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### SEDAR87-RD-15

November 2024



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# ICES Journal of Marine Science

ICES Journal of Marine Science (2017), 74(5), 1311-1321. doi:10.1093/icesjms/fsw193

## **Original Article**

## Comparing estimates of abundance trends and distribution shifts using single- and multispecies models of fishes and biogenic habitat

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Thorson, J. T., and Barnett, L. A. K. Comparing estimates of abundance trends and distribution shifts using single- and multispecies models of fishes and biogenic habitat. – ICES Journal of Marine Science, 74: 1311–1321.

Received 30 August 2016; revised 7 October 2016; accepted 10 October 2016; advance access publication 14 January 2017.

Several approaches have been developed over the last decade to simultaneously estimate distribution or density for multiple species (e.g. "joint species distribution" or "multispecies occupancy" models). However, there has been little research comparing estimates of abundance trends or distribution shifts from these multispecies models with similar single-species estimates. We seek to determine whether a model including correlations among species (and particularly species that may affect habitat quality, termed "biogenic habitat") improves predictive performance or decreases standard errors for estimates of total biomass and distribution shift relative to similar single-species models. To accomplish this objective, we apply a vector-autoregressive spatio-temporal (VAST) model that simultaneously estimates spatio-temporal variation in density for multiple species, and present an application of this model using data for eight US Pacific Coast rockfishes (Sebastes spp.), thornyheads (Sebastolobus spp.), and structure-forming invertebrates (SFIs). We identified three fish groups having similar spatial distribution (northern Sebastes, coastwide Sebastes, and Sebastolobus species), and estimated differences among groups in their association with SFI. The multispecies model was more parsimonious and had better predictive performance than fitting a single-species model to each taxon individually, and estimated fine-scale variation in density even for species with relatively few encounters (which the single-species model was unable to do). However, the single-species models showed similar abundance trends and distribution shifts to those of the multispecies model, with slightly smaller standard errors. Therefore, we conclude that spatial variation in density (and annual variation in these patterns) is correlated among fishes and SFI, with congeneric fishes more correlated than species from different genera. However, explicitly modelling correlations among fishes and biogenic habitat does not seem to improve precision for estimates of abundance trends or distribution shifts for these fishes.

**Keywords:** fish distribution shift, index standardization, joint dynamic species distribution model (JDSDM), Pacific rockfish, spatio-temporal model, structure-forming invertebrates.

#### Introduction

There are several benefits to simultaneously analysing the distribution and density of multiple species within a natural community. Multispecies models of spatial distribution can estimate associations among species (Latimer *et al.*, 2009; Ovaskainen *et al.*, 2016; Thorson *et al.*, 2015a, 2016a), such that the presence

or absence of a given species can be used as an indicator of habitat for other species when reliable habitat variables are otherwise lacking (Ovaskainen *et al.*, 2010). Multispecies models fitted to presence/absence data (termed "multispecies occupancy models") can also be used in some cases to identify the impact of management actions more efficiently than using single-species occupancy

Published by International Council for the Exploration of the Sea 2017. This work is written by US Government employees and is in the public domain in the US. International Council for the Exploration of the Sea models (Zipkin *et al.*, 2010). Furthermore, research shows that estimating the distribution for each species individually and then summarizing community-level properties by stacking results from single-species analyses can result in improper inference about ecological communities (Clark *et al.*, 2014).

The predictive performance of species distribution models is often improved when available covariates are included that are informative about habitat quality. Unfortunately, environmental variables associated with habitat quality are difficult to measure for many species, including demersal marine fishes. To overcome this difficulty, new species distribution modelling (SDM) techniques may allow differences in habitat to be inferred from spatial variation in the density of species with similar habitat requirements (Latimer et al., 2009; Ovaskainen et al., 2010). For example, joint species distribution models have previously been used to show strong covariation in population density among US Pacific Coast rockfishes and thornyheads (Sebastes and Sebastolobus spp.), and these correlations imply that the population density of one species is informative about the density of correlated species (Thorson et al., 2015a). Similarly, joint dynamic species distribution models (JDSDMs) can estimate abundance trends for infrequently encountered species and have revealed similarities in spatio-temporal dynamics among related butterfly species (Thorson et al., 2016a,b). However, JDSDMs have not previously been used to explore associations between fishes and species that are associated with specific habitat features (e.g. structure-forming invertebrates, SFI).

Marine fishes are intensively managed in many parts of the developed world, and the management of marine fisheries is strongly linked to estimates of population status and productivity from population models (termed "stock assessment models") throughout North America and Europe (Methot, 2009; Maunder and Punt, 2013). Although these stock assessment models often integrate many different types of information, time-series that are proportional to population abundance ("abundance indices") are often among the most critical (Francis, 2011). For this reason, there is considerable research regarding best practices for minimizing error when estimating abundance indices for fishes from survey data (Walters, 2003; Maunder and Punt, 2004; Shelton et al., 2014). Similarly, survey data are increasingly used to estimate shifts in fish distribution over time (e.g. due to climate change), and distribution shifts are often measured by estimating the centroid of the population's distribution and shifts in this centroid over time (Perry et al., 2005; Pinsky et al., 2013). Research suggests that spatio-temporal models are statistically efficient and can improve precision for estimates of abundance indices or distribution shifts relative to nonspatial models, given limited available data (Thorson et al., 2015b, 2016b). Recently, novel methods have been proposed for estimating abundance indices by simultaneously fitting a JDSDM to data for multiple species (Thorson et al., 2016). However, there is little research comparing the single- and multispecies approaches to estimating abundance indices for marine fishes.

