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Data-moderate assessment approaches for Southwest Pacific Ocean (SWPO) striped marlin

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1 Introduction

Assessment of Southwest Pacific Ocean (SWPO) striped marlin (*Kajikia audax*) has been challenging. As documented in the joint NOAA-SPC modeling meeting report (Ducharme-Barth et al., 2025), key issues identified by WCPFC SC20 included poor fits to size composition data and relative abundance indices, conflicts between different data sources, and difficulties in estimating model initial conditions.

Discussions at the 2025 SPC Pre-Assessment Workshop also highlighted challenges related to the estimation of total population scale, where the low biomass scale indicated by previous versions of the assessment appears largely driven by the size composition from multiple fisheries. While estimation of low biomass scale is not inherently problematic, the resulting high ratio of fishing mortality (F) to natural mortality (M) that is maintained over multiple decades in the face of relative stability in catches and catch-per-unit-effort (CPUE) is suspicious and potentially indicative of additional model mis-specification.

The complexity of integrated age-structured models, while providing detailed population dynamics representation, can become problematic when fundamental data conflicts exist or when key biological parameters are uncertain. For SWPO striped marlin, these challenges are compounded by spatiotemporal heterogeneity in fleet coverage, potential non-representativeness in mixed-fleet composition data, and limitations in age data from opportunistic sampling. In the current case, changes to productivity (growth, natural mortality, maturity, and/or steepness) or selectivity assumptions will impact biomass scaling through predictions of expected size composition data.

Scaling back model complexity to a *data-moderate* approach, like Bayesian surplus production models (BSPMs), offers analysts a simplified yet robust alternative for stock assessment when data limitations or conflicts present challenges for more complex models. These models have proven effective in recent pelagic fish stock assessments (Winker et al., 2018), and are routinely applied for WCPFC shark assessments (ISC, 2024; Neubauer et al., 2019, 2024). They also facilitate a more tractable exploration of model assumptions by distilling productivity and fishing assumptions into a restricted parameter subset. The BSPM framework allows for explicit incorporation of parameter uncertainty through informative priors while maintaining computational efficiency and interpretability.

One of the advantages of the Bayesian approach is the through the use of prior information to propagate uncertainty in model assumptions, and biological simulation approaches can provide a robust methods for parameterizing key population dynamics parameters (ISC, 2024; Pardo et al., 2016). For species with complex life histories like striped marlin, these simulation-based priors can incorporate uncertainty in growth, maturity, natural mortality, and reproductive parameters to derive realistic distributions for maximum intrinsic rate of increase and carrying capacity. Additionally, prior pushforward checks can be used to further refine parameter distributions by identifying parameter combinations that produce clearly improbable outcomes.

This analysis presents a data-moderate approach for SWPO striped marlin that addresses the key technical challenges identified in previous assessments while providing a robust framework for stock status evaluation. The model incorporates biological uncertainty through simulation-based priors and includes approaches to address catch uncertainty and population depletion.

2 Methods

2.1 Model Framework

A series of Bayesian state-space surplus production models (BSPMs) spanning the period 1952-2022 were developed following the Fletcher-Schaefer production model framework (Edwards, 2024; Winker et al., 2020). Development of the BSPM followed the approach of (ISC, 2024; Neubauer et al., 2019) and incorporated recent best practices for surplus production models (Kokkalis et al., 2024) and Bayesian workflows for stock assessment (Monnahan, 2024) as guides for model development, analysis, and presentation.

The models were implemented in the Stan probabilistic programming language (Team, 2024b) to take advantage of enhanced convergence diagnostics, greater efficiency in posterior sampling through the no-U-turn (NUTS) Hamiltonian Monte Carlo algorithm (Betancourt & Girolami, 2013), and greater flexibility with model configuration and prior specification. Implementation in R using the *rstan* package (Team, 2024a) provides connection to an ecosystem of additional R packages for model diagnostics and validation: *bayesplot* (Gabry & Mahr, 2024) for posterior visualization and *loo* (Vehtari et al., 2024).

