# A spatial-temporal approach to modeling somatic growth across inland recreational fisheries landscapes 

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#### Abstract

We develop a mechanistically motivated von Bertalanffy growth model to estimate growth rate and its predictors from spatial-temporal data and compare this model's performance with a suite of commonly used mixed-effects growth models. We test these models with simulated data and then apply them to test whether concerns that high density is causing growth suppression of walleye (Sander vitreus) in Alberta, Canada, are supported using data collected during 2000-2017. Simulation experiments demonstrated that models that failed to account for complex dependency structures often resulted in growth rate estimates that were less accurate and biased low as judged by median absolute relative error and median relative error, respectively. The magnitude of this bias depended on the parameter values used for simulation. For the case study, a spatialtemporal model was more parsimonious and had higher predictive performance relative to simpler models and did not support the slow-growing walleye hypothesis in Alberta. These findings demonstrate the importance of considering spatial-temporal correlation in analyses that rely on surveillance-style monitoring datasets, particularly when examining relationships between life-history traits and environmental characteristics.

Résumé : Nous développons un modèle de croissance de von Bertalanffy à relations mécanistes pour estimer le taux de croissance et ses variables prédictives à partir de données spatiotemporelles et comparons la performance de ce modèle à une série de modèles de croissance à effets mixtes couramment utilisés. Nous mettons ces modèles à l'essai en utilisant des données simulées et les appliquons ensuite pour vérifier si des données recueillies de 2000 à 2017 appuient l'interprétation proposée qu'une forte densité causerait une suppression de la croissance de dorés jaunes (Sander vitreus) en Alberta (Canada). Des expériences de simulation démontrent que les modèles qui ne tiennent pas compte de structures de dépendance complexes produisent souvent des estimations des taux de croissance moins exactes et biaisées vers le bas, comme indiqué par l'erreur relative médiane absolue et l'erreur relative médiane, respectivement. La magnitude de ce biais dépend des valeurs des paramètres utilisées pour la simulation. Pour l'étude de cas, un modèle spatiotemporel s'avère plus parcimonieux et présente une meilleure performance prédictive que les modèles plus simples et n'appuie pas l'hypothèse des dorés à croissance lente en Alberta. Ces résultats démontrent l'importance de tenir compte des corrélations spatiotemporelles dans les analyses qui reposent sur des ensembles de données de type surveillance, en particulier pour l'examen des relations entre des caractères du cycle biologique et des caractéristiques du milieu ambiant. [Traduit par la Rédaction]


## Introduction

Inland recreational fisheries are embedded in complex socialecological systems (see Carruthers et al. 2019), generate $\sim$ US $\$ 51$ billion $\cdot$ year $^{-1}$ in North America (Funge-Smith 2018), and support important ecosystem services when effectively managed (Lynch et al. 2016). However, those tasked with assessing and managing inland recreational fisheries face unique challenges. For example, the status of inland recreational fisheries is often determined from snapshots of monitoring data that have been collected across space and time (e.g., Lester et al. 2003; Post 2013; Lorenzen et al. 2016), which precludes the use of many traditional assessment methods (e.g., see methods in Hilborn and Walters
1992). Thus, a key challenge of managing lakes across landscapes is that it is impossible to carry out in-depth assessments on every lake within a jurisdiction and hence to manage on a lake-by-lake basis (Shuter et al. 1998). Consequently, ecologists often use monitoring data to relate life-history traits to environmental characteristics to inform ecological processes, identify vulnerable populations, and guide management actions across landscapes (e.g., Shuter et al. 1998; Lester et al. 2014; Wilson et al. 2019a).

Little attention has been paid to the issue that the scale at which many inland recreational fisheries operate may introduce spatial and temporal correlation to the monitoring data that management agencies collect from these systems. This is important be-

Received 9 December 2019. Accepted 3 August 2020.
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cause many landscape-scale datasets are observational in nature (Nichols and Williams 2006) and because studies conducted using such datasets need to be vigilant when inferring process from pattern, as a lack of controls, interspersion of treatments, and (or) missing variables can all bias inference (Hurlbert 1984). Thus, the price one pays to analyze landscape-scale data is that there exists the potential for complex correlation structures within those data, which can bias statistical and ecological inference in unexpected ways (Zuur et al. 2017).

Hierarchical modeling has emerged as a powerful tool for disentangling the ecological process of interest from uncontrolled sources of observation error (Royle and Dorazio 2008); however, such modeling has typically assumed independence among lakes or years in the context of inland recreational fisheries. For instance, many studies treat lakes as discrete grouping levels for normally distributed random effects to inform fisheries ecology and management across landscapes (e.g., Helser and Lai 2004; Tsehaye et al. 2016; Wilson et al. 2019a). Whereas lakes are often treated as independent units in mixed-effects models for computational ease, there are intuitive reasons to suspect this assumption may not always hold. For example, both the landscape-scale generalizations and social-ecological systems approaches to inland fisheries recognize that the correct scale of assessment and management is not at the scale of the individual lake - either because of correlated biological or environmental characteristics (e.g., species productivity, primary productivity, temperature, etc.; Myers et al. 1997; Shuter et al. 1998) or because angler movements and hence fish dynamics may be correlated among lakes (Post et al. 2008; Kaemingk et al. 2018). Both views imply that data collected from inland recreational fisheries may be correlated in space or time, and thus methods that account for spatial-temporal correlation may help to separate signal from noise (Cressie and Wikle 2015).

In Alberta, Canada, an 18 -year province-wide monitoring dataset on walleye (Sander vitreus) provides a useful case study with which to explore the challenges of analyzing monitoring data and providing management advice in the presence of spatial-temporal correlation. Walleye are among the most widely distributed and sought-after gamefishes in North America (Bozek et al. 2011), and in Alberta high angling effort and low population productivity resulted in walleye declines and collapsed fisheries by the 1990s (Post et al. 2002). In response to these declines, managers implemented stringent regulations to reduce fishing mortality (i.e., catch-and-release, high minimum length limits), which promoted the partial recovery of walleye in many lakes by the early 2000s (Sullivan 2003). In both January 2018 and 2020, public consultations on fisheries management practices were held throughout the province. A criticism raised by many stakeholders during these meetings was that walleye populations were overabundant and hence growing slowly, termed the Slow-Growing Walleye Hypothesis (SGWH). While these criticisms were anecdotal, ample evidence of density-dependent somatic growth exists in the literature (Post et al. 1999; Lester et al. 2014). Consequently, the SGWH is important to test because angler satisfaction is driven in part by the size of fish caught (Beardmore et al. 2014), and because it implies that managers may need to strike a balance between producing a few large and many small fish (Walters and Post 1993).