For three reasons, Pacific rockfishes and their close relatives provide an interesting example when studying associations between fishes and species that affect habitat suitability ("biogenic habitat") or the potential benefit of these associations when estimating abundance indices or distribution shifts. Most importantly, Pacific rockfishes manifest an astounding diversity of species, with more than 65 species co-occurring in the Northeast Pacific (Hyde and Vetter, 2007) and exhibit a wide range of life history strategies (Love et al., 2002; Mangel et al., 2007). Given this life history diversity, rockfishes likely include species whose spatial distributions are both strongly correlated and relatively uncorrelated with SFI. Second, Pacific rockfishes differ in functional traits related to the feeding type and efficiency [eye and gill raker size, Ingram and Shurin, 2009], so species with similar spatial distribution and feeding types might exhibit correlated changes in productivity over time in response to variable food supply. Therefore, bottom-up drivers of abundance or distribution changes would result in correlated abundance or distribution changes over time for species with similar feeding types. Third, many Pacific rockfishes have low and extremely variable population densities (Thorson et al., 2011), such that single-species estimates of trends in population abundance or population distribution are frequently imprecise (Thorson et al., 2015b, 2016b). Given these characteristics of the rockfish assemblage, the inclusion of information about species associations and biogenic habitat when estimating population abundance may increase precision and thereby improve stock assessments.

Given the potential benefit of estimating habitat quality from the density of co-occurring marine species when estimating abundance indices, we seek to simultaneously estimate the density of Pacific rockfishes and structure-forming invertebrates at a coastwide scale. Specifically, we seek to answer three questions: (i) do Pacific rockfishes have an association with structure-forming invertebrates on the US West Coast? (ii) Is this association similar or variable among rockfish species? and (iii) Does the inclusion of information regarding co-occurrence (either among rockfishes or between rockfish and SFI) improve predictions of local rockfish density or increase precision when estimating rockfish abundance trends or population distribution? To address these questions, we develop a vector-autoregressive spatio-temporal (VAST) model for jointly analysing catch-rate data for fish and structure-forming invertebrates and apply the model to data for eight rockfish species and SFI during 2003-2014.

#### Methods Pacific rockfishes

Pacific rockfishes (genus Sebastes) and thornyheads (genus Sebastolobus), hereafter collectively called "rockfishes", are one of the dominant species groups within the assemblage of bottomassociated fishes off the US West Coast. Pacific rockfishes in this region are monitored by the West Coast groundfish bottom trawl survey (WCGBTS) conducted annually by the Northwest Fisheries Science Center since 2003 (Bradburn et al., 2011). The WCGBTS covers areas between the Canada and Mexico borders in 55-1280 m depth, and survey stations for each year are chosen at random within strata defined by depth and latitude (two regions divided at Point Conception, CA). Four commercial vessels (20–28 m length) are chartered each year to sample from mid-May to late October, conducting ca. 15-min tows at a speed of 2 knots using a standard Aberdeen-type trawl with a 3.8-cm mesh codend liner, 25.9-m headrope, and 31.7-m footrope. All fishes and invertebrates are sorted at sea to the lowest possible taxon, and their wet weight is measured. For the purposes of our analysis, we take the midpoint of each haul to represent the location of each biological sample.

We analyse these survey data between the years 2003 and 2014, focusing on structure-forming invertebrates and eight species of Pacific rockfish (Table 1) that are frequently captured within the survey and for which there was previous documentation of

**Table 1.** List of taxa (common and scientific name), the abbreviations used to indicate taxa in plots, and the total number of encounters during 2003–2014.

Common name	Scientific name	Plotting code	Encounters	
Structure-forming invertebrates (SFIs)	_	SFI	6383	
Longspine thornyhead	Sebastolobus altivelis	L. spine	2758	
Shortspine thornyhead	Sebastolobus alascanus	S. spine	3891	
Darkblotched rockfish	Sebastes crameri	Dark	1338	
Pacific Ocean perch	Sebastes alutus	POP	547	
Sharpchin rockfish	Sebastes zacentrus	Sharp	490	
Splitnose rockfish	Sebastes diploproa	Split	1619	
Stripetail rockfish	Sebastes saxicola	Stripe	1630	
Greenspotted rockfish	Sebastes chlorostictus	Green	434	

association with structure-forming invertebrates at fine spatial scales (Love et al., 2002). We aggregate the structure-forming invertebrate taxa into a single grouping to obtain adequate encounter rates for estimating the distribution for structure-forming invertebrates. This SFI group primarily consists of sponges (phylum Porifera), anemones (order Actiniaria), and sea pens (order Pennatulacea), along with fewer observations of true corals (subclass Hexacorallia) and other soft corals (subclass Octocorallia). Although the survey is primarily designed to capture demersal fishes and is not as effective as visual methods for assessing structure-forming invertebrates, it is the primary source of data for estimating spatio-temporal associations between demersal fishes and biogenic habitat at large spatial and temporal scales off the US West Coast. Bottom-trawl samples have been shown to be a good predictor of biogenic habitat distribution in areas such as the eastern Bering Sea based on validation using camera surveys (Rooper et al., 2016).