The model begins from an assumed unfished state in 1952 and incorporates process error in population dynamics, observation error in abundance indices, and uncertainty in biological parameters. Two model variants were developed: a baseline model and a depletion-constrained model that incorporates additional information about historical stock status from mean size data.

2.1.1 Input data

The BSPM was fit to catch and standardized CPUE data for SWPO striped marlin. Annual catch data (in numbers) spanning 1952-2022 were compiled from the 2024 assessment (Castillo-Jordan et al., 2024) sources and aggregated into total removals by year.

The catch series (Figure 1) shows initial low removals in 1952-1953, followed by a high peak in 1954. Overall, however, catches have been relatively stable though catches in recent decades show a slight decline. A single standardized CPUE index (1988-2022) from the diagnostic case of the 2024 assessment (Castillo-Jordan et al., 2024) was used. The index is very noisy (Figure 2). In general, however, there is a slight declining trend over the index period. The decline was most pronounced after 2000, though the trend stabilized somewhat, prior to showing a slight increase in recent years.

2.1.2 Population Dynamics

The population dynamics follow a Fletcher-Schaefer surplus production model with state-space formulation where population depletion x_t (relative to carrying capacity K) evolves according to:

$$x_{t} = \begin{cases} (x_{t-1} + R_{max}x_{t-1}(1 - \frac{x_{t-1}}{h}) - C_{t-1}) \times \epsilon_{t}, & x_{t-1} \leq D_{MSY} \\ (x_{t-1} + x_{t-1}(\gamma \times m)(1 - x_{t-1}^{n-1}) - C_{t-1}) \times \epsilon_{t}, & x_{t-1} > D_{MSY} \end{cases}$$

where R_{max} is the maximum intrinsic rate of increase, n is the shape parameter, and ϵ_t represents multiplicative process error with $\epsilon_t = \exp(\delta_t - \sigma_P^2/2)$ and $\delta_t \sim N(0, \sigma_P)$. The intermediate parameters are: $D_{MSY} = (1/n)^{1/(n-1)}$, $h = 2D_{MSY}$, $m = R_{max}h/4$, and $\gamma = n^{n/(n-1)}/(n-1)$.

2.1.3 Observation Model

The model fits to a standardized CPUE index using a lognormal likelihood:

$$I_t \sim \text{Lognormal}(\log(q \times x_t) - \frac{\sigma_{O,t}^2}{2}, \sigma_{O,t})$$

where catchability q is analytically derived assuming an uninformative uniform prior, and total observation error $\sigma_{O,t} = \sigma_{O,input} + \sigma_{O,add}$ combines fixed input uncertainty with an estimated additional error component.

2.1.4 Catch Treatment

Annual fishing mortality is estimated directly $F_t \sim N^+(0, \sigma_F)$ with catch fitted using lognormal likelihood with uncertainty $\sigma_C = 0.2$

Catch is predicted as:

$$C_t = (x_t + \text{surplus production}_t) \times (1 - \exp(-F_t)) \times \epsilon_t \times K$$

2.2 Prior Development

2.2.1 Biological Simulation Framework

Informative priors for key population parameters were developed through biological simulation using Monte Carlo sampling from biologically plausible parameter distributions. The simulation framework incorporated uncertainty and correlations in life history parameters:

- Growth parameters: Francis parameterization with independent sampling
- Natural mortality: Reference mortality M_{ref} negatively correlated with maximum age (r = -0.3)
- Maturity: Length-based logistic function parameters
- Weight-length relationship: Correlated allometric parameters (r = -0.5) with biological constraints
- Fishery selectivity: Length at first capture for depletion prior

Parameter combinations (Table 1) were filtered to ensure biological realism, with weight-length relationships constrained to produce 150-350 kg individuals at 300 cm length. Correlations between life history parameters reflect biological trade-offs, with faster-growing species typically having higher natural mortality rates and shorter lifespans.