This paper addresses the challenge of analyzing landscape-scale fisheries data in the presence of spatial-temporal correlation. To do so, we demonstrate a linkage between a commonly used somatic growth model and its underlying bioenergetics parameters. We then use this derivation to motivate a spatial-temporal hierarchical growth model and compare it with a suite of simpler mixed-effects models using simulated data where truth is known, and we measure model performance using several performance measures and a decision analysis. Finally, we demonstrate the unpredictable outcomes that can occur when ignoring spatialtemporal dependency by applying each of these methods to an

18-year datase to test the SGWH in Alberta. This comparison between simulation experiment and case study application helps demonstrate the potential pitfalls of mis-specifying mixed-effects models when analyzing landscape-scale fisheries data.

## Materials and methods

## Bioenergetics derivation of the von Bertalanffy growth model

The von Bertalanffy growth model is often used by fisheries managers to report fish growth patterns and can be used to test for the effects of covariates on growth rate early in life (Shuter et al. 1998). This growth model is derived from a bioenergetics model specifying

$$
\begin{equation*}
\frac{\mathrm{d} W}{\mathrm{~d} t}=H W_{\mathrm{t}}^{\frac{2}{3}}-m W_{\mathrm{t}} \tag{1}
\end{equation*}
$$

where $W$ is weight at time $t$, and $H$ and $m$ are mass-specific anabolic (building tissue) and catabolic (breaking down tissue) terms, respectively. Solving eq. 1 and transforming from weight to length yields the von Bertalanffy growth function:

$$
\begin{equation*}
\hat{l}_{i}=l_{\infty}\left\{1-\mathrm{e}^{\left[-(\mathrm{K})\left(a_{i}-t_{0}\right)\right]}\right\} \tag{2}
\end{equation*}
$$

where $a_{i}$ and $\hat{l}_{i}$ are the age and predicted length of fish $i$, respectively, $l_{\infty}$ is a parameter that represents the average asymptotic maximum length of fish, $K$ is the unitless Brody "growth" coefficient, and $t_{0}$ is a nuisance parameter that describes age when length is hypothetically zero. This growth model implies that

$$
\begin{equation*}
l_{\infty}=\frac{H}{m} \cdot \alpha^{-\frac{1}{3}} \tag{3}
\end{equation*}
$$

where $\alpha$ is the shape parameter from a weight-length relationship ( $W=\alpha \cdot$ Length ${ }^{\beta}$; see van Poorten and Walters 2016 eqs. 1-6). Furthermore, it can be shown that

$$
\begin{equation*}
K=\frac{m}{3} \tag{4}
\end{equation*}
$$

Thus, it is commonly recommended that studies examining the effect of food availability on growth should focus primarily on $l_{\infty}$ rather than $K$ when using the von Bertalanffy growth model because $l_{\infty}$ is related to the anabolic parameter $H$, whereas $K$ is not (e.g., Walters and Post 1993; van Poorten and Walters 2016). However, a reparameterization of the von Bertalanffy growth model introduces a new parameter, $\omega=K \cdot l_{\infty}$, which represents the growth rate of length measurement units per year in early life and reduces correlation among parameters (Gallucci and Quinn 1979). Combining eqs. 3-4 to recast $\omega$ in terms of the underlying bioenergetics relationships shows that

$$
\begin{equation*}
\omega=\frac{H \cdot \alpha^{-\frac{1}{3}}}{3} \tag{5}
\end{equation*}
$$

Thus, growth rate in early life is exclusively a function of anabolic gains, which scales with feeding rate or food availability (Beverton and Holt 1957, pp. 105-106; Charnov 2010). Additionally, positive covariation between $H$ and $m$ implies that $l_{\infty}$ varies much less with changes in $H$ than would be expected if $H$ and $m$ were independent. Furthermore, while anglers may not care about growth in early life history per se, they do care about the size of their catch (Beardmore et al. 2014), and variation in adult size is linked to growth rate in early life (Lester et al. 2004). Consequently, we
focus on $\omega$ as a proxy for food intake and density-related effects throughout this paper.

## Mixed-effects von Bertalanffy growth models

Here we develop four mixed-effects versions of the von Bertalanffy growth model derived above to use in both our simulation experiment and case study below. The Gallucci and Quinn (1979) von Bertalanffy growth model parameterization is

$$
\begin{align*}
& \left.\hat{\imath}_{i}=l_{\infty}\left\{1-\mathrm{e}^{\left[-\left(\frac{\omega}{l_{\infty}}\right)\left(a_{i}-t_{0}\right)\right.}\right]\right\} \mathrm{e}^{\varepsilon_{i}}  \tag{6}\\
& \varepsilon_{i} \sim \operatorname{Normal}\left(0, \sigma^{2}\right)
\end{align*}
$$

where parameters are defined in the previous section. This equation also assumes that observed lengths follow a lognormal distribution with variance $\sigma^{2}$. A common extension to this model is to incorporate a normally distributed random effect that accounts for unstructured, among-lake heterogeneity. For simplicity, we assume this random effect influences $\omega$ rather than all parameters simultaneously.