#### Vector-autoregressive spatio-temporal (VAST) model

We seek to estimate the association among fishes and structureforming invertebrates and, therefore, model correlations among density d(s, c, t) for each taxon c (indicating fish species or the SFI group) at location s and time t (all symbols are summarized in Table 2). To do so, we build upon recent research regarding JDSDMs. In particular, we propose a VAST model, where the probability distribution for catch data  $b_i$  is decomposed into two components representing (i) the probability of encounter  $p(s_i, c_i, t_i)$  for the location  $s_i$ , taxon  $c_i$ , and year  $t_i$  of the  $i^{\text{th}}$ sample, and (ii) the expected catch rate  $r(s_i, c_i, t_i)$ , given that taxon ci is encountered. Decomposing catch rates into encounter-probability p and positive catch rates r is commonly conducted using delta models (Maunder and Punt, 2004; Martin et al., 2005), although delta models have not previously been used within JDSDMs. Using a delta model allows us to separately identify species with similar distribution (similarities in occupied habitat) vs. similar density (similarities in hotspots within their distribution). Therefore, we specify:

$$\Pr(b_i = B) = \begin{cases} 1 - p(s_i, c_i, t_i) & \text{if } B = 0\\ p(s_i, c_i, t_i) \times \operatorname{Lognormal}\{B|\log[w_i \times r(s_i, c_i, t_i)], \sigma_c^2\} & \text{if } B > 0 \end{cases}$$
(1)

where Lognormal( $x|\mu, \sigma^2$ ) is a lognormal probability distribution function for value x, given a log-mean of  $\mu$  and a variance of  $\sigma^2$ , and  $w_i$  is the area swept for the *i*<sup>th</sup> sample.

Using this delta model, we separately develop a spatiotemporal model for encounter probabilities p and positive catch rates r. We approximate spatio-temporal variation in encounter probability  $p(s_i, c_i, t_i)$  using a logit-linked linear predictor:

$$\operatorname{logit}[p(s_i, c_i, t_i)] = \gamma_p(c_i, t_i) + \varepsilon_p(s_i, c_i, t_i) + \delta_p(c_i, v_i)$$
(2)

where  $\gamma_p(c_i, t_i)$  is an intercept for encounter probability for each taxon *c* and time *t*,  $\varepsilon_p(s_i, c_i, t_i)$  approximates spatio-temporal variation in encounter probability (in logit-space), and  $\delta_p(c_i, v_i)$  is a "vessel effect" for the vessel  $v_i$  conducting the *i*<sup>th</sup> sample when catching taxon  $c_i$ . Vessel effects are included because the WCGBTS is obtained using 3–4 different vessels per year, and previous research indicates that vessels in each year have small, but important, variation in fishing behaviour and resulting catch rates (Helser *et al.*, 2004; Thorson and Ward, 2014). Expected catch rates when a species is encountered  $r(s_i, c_i, t_i)$  are similarly approximated using a log-linked linear predictor:

$$\log[r(s_i, c_i, t_i)] = \gamma_r(c_i, t_i) + \varepsilon_r(s_i, c_i, t_i) + \delta_r(c_i, v_i)$$
(3)

where  $\gamma_r(t_i, c_i)$  is an intercept,  $\varepsilon_r(s_i, c_i, t_i)$  is the spatio-temporal variation, and  $\delta_r(c_i, v_i)$  is a vessel effect for expected catch rates.  $\varepsilon_r(s_i, c_i, t_i)$ ,  $\varepsilon_p(s_i, c_i, t_i)$ ,  $\delta_p(c_i, v_i)$ , and  $\delta_r(c_i, v_i)$  approximate processes that affect density and catchability, respectively, but are not otherwise modelled explicitly (Thorson *et al.*, 2016). Including these effects in the VAST model allows catch data  $b_i$  for nearby locations to be correlated (via correlations in encounter probability *p* or positive catch rate *r*) and also improves density predictions at locations that otherwise have little data (Shelton *et al.*, 2014).

The VAST model involves specifying a probability distribution for spatio-temporal variation ( $\varepsilon_p(s, c, t)$  and  $\varepsilon_r(s, c, t)$ ) and vessel effects ( $\delta_p(c, v)$  and  $\delta_r(c, v)$ ). For each modelled year, we therefore specify a three-dimensional Gaussian process for spatio-temporal variation:

$$\operatorname{vec}[\mathbf{E}_p(t)] \sim \operatorname{MVN}(0, \mathbf{R}_p \otimes \mathbf{V}_{\varepsilon p})$$
 (4)

where  $\mathbf{E}_p(t)$  is a matrix composed of  $\varepsilon_p(s, c, t)$  at every modelled location *s* and taxon *c* in a given year *t*,  $\operatorname{vec}[\mathbf{E}_p(t)]$  is a vector composed of stacking every column of  $\mathbf{E}_p(t)$ ,  $\mathbf{R}_p$  is the correlation matrix approximating similar encounter probability among locations,  $\mathbf{V}_{\varepsilon p}$  is the covariance matrix approximating similar encounter probability among species, and  $\otimes$  is the Kronecker product such that  $\mathbf{R}_p \otimes \mathbf{V}_{\varepsilon p}$  is a covariance matrix between any taxon *c* and location *s* in year *t* [ $\mathbf{E}_r(t)$  follows an identical distribution, but with  $\mathbf{R}_r$  and  $\mathbf{V}_{\varepsilon r}$  in place of  $R_p$  and  $\mathbf{V}_{\varepsilon p}$ ].