2.2.2 Maximum Intrinsic Rate of Increase (R_{max}) Prior

 R_{max} was calculated by numerically solving the Euler-Lotka equation using bounded optimization (nlminb function in R) to find the value of R_{max} that satisfies the equation within convergence criteria. Starting values and bounds were set to ensure biologically realistic solutions. Formulation of the Euler-Lotka equation followed the approach implemented in openMSE (Hordyk et al., 2024; Stanley et al., 2009):

$$\alpha \sum_{a=1}^{A_{max}} l_a f_a \exp(-R_{max} \times a) = 1$$

where α is the slope at the origin of the stock-recruitment curve, l_a is the probability of survival to age a, and f_a is the fecundity (reproductive output) at age a. Additional technical detail can be found in Section 7.1 of the Technical Annex.

Repeating this calculation across each of the 500,000 parameter combinations resulted in a prior distribution of R_{max} conditioned on plausible biology and life-history characteristics. The resulting distribution was first filtered to values of $R_{max} < 1.5$ representative of what the literature (Gravel et al., 2024; Hutchings et al., 2012) indicates is most likely for teleosts. A prior pushforward analysis, similar to (ISC, 2024), was used to further refine this distribution and develop a lognormal prior for R_{max} . Briefly, random values of R_{max} along with random values of the other leading parameters of the BSPM (drawn from the prior distributions described in Table 2) were used to drive simulated populations forward subject to direct removal of the observed catch (Figure 1). Based on the results of the prior pushforward (Figure 3), the R_{max} distribution was further filtered based on the following criteria:

- Depletion in the terminal year > 2% and $R_{max} < 1$
- Depletion in the terminal year > 2%, $R_{max} < 1$, and trend in depletion over the last decade between -5% and 10%

This last filtering step approximates the recent relative trend seen in the index (Figure 2). Fitting a lognormal distribution to the remaining R_{max} values results in the prior distribution specified in Table 2 and shown in Figure 4. Note that though initial filtering restricted $R_{max} < 1$ the final lognormal prior still assigns some prior weight to $R_{max} > 1$.

2.2.3 Depletion Prior

It is not possible to fit to size composition data in a BSPM framework, however information on fishing mortality and depletion may be contained in the long time series of New Zealand recreational fishing data. This fishery routinely catches the largest individuals and shows a decline in average size since the start of the model period [FIGURE: NZ weight data and depletion estimation], prior to which limited large-scale commercial fishing occurred. A supplementary analysis estimated initial population depletion in 1988 using New Zealand recreational fishing weight data. A modified Gedamke-Hoenig approach (Gedamke & Hoenig, 2006) for using size composition data to estimate mortality in non-equilibrium situations was applied. The method assumed a two-period mortality model where total mortality increased from Z_1 (initial period) to Z_2 (final period) at a known change point (1952) when industrial fishing commenced at the start of the model period (more detail in Technical Annex Section 7.2). For each successful parameter combination, relative spawning stock biomass depletion was calculated as the ratio of spawning potential under the two mortality regimes:

$$\text{Relative Depletion} = \frac{SPR_{Z_2}}{SPR_{Z_1}} = \frac{\sum_{a=1}^{A_{max}} l_{a,Z_2} f_a}{\sum_{a=1}^{A_{max}} l_{a,Z_1} f_a}$$

where $l_{a,Z}$ is survival to age a under total mortality Z, and f_a is reproductive output at age a (incorporating maturity, weight, sex ratio, and reproductive cycle). Additional technical detail on the population dynamics components of this equation can be found in Section 7.1 of the Technical Annex.

The resulting depletion estimates across successful parameter combinations were used to construct an

informative lognormal prior for stock depletion that could be applied in the BSPM in 1988.

2.3 Model Implementation

2.3.1 Computational Setup

Models were implemented using the No-U-Turn Sampler (NUTS) in Stan with the following configuration: - 5 chains with random starting points - 3,000 iterations per chain (1,000 warmup) - Thinning: every 10th sample retained - Adaptation parameters: adapt_delta = 0.99, max_treedepth = 15 - Final sample size: 1,000 posterior samples

2.3.2 Model Configurations

Two primary model configurations were developed:

- 1. Base BSPM (bspm estF.stan): Estimated fishing mortality without depletion prior
- 2. Depletion-constrained BSPM (bspm_estF_softdep.stan): Includes 1988 depletion prior

Both models used the same prior parameterizations (Table 2) based on the biological simulation filtering.