## "By-lake" random effects

One mixed-effects parameterization of this model is
(7) $\quad \omega_{\text {lake }} \sim \mathrm{e}^{\omega_{0}+\operatorname{Normal}\left(0, \sigma_{\text {Iake }}^{2}\right)}$
where lake represents a lake-specific deviation from the global intercept $\omega_{0}$, and $\sigma_{\text {lake }}^{2}$ describes the variance in the lake-specific deviations. This "by-lake" random-effects structure is often employed in inland fisheries to assess growth and infer ecological processes from data collected across multiple lakes (see related models in Ogle et al. 2017; Wilson et al. 2019a; Höhne et al. 2020) and represents our first model.
"By-time" random effects
A second version of the von Bertalanffy growth model is

$$
\begin{equation*}
\omega_{\text {year }} \sim \mathrm{e}^{\omega_{0}+\operatorname{Normal}\left(0, \sigma_{\text {year }}^{2}\right)} \tag{8}
\end{equation*}
$$

where year represents a year-specific deviation from the global average $\omega_{0}$, and $\sigma_{\text {year }}^{2}$ now describes the variance of the yearspecific deviations. This "by-time" model applies a single deviation to the average growth rate across all lakes at each time step (see related random effects structures in Matthias et al. 2018; Pedersen et al. 2018). The "by-time" model serves as an intermediate step between the "by-lake" model and the following model.

## "Both" lake and time random effects

A third "both" model features lake-specific random effects and year-specific random effects and thus is more complex than either of the previous two mixed-effects models:

$$
\begin{equation*}
\omega_{\text {lake, year }}=\mathrm{e}^{\omega_{0}+\operatorname{Normal}\left(0, \sigma_{\text {aike }}^{2}\right)+\operatorname{Normal}\left(0, \sigma_{\text {year }}^{2}\right)} \tag{9}
\end{equation*}
$$

Extensions of this third "both" model have also appeared in the literature (see related models in Wagner et al. 2007; Tsehaye et al. 2016; Embke et al. 2019). The models we have noted thus far assume that the random-effect deviations are independent for lakes and (or) time steps.

## Autoregressive lag-1 spatial-temporal (ar1-st) random effects

Next, we introduce Gaussian random fields (GRFs) to extend the models above to explicitly account for structured correlation or
dependency in space and time. GRFs can also help account for the effect(s) of unmeasured covariates on the response variable (Zuur et al. 2017) and have not yet been applied to the von Bertalanffy growth function. A GRF is a multidimensional version of a Gaussian random variable defined by the expectation, variance, and covariance of a multivariate normal distribution (Cressie and Wikle 2015). Specification of a GRF determines the value of a random variable corresponding to any point in space $s$ or spacetime $s, t$ defined on a grid (e.g., an $x-y$ grid for spatial models or an $x-y$-t grid for space-time models).

One often approximates a GRF because full estimation can be impossible, particularly for spatial-temporal models and large datasets such as those collected across landscapes by fisheries management agencies. One such approximation to a GRF is the Stochastic Partial Differential Equation (SPDE) approach (Lindgren et al. 2011), which assumes that a GRF is Markovian in nature (i.e., a Gaussian Markov random field or GMRF), and that the spatial correlation follows a Matérn covariance function (Zuur et al. 2017). The Matérn covariance function requires estimating two parameters: Matérn $\kappa$, which describes the decorrelation rate with distance, and $\tau$, which controls the variance of the spatial noise. The Matérn parameters $\kappa$ and $\tau$ provide values for the spatial covariance matrix $\boldsymbol{\Sigma}$, and then the spatial precision matrix $\mathbf{\Sigma}^{-1}$ is calculated via the analytical methods in Lindgren et al. (2011). The SPDE method requires laying a "mesh" across the study area, and the nodes of this mesh are then estimated as spatially correlated random effects. Spatial range (defined as the distance below which spatial correlation is approximately $\leq 13 \%$ ) and the marginal standard deviation of the spatial process $\sigma_{0}$ are then calculated as derived variables (Lindgren and Rue 2015):
(10) $\quad$ spatial range $=\frac{\sqrt{8}}{\kappa}$
and

$$
\begin{equation*}
\sigma_{0}=\frac{1}{\tau \kappa \sqrt{4 \pi}} \tag{11}
\end{equation*}
$$

We can extend the von Bertalanffy modeling framework to explicitly account for first-order autoregressive spatial-temporal correlation via

$$
\begin{equation*}
\omega_{s, t}=\mathrm{e}^{\omega_{0}+v_{s, t}} \tag{12}
\end{equation*}
$$

where

$$
\begin{align*}
& v_{s, t}=\rho v_{s, t-1}+u_{s, t}  \tag{13}\\
& u_{s, t} \sim \operatorname{GMRF}(0, \mathbf{\Sigma}) \tag{14}
\end{align*}
$$

and

$$
\begin{equation*}
v_{s, t=1} \sim \operatorname{Normal}\left(0, \frac{\sigma_{0}^{2}}{1-\rho^{2}}\right) \tag{15}
\end{equation*}
$$

Here, $v_{s, t}$ represents correlated site $s$ and correlated temporal $t$ deviations for each location and year. A third parameter $\rho$ is constrained to $[-1,1]$ and describes autoregressive temporal correlation between $t$ independent realizations of an isotropic spatial random field $u_{s, t}$, while the term $1-\rho^{2}$ enforces stationarity of the spatial-temporal random field (see separable ar1-st random effects in Cameletti et al. 2011; Blangiardo and Cameletti 2015; Zuur et al. 2017). We use $s$ and $t$ to distinguish between spatial-temporal

Table 1. Parameters used for the von Bertalanffy growth model simulation experiment.

| Parameter | Description | Value |
| :--- | :--- | :--- |
| No. of years | No. of years for which data were simulated | 10 |
| No. of lakes | No. of simulated lakes | 15 |
| No. of fish | No. of simulated fish per lake | 20 |
| Age range | Range of ages | $[0,25]$ |
| $l_{\infty}$ | Average asymptotic maximum size | 55 |
| $\omega_{0}$ | Juvenile growth rate | 14 |
| $t_{0}$ | Hypothetical age when length is zero | -1 |
| $\sigma$ | Standard deviation (SD) in log space of the likelihood function | 0.2 |
| $\sigma_{\text {lake }}$ | SD of the lake random effects | $0.2,0.5,0.8$ |
| $\sigma_{\text {year }}$ | SD of the year random effects | $0.2,0.5,0.8$ |
| $\sigma_{0}$ | SD of the spatial-temporal random effects | $0.2,0.5,0.8$ |
| $\rho$ | Temporal correlation | $0.1,0.5,0.9$ |
| Spatial range | Distance at which spatial correlation decreases to | $10 \%, 50 \%, 90 \%$ of grid |
|  | approximately $\leq 13 \%$ |  |