Name	Symbol	Dimension	Туре
Sample	i		Index
Taxon	с	_	Index
Location	S	_	Index
Year	t	_	Index
Taxon (species or species-group)	с	_	Index
Vessel	ν	_	Index
Probability distribution for catches	В	_	Random variable
Catch data	bi	ni	Data
Area-swept for each sample	Wi	ni	Data
Area associated with each location	a(s)	ns	Data
Statistic associated with each location	$\mathbf{x}(\mathbf{s})$	n <sub>s</sub>	Data
Variance in positive catch rates	$\sigma^2$	_	Fixed effect
Intercept for p	$\gamma_n(c,t)$	$n_c \times n_t$	Fixed effect
Intercept for r	$\gamma_r(c,t)$	$n_c \times n_t$	Fixed effect
Decorrelation distance in $\varepsilon_n(s, c, t)$	κ <sub>n</sub>	_	Fixed effect
Decorrelation distance in $\varepsilon_r(s, c, t)$	κ <sub>r</sub>	_	Fixed effect
Geometric anisotropy	Н	$2 \times 2$	Fixed effect
Factor approximation to $V_{\varepsilon n}$	$\mathbf{L}_{\mathrm{s}p}$	$n_c \times n_f$	Fixed effect
Factor approximation to $V_{sr}$	$L_{sr}$	$n_c \times n_f$	Fixed effect
Factor approximation to $\mathbf{V}_{\delta n}$	$\mathbf{L}_{\delta p}$	$n_c \times n_f$	Fixed effect
Factor approximation to $V_{\delta r}$	$L_{\delta r}$	$n_c \times n_f$	Fixed effect
Spatio-temporal variation in p	$\varepsilon_p(s,c,t)$	$n_s \times n_c \times n_t$	Random effect
Spatio-temporal variation in r	$\varepsilon_r(s,c,t)$	$n_{s} \times n_{c} \times n_{t}$	Random effect
Vessel effect for p	$\delta_p(c, v)$	$n_c \times n_v$	Random effect
Vessel effect for r	$\delta_r(c, v)$	$n_c \times n_v$	Random effect
Encounter probability	p(s, c, t)	$n_{s} \times n_{c} \times n_{t}$	Derived quantity
Positive catch rates	r(s,c,t)	$n_s \times n_c \times n_t$	Derived quantity
Local density	d(s, c, t)	$n_{s}  imes n_{c}  imes n_{t}$	Derived quantity
Spatio-temporal variation in <i>p</i> in year <i>t</i>	$E_{p}(t)$	$n_s  imes n_c$	Derived quantity
Spatio-temporal variation in <i>r</i> in year <i>t</i>	$\vec{E_r(t)}$	$n_s  imes n_c$	Derived quantity
Spatial correlation in $\varepsilon_p(s, c, t)$	$\mathbf{R}_{p}$	$n_{\rm s}  imes n_{\rm s}$	Derived quantity
Spatial correlation in $\varepsilon_r(s, c, t)$	R <sub>r</sub>	$n_{\rm s}  imes n_{\rm s}$	Derived quantity
Correlation among species in $\varepsilon_p(s, c, t)$	$\mathbf{V}_{\varepsilon p}$	$n_c  imes n_c$	Derived quantity
Correlation among species in $\varepsilon_r(s, c, t)$	$\mathbf{V}_{arepsilon'}$	$n_c  imes n_c$	Derived quantity
Correlation among species in $\delta_p(c, v)$	$\mathbf{V}_{\delta p}$	$n_c  imes n_c$	Derived quantity
Correlation among species in $\delta_r(c, v)$	ν <sub>δr</sub>	$n_c  imes n_c$	Derived quantity
Index of abundance	I(c,t)	$n_c  imes n_t$	Derived quantity
Center of spatial distribution	X(c,t)	$n_c  imes n_t$	Derived quantity

Table 2. List of symbols representing indices, data, fixed effects, random effects, and derived quantities defined in the main text.

Throughout, we use subscripts to indicate either properties of the *i*<sup>th</sup> sample (e.g. area-swept  $w_i$ , location  $s_i$ ), or parameters associated with encounter probability or positive catch rates (e.g.  $\kappa_p$  vs.  $\kappa_r$ ). Indices or scalars have dimension of zero, indicated using a "—" symbol.

Spatial correlation  $\mathbf{R}_p$  between location *s* and location s + h generally declines with an increased distance |h| between the two locations [sometimes termed Tobler's law of geography; Tobler, 1970]. We specify a Matérn function for this correlation, which includes a parameter  $\kappa_p$  governing the distance at which locations are essentially uncorrelated (increased  $\kappa_p$  leads to a decreased decorrelation distance) as well as a transformation matrix **H** representing geometric anisotropy (the tendency for correlations to decline faster in one direction than another):

$$\mathbf{R}_{p}(s,s+h) = \frac{1}{2^{\nu-1}\Gamma(n)} \times \left(\kappa_{p}|h\mathbf{H}|\right)^{n} \times K_{\nu}(\kappa_{n}|h\mathbf{H}|)$$
(5)

where *n* is a smoothness parameter [fixed at 1.0; Simpson *et al.*, 2012] and  $K_n$  is the Bessel function ( $\mathbf{R}_r$  is defined identically but with  $\kappa_r$  in place of  $\kappa_p$ ). Including geometric anisotropy is generally important for fishes along a narrow continental shelf like the US West Coast, where correlations decline faster moving onshore–offshore rather than moving alongshore (Thorson *et al.*, 2015b).