2.3.3 Model Diagnostics

Model performance was assessed using: - Convergence diagnostics: $\hat{R} < 1.01$, effective sample size > 500 -Posterior predictive checks for index and catch fits - Parameter identifiability through posterior examination

[TABLE: model convergence diagnostics]

2.4 Uncertainty Characterization

The analysis explicitly incorporated multiple sources of uncertainty:

- 1. Biological uncertainty: Through comprehensive life history parameter simulation
- 2. Catch uncertainty: Via lognormal catch likelihood with $\sigma_C = 0.2$
- 3. Observation uncertainty: Through time-varying index uncertainty plus additional error
- 4. Process uncertainty: Through estimated recruitment/population variability

Results are presented as posterior distributions reflecting the full uncertainty structure, with management quantities calculated from the joint posterior distribution of all model parameters.

[FIGURE: posterior parameter distributions and correlations]

3 Declaration of Generative AI use

A generative artificial intelligence (AI) assistant, Anthropic Claude Sonnet 4.0, was used to parse model code in the preparation of an initial draft of this report.

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5 Tables

Table 1: Biological parameter distributions and sources used in simulation framework

Parameter	Symbol	Distribution	Mean/Mode	CV/Range	Correlation	Source/Notes
Length at age 1	L_1	Lognormal	60 cm	0.2	Independent	(Ducharme-Barth et al., 2025)
Length at age 10	L_2	Lognormal	210 cm	0.2	Independent	(Ducharme-Barth et al., 2025)
Growth coefficient	k	Beta(6.5, 3.5)	~0.65	-	Independent	(Ducharme-Barth et al., 2025)
Length CV	cv_{len}	Uniform	_	[0.05, 0.25]	-	Growth variability
Maximum age	A_{max}	Lognormal	15 years	0.2	$r = -0.3$ with M_{ref}	(Farley et al., 2021)
Reference mortality	M_{ref}	Lognormal	$0.36 \ yr^{-1}$	0.44	$r = -0.3$ with A_{max}	(Hamel & Cope, 2022) $(5.4/A_{max})$
Maturity slope	a_{mat}	Normal	-20	0.2	-	Logistic steepness
Length at 50% maturity	L_{50}	Lognormal	184 cm	0.2	-	(Farley et al., 2021)
Weight coefficient	a_w	Lognormal	5.4×10^{-7}	0.05	$r = -0.5$ with b_w	(Castillo-Jordan et al., 2024)
Weight exponent	b_w	Normal	3.58	0.05	$r = -0.5$ with a_w	(Castillo-Jordan et al., 2024) Constrained: 150-350 kg at 300 cm
Steepness	h	Beta(3, 1.5)	~0.73	-	-	Censored to $[0.2, 1.0]$
Sex ratio (prop. female)	s	Normal	0.5	0.05	-	Constrained [0.01, 0.99]
Reproductive cycle	-	Fixed	1 year	-	-	Annual spawning
Length at first capture	L_c	Lognormal	$140~\mathrm{cm}$	0.125		Fishery selectivity

Parameter	Symbol	Distribution	Mean/Mode	CV/Range	Correlation	Source/Notes
Reference ages	age_1, age_2	Fixed	0, 10 years	-	-	Francis
						parameterization

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Table 2: Prior distributions for Bayesian surplus production model parameters

Parameter	Distribution	Mean	SD	Description			
Core Population Parameters							
K	Lognormal	$\log(1,049,036)$	0.46	Carrying capacity			
R_{max}	Lognormal	$\log(0.5099)$	0.46	Maximum intrinsic rate of increase (from biological simulation)			
n	Lognormal	$\log(2)$	0.1	Shape parameter (Pella-Tomlinson production function); $n=2$ is a Schaefer model			
Process and Observation Error							
σ_P	Lognormal	$\log(0.0533)$	0.27	Process error standard deviation (Winker et al., 2018)			
$\sigma_{O,add}$	Half-Normal	0	0.2	Additional observation error			
σ_F	Half-Normal	0	0.025	Fishing mortality variability			
Depletion Constraint							
(Constrained Model Only)							
x_{1988}	Lognormal	$\log(0.6374)$	0.2	Initial depletion at $t = 37$ (this analysis)			