Note: Parameters with multiple values were varied individually depending on the scenario, but were otherwise held at middle values.
correlated random effects and their independent lake and year random effect counterparts (eqs. 7-9), although we note that for our purposes lake and $s$ correspond to the same location (i.e., a given lake) and that year and $t$ correspond to the same moments in time (i.e., a given year). The covariance matrix $\mathbf{\Sigma}$ is obtained using the estimated parameters $\kappa$ and $\tau$ via the SPDE approach of Lindgren et al. (2011). This "ar1-st" model is formulated such that random effects for $\omega$ are estimated for each lake-year combination, and this model allows growth rates at nearby points in space and (or) time to be more similar than growth rates at locations further away in space and (or) time. Thus, the ar1-st model relaxes the assumptions of lake and year independence assumed by the models presented above (i.e., the by-lake, by-time, and both models).

## Simulation experiment

We conducted a four-by-four factorial experiment to demonstrate the performance of each of the models described in the previous section. Each of the four models (by-lake, by-time, both, and ar1-st) were used to simulate data, and each of the four models was then fitted to each of the simulated datasets. All simulations assumed single values for both $t_{0}$ and $l_{\infty}$ to simplify comparison of patterns among scenarios. For each set of simulations, we simulated a collection of 15 lakes for 10 years and sampled 20 fish from each lake-year combination. Random effects for the by-lake, by-time, and both models were simulated in the R statistical environment (eqs. 7-9; R Core Team 2019), while Gaussian spatialtemporal fields for the ar1-st model were simulated using the GMRFlib package in C (Rue and Follestad 2001). For the ar1-st model, we drew the centroid of each lake randomly from a uniform $[0,10]$ distribution. We obtained fish ages within lakes by randomly sampling integers between 0 and 25 and calculated lengths by then applying the corresponding random effect values and von Bertalanffy growth parameters. We then used calculated lengths to generate observed lengths by adding lognormal noise according to eq. 6. For each model and parameter combination, we generated 300 replicate datasets and then fit all four models to each replicate dataset. We recorded the marginal maximum likelihood estimate of $\omega_{0}$ from each iteration and determined bias and accuracy of models by calculating the median relative error (MRE) and the median absolute relative error (MARE) across iterations for each scenario. We repeated the entire four-by-four factorial experiment nine times (i.e., 43200 hierarchical model fits in total) to explore the effects of varying random effect parameter values on our results, and we varied values of $\sigma, \sigma_{\text {year }}, \sigma_{0}, \rho$, and spatial range from low to high values. We chose parameter values to be representative of walleye life history (Table 1), and all four random-effects structures considered in the simulation experiment are listed in Table 2.

In addition to the performance metrics above, we also conducted a simple decision analysis that identified the MinMax solution, which is the estimation model that resulted in the minimum value of the maximum MARE across all simulation scenarios. MinMax is a tool used in game theory and decision analysis that is useful for identifying an option (i.e., model) that is likely to perform best given uncertainty in or incorrect assumptions about the underlying dynamics of a system (McGilliard et al. 2015). Phrased differently, the MinMax solution represents the best model to use when one does not know which of the simulation scenarios most accurately reflect truth, which is likely the case in most landscape-scale age and growth analyses. We used MARE for the MinMax calculations because it incorporates both bias and variance in a single measure (see also McGilliard et al. 2015).

## Case study application to the Alberta walleye fishery

We then applied each of the models above to Alberta's standardized Fall Walleye Index Netting (FWIN) program data (Fig. 1). Data were collected according to the methods in Morgan (2002). Multimesh gillnets were eight panels $\times 7.6 \mathrm{~m}$ in length, set in randomly stratified locations across the 2-5 and 5-15 m depth strata, and fished perpendicular to shore for 21-27 h during September when water temperatures were $10-15^{\circ} \mathrm{C}$. Twenty-three percent of the FWIN surveys used half the standard net length but the same proportion of mesh sizes. We doubled catch rates in half nets to make them comparable to the standard nets for our effective density calculations (below). Age was estimated using fin rays or otoliths, and we excluded fin ray age estimates $>6$ (Koenigs et al. 2015). Sex was determined via inspection of the gonads, and we retained lakes with a minimum of 50 age-length samples for analysis. A total of 251 FWIN surveys were conducted in 81 lakes during 2000-2017 (surveys per lake: min. $=1$, median $=3$, max. $=$ 12), and 36798 fish were included in the analysis ( 18264 females; 18534 males).

We extended the estimation model from the simulations to incorporate fixed effects to test the SGWH in Alberta. We estimated the degree to which $\omega$ was related to intraspecific effective density, growing degree days (GDD) $>5^{\circ} \mathrm{C}$, interspecific effective density, and an interaction term between intra- and interspecific density:

$$
\begin{equation*}
\omega=\mathrm{e}^{\omega_{0}+\beta_{1} \text { intraspecific }+\beta_{2} \mathrm{GDD}+\beta_{3} \text { interspecific }+\beta_{4} \text { interaction }} \tag{16}
\end{equation*}
$$

plus the random-effect structures in the by-lake, by-time, both, or ar1-st models. We used the intraspecific effective density covariate and its corresponding $95 \%$ confidence intervals from these models as our test of the SGWH in Alberta. We used average effective density as our measure of density because it strikes a balance

Table 2. Random effects structures used in the simulation experiment and in the Alberta walleye case study.