We do not know *a priori* which taxa are likely to be more or less correlated. Therefore, we model covariance  $\mathbf{V}_{\varepsilon p}$  among species using a factor-analysis decomposition:

$$\mathbf{V}_{\varepsilon p} = \mathbf{L}_{\varepsilon p} \mathbf{L}_{\varepsilon p}^{T} \tag{6}$$

where  $\mathbf{L}_{\varepsilon p}$  is a  $n_c$  by  $n_f$  matrix defining the first  $n_f$  columns of the Cholesky decomposition of covariance matrix  $\mathbf{V}_{\varepsilon p}$ ,  $\mathbf{L}_{\varepsilon p}^T$  is the matrix-transpose of  $\mathbf{L}_{\varepsilon p}$ , and  $\mathbf{V}_{\varepsilon r}$  is defined identically but with  $\mathbf{L}_{\varepsilon r}$  in place of  $\mathbf{L}_{\varepsilon p}$  [see Thorson *et al.*, 2016a,b and Warton *et al.*, 2015 for details regarding this factor-analysis decomposition]. Similarly, we specify a factor-analysis decomposition for the covariance  $\mathbf{V}_{\delta p} = \mathbf{L}_{\delta p} \mathbf{L}_{\delta p}^{T}$  among vessel effects:

$$\boldsymbol{\delta}_{\mathrm{p}}(\boldsymbol{\nu}) \sim \mathrm{MVN}(0, \mathbf{V}_{\delta p})$$
 (7)

where  $\delta_p(v)$  is the vector of vessel effects affecting encounter probability  $\delta_p(c, v)$  for all taxa *c* and a given vessel *v* (and where



**Figure 1.** Comparison of estimated density functions (averaged across all years for each taxon) for structure-forming invertebrates, six *Sebastes*, and two *Sebastolobus* species using the VAST multispecies model (bottom row) compared with a single-species estimate for each taxon (top row; see Table 1 for plotting code for taxa; inset colour bar shows average log-density in kg km<sup>-2</sup>, and colours are defined identically for single- and multispecies models for each taxon).

 $\mathbf{V}_{\delta r}$  is defined identically, but with  $\mathbf{L}_{\delta r}$  in place of  $\mathbf{L}_{\delta p}$ ). This factor-analysis decomposition allows the analyst to select an appropriate number of factors  $n_f$  for approximating spatio-temporal covariation or covariation among vessels, where  $0 < n_f \leq n_c$ . Specifying a reduced number of factors  $(n_f < n_c)$  decreases the number of estimated parameters and, therefore, may result in smaller standard errors for other parameters or more precise predictions of local density (Thorson *et al.*, 2015a). However, reducing the number of factors could also result in biased estimates of abundance trends (e.g. by shrinking dynamics for all species onto a small number of dimensions). We leave exploration of this bias-variance trade-off as a topic for future research and instead specify full rank for each covariance  $(n_f = n_c)$  to eliminate this source of potential bias.

Parameters are estimated for the VAST model by maximizing the marginal likelihood of fixed effects given available data. We treat the intercept parameters for each species ( $\gamma_p(c, t)$  and  $\gamma_r(c, t)$ ), the spatial scale of spatio-temporal variation ( $\kappa_{\lambda}$  and  $\kappa_r$ ), the shape of geometric anisotropy (two parameters in **H**), the covariation among species ( $\mathbf{L}_{\varepsilon p}$  and  $\mathbf{L}_{\varepsilon r}$ ), the covariation among vessels ( $\mathbf{L}_{\delta p}$  and  $\mathbf{L}_{\delta r}$ ), and the magnitude of residual variation in positive catch rates for each species ( $\sigma_c^2$ ) as fixed effects. We treat spatio-temporal variation [ $\varepsilon_p(s, c, t)$  and  $\varepsilon_r(s, c, t)$ ] and catchability variation [ $\delta_p(c, \nu)$  and  $\delta_r(c, \nu)$ ]as random effects (Thorson

and Minto, 2015). We define the joint likelihood as the product of the probability of random effects (given fixed effects) and the probability of the data (given random and fixed effects). We then calculate the marginal likelihood of fixed effects while integrating the joint likelihood with respect to random effects. We specifically use the Laplace approximation to approximate the multidimensional integral required to calculate the marginal likelihood (Skaug and Fournier, 2006). The Laplace approximation is implemented using Template Model Builder (Kristensen et al., 2016), and Template Model Builder also provides the gradient of the approximated marginal likelihood with respect to all fixed effects. We use a gradient-based nonlinear minimizer within the R statistical environment (R Core Team, 2015) to identify maximum-likelihood estimate (MLE) of fixed effects. To improve computational efficiency, we use Revolution Open R for low-level parallelization of matrix computations (http://www.revolutiona nalytics.com/revolution-r-open), a stochastic partial differential equation (SPDE) approximation for all spatial processes (Lindgren et al., 2011), and the R-INLA software (Lindgren, 2012) to compute the triangulated mesh used in the SPDE approximation. We distribute code for applying the VAST model to other datasets as an R package on the author's website (www. github.com/james-thorson/VAST) and have confirmed that the VAST model provides identical parameter estimates to a

