \in (\frac{1}{e}, 1)
\end{cases}$$
(35)

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

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