| Model | Description | Random effects structure |
| :---: | :---: | :---: |
| by-lake | Assumes variation around growth rates in lakes is a normally distributed variable. | $\operatorname{Normal}\left(0, \sigma_{\text {lake }}^{2}\right)$ |
| by-time | Assumes growth rates in lakes share temporal deviations, which are assumed to be a normally distributed variable. | $\operatorname{Normal}\left(0, \sigma_{\text {year }}^{2}\right)$ |
| both | Assumes growth rates in lakes vary randomly, while also sharing common temporal deviations among lakes. All random deviations are assumed to be normally distributed. | $\begin{aligned} & \operatorname{Normal}\left(0, \sigma_{\text {lake }}^{2}\right) \\ & \operatorname{Normal}\left(0, \sigma_{\text {year }}^{2}\right) \end{aligned}$ |
| ar1-st | Relaxes the assumptions of the above models and allows growth rate deviations at locations nearby in space and (or) time to be more closely related than those observed at locations further away in space and (or) time. | $\begin{aligned} & v_{s, t}=\rho v_{s, t-1}+u_{s, t} \\ & u_{s, t} \sim \operatorname{GMRF}(0, \Sigma) \\ & v_{s, t=1} \sim \operatorname{Normal}\left(0, \frac{\sigma_{0}^{2}}{1-\rho^{2}}\right) \end{aligned}$ |

Note: $\sigma_{\text {lake }}^{2}=$ variance of the lake-specific random effects; $\sigma_{\text {year }}^{2}=$ variance of the year-specific random effects; $\sigma_{0}^{2}=$ variance of the spatial-temporal field; $\rho=$ temporal correlation; GMRF = Gaussian Markov random field; $\mathbf{\Sigma}=$ covariance matrix; $v_{s, t}=$ random effects correlated in space $s$ and time $t$. The covariance matrix is obtained using the Matérn parameters $\kappa$ and $\tau$, which are estimated using the stochastic partial differential equation approach (Lindgren et al. 2011).
between the number and consumption allometry of fish in a population and thus is a more appropriate measure of density-related effects on fish growth compared with biomass or numbers per hectare (Post et al. 1999). Effective density is defined as the sum of squared fish lengths (i.e., $\Sigma_{i=1}^{n}$ length ${ }^{2}$ ). This index approximates population consumption of prey and therefore represents the best measure of the intensity of exploitative competition (Walters and Post 1993; Post et al. 1999). We calculated average effective density for each lake-year combination as this value divided by the number of nets set in a given lake in a given year. We used all remaining covariates to explore additional variables known to influence walleye growth rate. We estimated GDD based on the latitude, longitude, and elevation of each lake's centroid using ClimateWNA (Wang et al. 2012), and we hypothesized this would have a positive effect on growth rate. We defined interspecific density as the effective density of all non-walleye fish within a FWIN net (see online Supplementary material S1 ${ }^{1}$ for species). Additionally, we included an interaction term between intra- and interspecific density to represent the hypothesis that lakes varied in terms of their overall carrying capacity, which may alter the relationship between growth rates and intraspecific density. We included an additional covariate $\beta$ Sex to account for differences in asymptotic size for males and females and allowed both asymptotic size and $t_{0}$ to vary randomly by lake:

$$
\begin{equation*}
l_{\infty} \sim \mathrm{e}^{l_{\operatorname{lgxploal}^{a}}+\beta S \operatorname{Sex}+\operatorname{Normal}\left(0, \sigma_{\text {lake }}^{2}\right)} \tag{17}
\end{equation*}
$$

and

$$
\begin{equation*}
t_{0} \sim t_{0_{\text {global }}}+\operatorname{Normal}\left(0, \sigma_{\text {lake } \epsilon_{0}}^{2}\right) \tag{18}
\end{equation*}
$$

Covariates were standardized prior to analysis by subtracting the mean and dividing by two times the standard deviation (Gelman 2008). We fit models with restricted maximum likelihood (REML) and maximum likelihood (ML) to facilitate comparison among models with different random effect structures and fixed effects, respectively (Pinheiro and Bates 2006). We also fit models with and without the interaction term to simplify comparisons among models.

We used R-INLA to construct a mesh required for the SPDE approximation (Lindgren and Rue 2015), which was laid over the study area for the ar1-st model (S2 ${ }^{1}$ ). We forced mesh nodes to coincide with lake centroids because we sought to estimate random effects for individual lakes rather than having these values be interpolated among mesh nodes using weighted averages (see Zuur et al. 2017). We also tested whether our case study results were sensitive to our choice of mesh (they were not).

We compared models using Akaike's information criterion (AIC) and predictive scores by conducting two cross-validation experiments to evaluate the predictive performance of our models. Cross-validation involves repeatedly splitting data into "training" and "testing" sets, where models are fitted to a training set and then used to predict the testing set. We used the fitted models to calculate the natural logarithm of the likelihood of data given the predictions for the testing set and divided this by the number of observations in the testing set to obtain a predictive score for each model. Models with the highest predictive scores were deemed most parsimonious for out-of-sample prediction (Gelman et al. 2014). We used two different cross-validation routines to test performance of our models and to safeguard against overfitting: leave-one-lake-out cross-validation, which tested for a model's ability to predict data in a new lake, and h -block cross validation, which tested how well the models predicted blocks of data in space (Roberts et al. 2017).

## Even more mixed-effects models: random slopes ar1-st model

In addition to our primary test of the SGWH using the randomintercept models above, we conducted an additional test of this hypothesis that allowed the intraspecific density covariate to vary by lake. We note that while most inland recreational fisheries random-effects analyses assume variants of the random-intercept model forms we described above (however see Wagner et al. 2007; Tsehaye et al. 2016; Hansen et al. 2020), experience and ecological intuition suggest that the impact of intraspecific density on walleye growth rates may vary by lake (Fig. 2; see also Gelman and Hill 2007). Furthermore, this parameterization provides useful information to fisheries biologists tasked with managing specific lakes (i.e., this model provides information on the relationship between effective walleye density and growth rate for a specific lake unlike the random intercept models above). We used a within-and-

[^0]Fig. 1. Distribution of known but unsampled walleye lakes in Alberta versus walleye lakes where surveys were conducted during 2000-2017 (open versus closed circles, respectively). Inset: location of study area in Canada. Map created in R using publicly available datasets. See Materials and methods for information on datasets.