	Spatio-temporal									
	1	2	3	4	5	6	7	8	9	
SFI-	1.0	0.9	0.4	-0.8	-0.7	-0.9	-0.8	-0.9	-0.9	
L. spine-	0.9	1.0	0.7	-0.8	-0.5	-0.7	-0.6	-0.9	-0.8	
S. spine-	0.4	0.7	1.0	-0.1	0.2	-0.1	0.1	-0.4	-0.5	ility
Dark-	-0.8	-0.8	-0.1	1.0	0.9	0.9	0.9	0.9	0.7	obab
POP-	-0.7	-0.5	0.2	0.9	1.0	0.9	0.9	0.8	0.5	er pr
Sharp-	-0.9	-0.7	-0.1	0.9	0.9	1.0	0.9	0.9	0.8	count
Split-	-0.8	-0.6	0.1	0.9	0.9	0.9	1.0	0.9	0.6	Enc
Stripe-	-0.9	-0.9	-0.4	0.9	0.8	0.9	0.9	1.0	0.9	
Green-	-0.9	-0.8	-0.5	0.7	0.5	0.8	0.6	0.9	1.0	
SFI-	1.0	0.1	0.3	0.1	0.2	0.0	0.2	-0.3	0.6	
L. spine-	0.1	1.0	0.5	0.5	0.6	0.6	0.4	0.3	0.4	
S. spine-	0.3	0.5	1.0	0.5	0.5	0.3	0.4	0.0	0.4	e
Dark-	0.1	0.5	0.5	1.0	0.9	0.9	0.6	0.5	0.7	ch rat
POP-	0.2	0.6	0.5	0.9	1.0	0.9	0.4	0.2	0.7	e cat
Sharp-	0.0	0.6	0.3	0.9	0.9	1.0	0.3	0.3	0.7	ositiv
Split-	0.2	0.4	0.4	0.6	0.4	0.3	1.0	0.6	0.5	Å
Stripe-	-0.3	0.3	0.0	0.5	0.2	0.3	0.6	1.0	0.2	
Green-	0.6	0.4	0.4	0.7	0.7	0.7	0.5	0.2	1.0	

**Figure 2.** Analytic estimate of correlation among fishes and SFI using the VAST model (see Table 1 for taxa abbreviations) for encounter probabilities ( $R_{\epsilon r}$ , top panel) or positive catch rates ( $R_{\epsilon p}$ , bottom panel). Numbered columns correspond to the species groups indicated by the row labels, ordered from top to bottom.

previous spatio-temporal index standardization model (SpatialDeltaGLMM, Thorson *et al.*, 2015b) when applied to data for a single species. However, the VAST model also incorporates most capabilities of spatial dynamic factor analysis for monitoring trends in community abundance or conducting species ordination (Thorson *et al.*, 2016a,b).

After parameters are estimated, we predict the value of random effects by identifying their values that maximize the joint likelihood, given the data and maximum likelihood estimates of fixed effects. We then use the predicted values for random effects to estimate total biomass I(c, t) for each taxon in each year [an "index of abundance"; Thorson *et al.*, 2015b]:

$$I(c,t) = \sum_{s=1}^{n_s} a(s) \times \text{logit}^{-1}[\gamma_p(c,t) + \varepsilon_p(s,c,t)] \\ \times \exp\left[\gamma_r(c,t) + \varepsilon_r(s,c,t)\right]$$
(8)

where a(s) is the area associated with location s[I(c, t)] does not include vessel effects  $\delta_p$  or  $\delta_r$ , because these are interpreted as

representing variation in catchability]. We also estimate the centroid of the distribution X(c, t) for each species in each year [termed "center of gravity"; Thorson *et al.*, 2016b]:

$$X(c,t) = \sum_{s=1}^{n_s} x(s) \begin{cases} a(s) \operatorname{logit}^{-1}[\gamma_p(c,t) + \varepsilon_p(s,c,t)] \\ \times \exp[\gamma_r(c,t) + \varepsilon_r(s,c,t)] \\ I(c,t) \end{cases}$$
(9)

where x(s) can be any statistic used to summarize distribution. We are particularly interested in estimating shifts in fish distribution northward or southward along the US West Coast, so we define x(s) as the distance north of the equator (in kilometres) for location *s*. Standard errors for I(c, t) and X(c, t) are then calculated by Template Model Builder using a generalization of the delta method.

We compare model performance when fitting all species simultaneously (the "multi-species analysis") to a conventional "singlespecies analysis", where each species is fitted individually using the VAST model. To compare performance between single- and multispecies models, we compute the Akaike information criterion (AIC) (Akaike, 1974). The AIC is a measure of model "parsimony" (see Figure 1.3 from Burnham and Anderson, 2002), which we use to identify the level of complexity that likely minimizes the combination of bias (from an overly simple model) and imprecision (from an overly complex model). We compute the "single-species" AIC as the sum of the AIC for the VAST model fitted to each individual species; this comparison is justified because the aggregate of single-species models is fitted to the same dataset as the multispecies model. We also conduct a tenfold cross-validation analysis to determine whether multi- or singlespecies analyses have greater predictive ability. To do so, we divide the data into 10 similarly sized partitions. For the first cross-validation, we estimate model parameters only using data in partitions 2-10, and calculate the probability of data in partition 1 using the predictive distribution, given estimated parameters. This process is repeated for all 10 partitions for the multispecies model. For the single-species model, we conduct this tenfold cross-validation for each species individually, and then sum the resulting log-predictive probabilities for each species.

#### Results

Inspection of density estimates for eight fishes and structureforming invertebrates using the multispecies VAST model shows that species are unevenly distributed throughout the California Current (Figure 1, bottom row). By distribution, the fishes can be broadly classified into three groups: coastwide *Sebastes* spp. (splitnose, stripetail, and greenspotted), northern *Sebastes* spp. (POP, sharpchin, and darkblotched), and *Sebastolobus* spp. (longspine and shortspine thornyheads). The thornyheads are distinguished by having increased densities in the deepest waters farthest from the US coast. Structure-forming invertebrates are found at highest densities offshore near northern Oregon, close to the Oregon– California border and offshore from the south of Monterey Bay through the Southern California Bight.