6 Figures

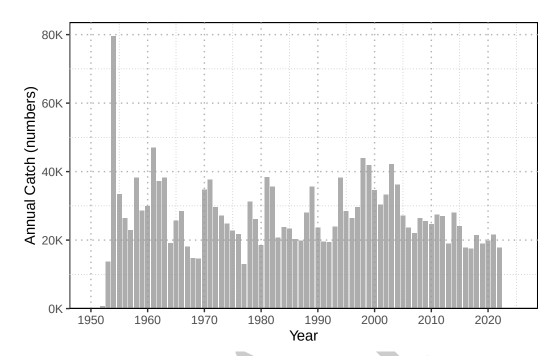


Figure 1: Annual catch of Southwest Pacific Ocean striped marlin (1952-2022). Catch data shows initial low removals in early years, a peak in 1954, followed by relatively stable catches with a slight decline in recent decades.

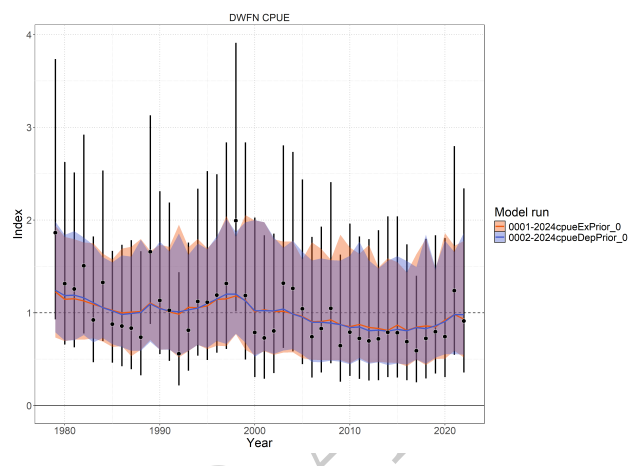


Figure 2: Standardized index (black points), observation error (black bars), and posterior predicted model fits (colored lines) with associated 95% credible interval (colored shading).

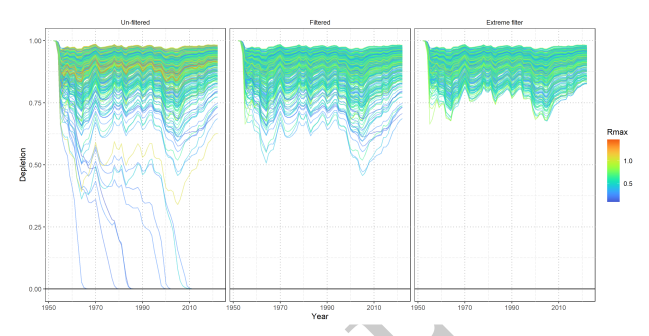


Figure 3: Prior pushforward check for population depletion trajectories under different biological parameter filtering scenarios. Each line represents a simulated population trajectory colored by maximum intrinsic rate of increase (R_{max}) . Left panel shows unfiltered R_{max} , middle panel shows filtered R_{max} , and right panel shows extreme filtering.

7 Technical Annex

Additional technical detail of population dynamics components used to solve the Euler-Lotka equation for R_{max} and implement the Gedamke-Hoenig approach (Gedamke & Hoenig, 2006) for using size composition data to estimate mortality in non-equilibrium situations.

7.1 Equilibrium Age-Structured Population Dynamics

7.1.1 Survival

Survival schedule was calculated assuming constant natural mortality:

$$l_a = \begin{cases} 1 & \text{if } a = 1 \\ l_{a-1} \times \exp(-M_{ref}) & \text{if } 1 < a < A_{max} \\ \frac{l_{A_{max}-1}}{1 - \exp(-M_{ref})} & \text{if } a = A_{max} \text{ (plus group)} \end{cases}$$

7.1.2 Fecundity

Reproductive output incorporated biological constraints:

$$f_a = \frac{\psi_a \times W_a \times s}{\rho}$$

where: $-\psi_a$ = proportion mature at age a - W_a = weight at age a (proxy for reproductive capacity) - s = sex ratio (proportion female) - ρ = reproductive cycle (years between spawning events)

Maturity-at-age was derived from length-based logistic maturity:

$$\psi_L = \frac{\exp(a_{mat} + b_{mat} \times L)}{1 + \exp(a_{mat} + b_{mat} \times L)}$$

where $b_{mat} = -a_{mat}/L_{50}$, then integrated over the length-at-age distribution to obtain ψ_a .