among-lake parameterization to explicitly separate the effect of walleye density within a given lake from the effective density effect among lakes (see Bell et al. 2018). We constructed an additional ar1-st model to allow the intraspecific density coefficients to vary randomly by group (lake):

$$
\begin{equation*}
\omega_{\mathrm{s}, \mathrm{t}}=\mathrm{e}^{\left.\omega_{0}+\beta_{1} \text { intra-among+( } \beta_{2}+\varepsilon_{\text {lake }}\right) \text { intra-within }+\beta_{3} \mathrm{GDD}+\beta_{4} \text { inter }} \tag{19}
\end{equation*}
$$

where

$$
\varepsilon_{\text {lake }} \sim \operatorname{Normal}\left(0, \sigma_{\text {lake }}^{2}\right)
$$

plus the ar1-st random effects. Here, intra-among represents a lake's mean effective walleye density, intra-within is the effective walleye density centered within each lake, $\beta_{2}$ represents the global slope estimate for walleye density, and $\varepsilon_{\text {lake }}$ represents the random deviation from $\beta_{2}$ for lake-specific intraspecific density effects. Thus, while our primary test of the SGWH in Alberta was
conducted using the random-intercept models above, we conducted this additional test to explore whether the SGWH was supported in any specific lake within the Alberta FWIN dataset. Thus, for this random-slopes ar1-st model we tested whether $\beta_{2}$ plus the lake-specific deviations for slope $\varepsilon_{\text {lake }}$ and the corresponding $95 \%$ confidence intervals overlapped zero.

## Estimation

We implemented all models in Template Model Builder (TMB; see Kristensen et al. 2015) and assessed statistical validity using methods in Zuur and Ieno (2016). TMB uses the Laplace approximation to calculate the marginal likelihood of the fixed effects conditional on the best estimates of the random effects (Skaug and Fournier 2006) and calculates the gradient of the marginal likelihood via automatic differentiation (Kristensen et al. 2015). These are then passed to the R statistical environment and optimized using maximum marginal likelihood via the nlminb() function ( R Core Team 2019). We used the R-INLA software to develop the sparse matrices required for the SPDE approximation for the ar1-st models (Lindgren and Rue 2015) and then passed this information to TMB for estimation. We note that this was necessary because R-INLA does not support nonlinear models such as the von Bertalanffy growth model. We ensured the maximum gradient of the objective function was $\leq 0.001$ and the Hessian matrix was positive-definite to test for consistency with convergence. Additionally, we encountered challenges fitting the ar1-st model to datasets simulated without spatial-temporal correlation. We placed vague penalties (i.e., priors) on both $\rho$ and $\tau$ to improve numerical performance and overcome these issues in the simulations. The penalty for $\rho$ was

$$
\begin{equation*}
\rho \sim \operatorname{Normal}(0,2) \tag{20}
\end{equation*}
$$

while the penalty for $\tau$ was

$$
\begin{equation*}
\log (\tau) \sim \operatorname{Normal}(0,3) \tag{21}
\end{equation*}
$$

Additionally, in situations where $\rho$ was estimated to be 1 , we set $\rho=1$ and re-estimated the ar1-st model so that the fitted model passed our convergence tests. These penalties and estimation techniques were not necessary for models in the case study. All code necessary to recreate our analyses and figures (including maps) is available at https://github.com/ChrisFishCahill/walleye_ growth. Polygons for our maps were taken from publicly available datasets (South 2017), and all other packages and software necessary to create our maps can be found at the link above.

## Results

## Simulation results

Simulations revealed that the ar1-st model was able to recover the true value of $\omega_{0}$ under most of the simulated random effects structures and parameter combinations (Fig. 3). Additionally, the ar1-st model featured low bias (median MRE near zero across scenarios; $\mathrm{S3}^{1}$ ) and demonstrated accuracy comparable to that of the data generation model as judged by MARE across most scenarios $\left(\mathrm{S} 4^{1}\right)$. In the presence of spatial-temporal correlation, the ar1-st model was often less biased and more accurate than all other models (S31; S41), though all models performed similarly well when random-effect noise was low (i.e., when $\sigma_{\text {lake }}, \sigma_{\text {year }}$, or $\sigma_{0}$ were at low levels; left middle panels of Fig. 3; S3 ${ }^{1}$; S4 ${ }^{1}$ ). Performance of all models typically decreased as $\rho$, random-effect noise, or spatial range increased from low to high values (i.e., rows from left to right Fig. 3; S3 ${ }^{1}$; S4 ${ }^{1}$ ).

The four-by-four factorial experiments revealed that estimation models that omitted a source of variation in terms of the true data generation process often resulted in $\omega_{0}$ estimates that were biased low; the magnitude of this bias depended on the parameter values

Fig. 2. Caricature demonstrating the importance of allowing coefficients to vary by group (i.e., lake) in mixed-effects models for landscapescale inland recreational fisheries analyses. In the plot on the left, lake was fitted as a random intercept, and thus lake-specific fits (thin lines) shared a common slope with the global fit (thick line). In the plot on the right, both intercept and slope were allowed to vary by lake (thin lines). Thus, while the latter model is more complex, it allows individual lakes to demonstrate neutral or even positive responses despite a negative global relationship between growth rate and effective density (thick line).


Fig. 3. Simulation results demonstrating the influence of random-effect structure on the estimation performance of von Bertalanffy growth models. For each panel, each model (i.e., "by-lake", "by-time", "both, and "ar1-st") was used to generate 300 datasets (data generation model indicated by $x$-axis groupings). All models were then fit to each replicate dataset, and distributions of the estimated growth rate $\omega_{0}$ are indicated by the point-ranges (circle: 50th quantile; inner range: 25th and 75th quantiles, outer range: 10th and 90th quantiles). Open circles indicate a match between the data generation and estimation model, while solid horizontal lines indicate the true value of $\omega_{0}$. Rows of panels indicate the influence of varying parameters singly on results; thus, row one panels demonstrate the result of increasing temporal correlation $\rho$ of the spatial-temporal field from low to high values, while values of random effect noise $\sigma_{0}$ and spatial correlation range were held at middle values. Note: $\sigma_{\text {lake }}$ and $\sigma_{\text {year }}$ were also varied from low to high values in the $\sigma_{0}$ row, but were otherwise held at middle values (see Table 1 for values). [Colour online.]