Our species classifications are supported by the estimated covariance matrices (Figure 2), where longspine and shortspine thornyheads have high pairwise correlations in both encounter probability and positive catch rates (0.5–0.7). Encounter probability of longspine is negatively associated with encounter



**Figure 3.** Relative log-biomass I(c, t) in units of log-metric tons (Equation 8) for each species using single-species (grey) or multispecies (red) modelling (note different *y*-axis ranges for each species and see Table 1 for taxa abbreviations; top row: SFI and *Sebastolobus* group; middle row: northern *Sebastes* group; bottom row: coastwide *Sebastes* group).

probability of Sebastes spp., while encounter probability of shortspine has both positive and negative associations with different Sebastes spp. This difference between thornyhead species is also apparent in distribution maps (Figure 1), where longspine has the deepest distribution of any fish in our analysis, whereas shortspine occupies a more shoreward distribution that overlaps with the spatial distribution for several Sebastes (e.g. darkblotched, POP, and splitnose). The Sebastes spp. all generally have high correlations (0.5-0.9) with one another for encounter probability  $(E_{v};$  Figure 2 top panel), but the northern vs. coastwide groups are strongly distinguished by correlations in positive catch rates ( $E_r$ ; Figure 2 bottom panel), where darkblotched, POP, and sharpchin have higher correlations with one another (0.9) than with splitnose, stripetail, and greenspotted (0.2-0.7). At this coastwide spatial scale, Sebastolobus spp. generally have increased encounter probability when structure-forming invertebrates are found, whereas coastwide and northern rockfish groups have somewhat decreased encounter probability in these cases. When fishes and SFI are encountered, however, an increased catch of SFI is associated with increased catch for all fishes except stripetail and sharpchin rockfish.

We next compare estimates of biomass trends using multispecies and single-species estimates (Figure 3). Biomass trends are broadly similar between models, and particularly for SFI, which shows a trend of increased biomass since 2008. Biomass trend estimates are most different between multi- and single-species models for the group of northern *Sebastes* spp. (darkblotched, sharpchin, and POP; Figure 3 middle row). For example, the multispecies model estimates lower abundance for POP in 2008 than the single-species model. This lower estimate for POP in 2008 using the multispecies model reflects a similar decrease in abundance for darkblotched rockfish in 2008 using either model – the estimate for POP in this year for the multispecies model is "shrunk" towards the estimate for darkblotched rockfish.

Estimates of variation and trends in center-of-gravity (COG) are also generally similar between multi- and single-species model outputs (Figure 4). The notable exceptions are again the northern *Sebastes* spp., specifically POP and sharpchin rockfish, which both have relatively few encounters relative to other species (ca. 500 each, see Table 1). For POP and sharpchin, the single-species estimates of COG are nearly 100 km farther south than COG estimates from the multispecies model (Figure 4 middle row). By



**Figure 4.** Northward center-of-gravity (COG), X(c, t), in units of kilometres north of the equator (Equation 9) for each species using single-species (grey) or multispecies (red) modelling (see Table 1 for taxa abbreviations, and Figure 3 caption for details).

sharing information about positive catch rates (as shown in the lower panel of Figure 2), the multispecies model estimates greater variation in density for these species between different locations off Oregon and Washington and, therefore, estimates a more northward distribution than the single-species model for POP and sharpchin (Figure 1 top row). We again interpret this as a consequence of statistical "shrinkage" for these species, where the multispecies model is sharing information among northern *Sebastes* spp. to infer density hotspots.

Finally, a comparison of standard errors (Figure 5) shows that the multispecies model generally has slightly larger standard error for estimating log-biomass (median 0.01 increase relative to single-species model) and center-of-gravity (median 2.4 km increase). This increased standard error presumably occurs because the multispecies model estimates greater spatial variation in density (Figure 5). For POP, for example, the single-species model estimates little spatial pattern except an increase in density moving northward along the coast, while the multispecies model estimates density hotspots in the same mid-depth areas off the Washington coast as it estimates as good habitat for splitnose and darkblotched rockfishes (Figure 1).

Despite estimating wider standard errors for abundance indices and distribution shifts, the multispecies model provides a more fit to available data. The multispecies model has an AIC score that is 5692.0 better than the combined AIC for singlespecies models, despite the multispecies model estimating an additional 112 parameters (409 fixed effects for the multispecies vs. 297 total among all single-species models). The improvement in fit for the multispecies model is also supported by the tenfold cross-validation analysis, where the multispecies model has a 4-5% greater predictive probability than when analysing each species individually (Table 3). This improvement in predictive score presumably arises because the multispecies model identifies finescale differences in species density for all taxa (e.g. comparing Fig. 1 top and bottom rows for splitnose), and these fine-scale density estimates are on average a useful prediction of variation in catch rates.

#### Discussion

We have used a JDSDM to illustrate strong associations (both positive and negative) between deep-water demersal fishes and structure-forming invertebrates at broad spatial scales along the



**Figure 5.** Comparison of standard error estimates for log-index of abundance,  $\ln[I(c, t)]$  in units of log-metric tonnes (Equation 8, top row) and centre-of-gravity (COG) X(c, t) in units of kilometres north of the equator (Equation 9, bottom row) from single-species and multispecies VAST models, where the scatterplots compare standard error estimates for each year t and taxon c (first column; x-axis shows single-species and y-axis shows multispecies standard errors) and the histograms show the difference between these standard error estimates (second column; where a positive value indicates that the multispecies model had a wider confidence interval than the single-species model for that taxon and year). The dotted line in each histogram indicates the median increase in log-standard error (top row) or standard error (bottom row) for the multispecies relative to the single-species model, and the number in the top-right corner indicates the median increase.