Length-at-age followed the Francis parameterization:

$$L_a = L_1 + (L_2 - L_1) \times \frac{1 - \exp(-k(a - age_1))}{1 - \exp(-k(age_2 - age_1))}$$

with length variability modeled using probability density functions linking length bins to ages.

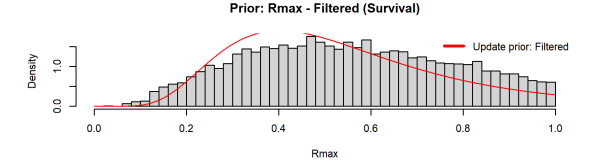
Weight-at-age used the allometric relationship:

$$W_a = a_w \times L_a^{b_w}$$

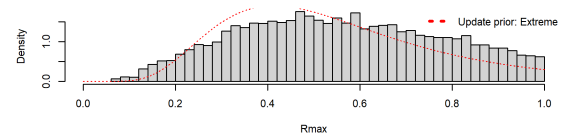
7.1.3 Stock-Recruitment Relationship

The slope at the origin of the stock-recruitment curve was:

$$\alpha = \frac{4h}{(1-h)\phi_0}$$







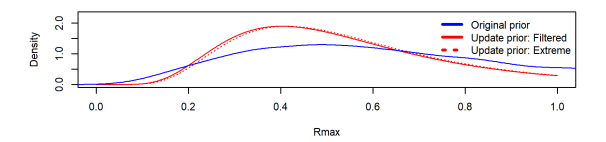


Figure 4: Prior distributions for maximum intrinsic rate of population increase R_{max} . Upper panel: Gray histogram is the R_{max} values from the numerical simulation which meet baseline filtering levels. Red line is fitted lognormal distribution. Middle panel: Gray histogram is the R_{max} values from the numerical simulation which meet extreme filtering levels. Dotted red line is fitted lognormal distribution. Bottom panel: Original distribution of R_{max} values from numerical simulation (gray), those from viable populations (blue), and the two lognormal priors (red).

The calculation incorporated steepness (h) through the Beverton-Holt stock-recruitment relationship. Spawners-per-recruit in the unfished state was:

$$\phi_0 = \sum_{a=1}^{A_{max}} l_a f_a$$

This α parameter links the intrinsic rate of increase to recruitment compensation, ensuring that R_{max} estimates reflect realistic population productivity under the assumed stock-recruitment dynamics.

Successful simulations (those yielding $R_{max} > 0$ and $R_{max} < 1.5$) were filtered to create a plausible prior distribution [FIGURE: pushforward]. The resulting distribution was fitted to a lognormal prior for use in the BSPM [FIGURE: Rmax prior].

7.2 Transitional Mean Weight Model

For each biological parameter combination, total mortality rates Z_1 and Z_2 were estimated by fitting a transitional mean weight model to observed temporal changes in recreational catch weights. The expected mean weight during the transition follows:

$$\bar{W}(d) = W_{\infty} - \frac{Z_1 Z_2 (W_{\infty} - W_c) [Z_1 + k + (Z_2 - Z_1) e^{-(Z_2 + k)d}]}{(Z_1 + k) (Z_2 + k) [Z_1 + (Z_2 - Z_1) e^{-Z_2 d}]}$$

where: - d = time since mortality change (years after 1952) - W_{∞} = asymptotic weight from growth parameters - W_c = weight at first capture (corresponding to L_c) - k = von Bertalanffy growth coefficient

The mortality parameters were estimated in a likelihood framework by maximizing the likelihood of the observed mean weights given a time series of predicted mean weights $\bar{W}(d)$. Optimization was constrained such that $Z_1, Z_2 \geq M_{ref}$ to ensure biologically realistic mortality estimates.