used in the simulation (Fig. 3; S3 ${ }^{1}$ ). For example, if data were simulated by lake but the by-time model was used to estimate the simulated data, the resulting $\omega_{0}$ estimates from the by-time model were generally biased low (Fig. 3; S3¹). The opposite was also true: if data were simulated using the by-time model but estimated
using the by-lake model, the resulting $\omega_{0}$ marginal maximum likelihood estimates from the by-lake model were often biased low (Fig. 3; S3 ${ }^{1}$ ). The both and ar1-st models were more flexible and able to partition the random-effect variation of the by-lake and by-time models and thus estimated $\omega_{0}$ with low bias and accuracy

Table 3. Information criteria for models fitted to the Alberta walleye data.

|  | Interaction | LOLO | h-block | CREML <br> Model | term |
| :--- | :--- | :--- | :--- | ---: | ---: |

Note: Akaike's information criterion (AIC) values were calculated using restricted maximum likelihood (REML; for comparing among models with different random effect structures) and maximum likelihood (ML; for comparing among models with and without interaction terms). Average predictive scores were used for leave-one-lake-out and spatial cross-validation (LOLO CV and h-block CV, respectively).
comparable to that of the true model ( $\mathrm{S} 3^{1} ; \mathrm{S} 4^{1}$ ). This general pattern remained as the true data generation process increased in complexity. For example, the ar1-st model often performed comparably well in terms of MRE and MARE to the both model when the both model was used to generate data, but the converse was not always true, as $\omega_{0}$ estimates from the both model were often biased low when the ar1-st model was used to generate data (Fig. 3; S3¹; S4¹). Results from the decision analysis showed that the ar1-st model was the MinMax solution, indicating that this model was the most robust choice across the potential axes of model misspecification considered in the simulation study (MinMax MARE values: by-lake: 13.8\%; by-time: 13.0\%; both: 13.1\%; ar1-st: $12.8 \%$ ).

## Case study results

Despite increased complexity, the ar1-st models were more parsimonious than simpler mixed effects models as judged by $\triangle$ REML AIC scores when fit to the case study data (Table 3). Additionally, the ar1-st models featured the highest out-of-sample predictive performance for both leave-one-lake-out and spatial h-block crossvalidation scores (Table 3). Comparisons of $\Delta$ ML AIC scores between models with and without interaction terms showed that the ar1-st model did not support inclusion of the interaction term, while $\Delta$ ML AIC scores for simpler models did support the inclusion of an interaction term between intra- and interspecific effective density.

The top-ranked model as determined by both AIC values and cross-validation scores was the ar1-st model without an interaction term (Table 3; Fig. 4; see $5^{1}$ for 251 lake-year fits to data). This model estimated that female walleye grew to an $l_{\infty_{\text {slobal }}}$ of 60.1 cm ( $95 \%$ CI: $58.2-61.9 \mathrm{~cm}$ ) versus 55.1 cm for males ( $95 \%$ CI: $53.4-$ 56.7 cm ), while $t_{\text {oglobal }}$ was -0.95 year ( $95 \%$ CI: -1.03 to -0.88 year). The global intercept for $\omega_{0}$ was $14.4 \mathrm{~cm} \cdot$ year $^{-1}(95 \%$ CI: $13.4-$ $15.5 \mathrm{~cm} \cdot$ year $^{-1}$ ), and lake- and year-specific growth rates estimated using the ar1-st model ranged from 7.9 to $26.3 \mathrm{~cm} \cdot$ year $^{-1}$ (Fig. 5; S5 ${ }^{1}$ ). The top-ranked model indicated that the SGWH was not supported in Alberta (point estimate: 0.03 ; $95 \%$ CI: $-0.01-0.07$ ), and none of the parameters hypothesized to affect $\omega$ were significantly different from zero (Fig. 6). This model also estimated that the spatial range of correlation among lakes was $52 \mathrm{~km}(95 \% \mathrm{CI}$ : $41-63 \mathrm{~km}$ ) and that $\rho$ was 0.93 , indicating there was high temporal persistence in the spatial-temporal field ( $95 \%$ CI: $0.90-0.95$ ). The ar1-st model with random slopes also supported the conclusions of the top-ranked model. The within-lake regression coefficient for intraspecific density using the random slopes model was 0.03 ( $95 \%$ CI: $-0.09-0.14$ ), and lake-specific estimates of the relationship between walleye growth rate and intraspecific density indicated that the SGWH was not supported in any of Alberta's lakes using FWIN data collected during 2000-2017 (Fig. 7). We present residual diagnostic plots for the top ranked ar1-st model in $\mathrm{S}^{1}$ and S7 ${ }^{1}$.

Fig. 4. Lake-year fits ( $n=251$ ) from the ar1-st model without an interaction term compared with the observed data collected during 2000-2017. The thick orange and blue lines represent the best fits for females (open circles) and males (closed circles), respectively, while thin lines represent lake- and year-specific deviations in growth for each sex. Note that dots are jittered for visualization. [Colour online.]


## Comparison of the by-lake and ar1-st models in the case study

Given the prevalence of the by-lake model in the literature, we also compared key parameter estimates and $95 \%$ CIs from this model to values from the top-ranked ar1-st model. Estimates and uncertainty intervals for $\omega_{0}$ from the by-lake model were shifted lower relative to the corresponding values from the ar1-st model (left panel, Fig. 6). These two models also differed in terms of the SGWH test (second panel, Fig. 6), and the by-lake model tended to provide point estimates that were farther from zero relative to the ar1-st model (right four panels; Fig. 6). The by-lake model also tended to estimate narrower 95\% CIs for parameters thought to influence growth rate relative to the ar1-st model (right four panels; Fig. 6).