Table 3. Predictive negative log-likelihood for left-out samples (where a low number indicates better fit for the multispecies model) from a
tenfold cross-validation experiment comparing single-species models to a multispecies VAST model that was estimated for all species
simultaneously, as well as the ratio of predictive probability for the multispecies model relative to the single-species model (a value >1.0
indicates better predictive performance for the multispecies model than the single-species model).

		Predictive negative log-likeli		
Partition	Number of cross-validation samples	Single-species VAST model	Multispecies VAST model	Ratio of predictive probability
1	6889	8459.71	8078.84	1.057
2	6832	8560.26	8209.90	1.053
3	6835	8768.53	8377.45	1.059
4	6890	8203.66	7815.19	1.058
5	6799	8335.34	8009.96	1.049
6	6828	8548.48	8154.91	1.059
7	6800	8426.61	8133.02	1.044
8	6997	8688.89	8335.62	1.052
9	6743	8439.27	8077.01	1.055
10	6859	8594.30	8271.42	1.048

US portion of the California Current. These associations vary substantially between two genera *Sebastolobus* (thornyheads) and *Sebastes*, where *Sebastes* can be further divided into northern and coastwide species. Previous work has shown phylogenetic signals in covariation among fishes (Thorson *et al.*, 2015a, 2016a,b) or other species (Ovaskainen *et al.*, 2010), but ours is the first study to (i) use a spatio-temporal statistical model to estimate

covariance between fishes and structure-forming invertebrates, and (ii) decompose this covariation into components representing encounter probabilities vs. positive catch rates [i.e. using the delta models that are conventional in fisheries science; Maunder and Punt, 2004]. Although the JDSDM was more parsimonious and had better predictive performance than single-species models (as shown by AIC and cross-validation analysis), the multispecies analysis resulted in slightly wider confidence interval estimates than analysing data for each species individually.

At a coastwide spatial scale, we estimate an increased encounter probability for Sebastolobus and a decreased encounter probability for Sebastes species where SFIs are present. In contrast, alternative visual sampling at fine spatial scales often shows a large increase in Sebastes density, given the presence of SFIs, and Sebastolobus densities are less often reported to be associated with biogenic habitat (Brodeur, 2001; Tissot et al., 2008; Yoklavich and O'Connell, 2008; du Preez and Tunnicliffe, 2011). Recent research suggests that correlations in distribution among species will often differ when looking at small and large scales (Ovaskainen et al., 2016), and this may explain why our results differ from those from fine-scale visual sampling. Alternatively, differences in results may arise because visual sampling often occurs in rocky habitats, whereas our analysis relies on bottom trawl data that are primarily available in soft-sediment habitats. We recommend future research combining data from small and coastwide scales (and both hard- and soft-bottom habitat) within a single spatio-temporal statistical model, where density-variation at fine scales could be obtained by either fishery-dependent catchrate data or direct observations (Jagielo et al., 2003; Shelton et al., 2014; Rooper et al., 2016; Thorson et al., 2016). We also recommend future research to include habitat variables and associations within size-structured spatio-temporal models (e.g. Kristensen et al., 2014; Nielsen et al., 2014). These models could then estimate separate habitat associations for juvenile and adult fishes and be used to target spatial management towards the more vulnerable or sensitive life stage for protected species.

Based on our results, we find that simultaneously modelling fishes and SFI yields more parsimonious predictions of density and also facilitates estimating variation in density at finer spatial scales than single-species models, even for species with few encounters (e.g. POP and sharpchin rockfishes). However, incorporating these associations when estimating trends in abundance or distribution does not shrink confidence intervals. For an ecologist conducting a stock assessment, incorporating multispecies data may complicate their description of estimated abundance indices, thereby decreasing stakeholder trust in the stock assessment process. Therefore, we imagine that our results will encourage many assessment scientists to continue using single-species models for estimating abundance indices. From a broader perspective, however, the increased parsimony and out-of-sample predictive ability of the multispecies model indicate that estimates of local density are generally improved by jointly modelling multiple species (including both fishes and biogenic habitat). Precise predictions of local density for rare species might be particularly useful for ecosystem modellers, who often initialize spatial ecosystem models using sparse sampling data for rare species or ecosystem components. These estimates of local density could also be used to prioritize areas for spatial management that have a high density of structure-forming invertebrates and fishes. Therefore, we suggest further research regarding the association of fished species and biogenic habitat, including the likely impact of spatial management on fishery productivity in the West Coast groundfish fishery.

#### Acknowledgements

We thank the NWFSC FRAM Fisheries Research Survey Team and the crew on the US West Coast Groundfish Bottom Trawl Survey for collecting the data, and we thank Michelle McClure, Trevor Branch, and Tim Essington for helpful comments and discussion. We also thank Mary Yoklavich, Joe Bizzarro, Chris Rooper, and two anonymous reviewers for comments on an earlier draft.

#### Funding

L.A.K.B. gratefully acknowledges funding from the Joint Institute for the Study of the Atmosphere and Ocean (JISAO) under NOAA Cooperative Agreement No. NA15OAR4320063, Contribution No. 2016-01-38.

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Handling editor: Emory Anderson