## Discussion

We have shown how growth rate in early life - as per the von Bertalanffy growth function - is linked to anabolism and then used this derivation to motivate a general model that estimates growth rate and its predictors from landscape-scale data with spatial-temporal correlation. Additionally, we demonstrated via simulation that this first-order autoregressive spatial-temporal

Fig. 5. Predictions of walleye growth rate $\left(\omega, \mathrm{cm} \cdot\right.$ year $\left.^{-1}\right)$ through space and time as estimated by the ar1-st model using Fall Walleye Index Netting data collected during 2000-2017. Maps created in R using publicly available datasets. See Materials and methods for information on datasets. [Colour online.]


Fig. 6. Comparison of key parameters (restricted maximum likelihood estimates $\pm 95 \%$ CI) estimated with the ar1-st and by-lake models using length-at-age data collected throughout Alberta during 2000-2017. Note: estimates from the first four panels were obtained from models fit without the interaction term, whereas the fifth panel was obtained from models fit with the interaction term included in the model. Solid horizontal lines indicate no effect.

(ar1-st) von Bertalanffy growth model provided superior growth rate estimates relative to a suite of simpler mixed-effects models when spatial-temporal correlation was present in the data. Borrowing a tool from decision analysis, we also showed the ar1-st
model was most likely to minimize the worst errors in situations when one does not know the underlying random-effects structure of a given dataset. Notably, our simulation experiments also revealed that failure to specify a random-effects structure that was

Fig. 7. Lake-specific estimates ( $\beta_{2}+\varepsilon_{\text {lake }}$ ) and $95 \%$ CIs of the effect of intraspecific effective density on walleye growth rate $\omega$ from the ar1-st random slopes model using Fall Walleye Index Netting data collected during 2000-2017.

flexible enough to capture the underlying complexity of the simulated data both decreased accuracy of growth rate estimates and resulted in estimates that were biased low relative to true values. Furthermore, for a landscape-scale case study on walleye in Alberta, we demonstrated that the ar1-st growth model vastly outperformed several simpler mixed-effects models in terms of both parsimony and out-of-sample prediction performance. Lastly, the ar1-st model found no evidence of density-dependent growth in the Alberta walleye case study, which contrasted the conclusions obtained from the simpler by-lake mixed-effects von Bertalanffy growth model and the claims of some anglers in the province. We explore these findings and their implications in detail below.

An important result from our simulations was that unmodeled dependency in mixed-effects von Bertalanffy growth models often results in growth rate estimates that are biased low; however, this bias largely disappeared when using the ar1-st growth model that explicitly accounted for complex random effects structures. We suggest this bias is related to a lack of independence (presence of pseudoreplication) in the simulated data, which increases both parameter bias and the type I error rate in regression models when not accounted for (Hurlbert 1984; Zuur and Ieno 2016). Furthermore, we speculate that this bias is negative due to nonlinearities in the von Bertalanffy growth model. We believe our findings parallel those of Miller et al. (2018), who showed that temporal correlation inflated the effect size of covariates thought to influence growth unless this source of dependency was explicitly accounted for using state-space modeling. This finding has broad implications given the prevalence of the von Bertalanffy growth model in the literature (e.g., Hilborn and Walters 1992; Walters and Essington 2010; Lee et al. 2017), the importance of obtaining accurate and reliable growth rate estimates for both stock assessments and basic ecological research (Lorenzen et al. 2016; Korman et al. 2017; Lorenzen and Camp 2019), and because the simpler hierarchical models we used here are often considered the de facto standard for analyzing age and growth data (e.g., Helser and Lai 2004; Ogle et al. 2017). Our findings demonstrate that ecologists should be wary when applying mixed-effects growth models to analyze landscape-scale datasets given the near ubiquity of spatial-temporal correlation in real-world data (Cressie and Wikle 2015) and because of the challenges of determining how such correlation will impact an analysis (Zuur et al. 2017; this study). Thus, we recommend that investigations examining relationships between life-history traits such as growth and environmental characteristics explicitly incorporate the effects of space and time and note that our findings are consistent with mounting evidence that spatial-temporal models improve estimates of density dependence (Thorson et al. 2015b), abundance (Royle et al. 2013), species distributions (Thorson et al. 2015a), and rare or extreme ecological events (Anderson and Ward 2019).

Our results from both the top-ranked ar1-st model and the random slopes ar1-st model show that the available data do not support the conclusion that walleye are experiencing densitydependent growth in Alberta. This is surprising given the range of densities observed due in part to the continued recovery of walleye in the province (Sullivan 2003), the wide range of growth rates we documented (e.g., see Figs. 4 and 5), and numerous studies demonstrating density-dependent juvenile growth with similar datasets in other fisheries (e.g., Lester et al. 2014; Ward et al. 2016). We expected the effects of density to be most pronounced during early life-history stages (Lorenzen and Camp 2019) and thus tested for density effects on a parameter that approximates maximum juvenile growth rate (Gallucci and Quinn 1979); however, there may be issues with this approach. First, the von Bertalanffy growth model has been criticized on theoretical grounds, as it has been suggested that this model cannot cleanly separate juvenile and adult growth rates (see Rennie et al. 2008). Nonetheless, the von Bertalanffy model we implemented provided an index of juvenile growth rate in centimetres per year, and we showed that
the growth rate parameter we focused on is a consequence of tissue-building processes using simple mechanistic relationships (see also van Poorten and Walters 2016). We believe that such a first-principles approach to modeling holds great potential for gaining new insights in complex ecological systems, similar to recommendations for both population ecology and spatialtemporal modeling more generally (Brännström and Sumpter 2005; Cressie and Wikle 2015). With this in mind, future studies could adapt more complex growth models to explicitly incorporate spatial-temporal random effects using a predictive process approach (see models in Latimer et al. 2009). Second, the approach we used treated $t_{0}$ and $l_{\infty}$ as standard, lake-specific random effects instead of spatially or time-varying quantities, which may not be warranted (Lee et al. 2017; this study). Spatial-temporal effects for these portions of the growth model were inestimable given the available data, although informative priors could be used to improve estimation of these quantities in the future. Lastly, it is possible that density dependence affects walleye in life stages other than early growth or that density effects could interact with lifetime growth schedules in ways we did not consider (see Lorenzen and Camp 2019).
The finding that walleye in the province were not experiencing density-dependent growth rate suppression during 2000-2017 contrasts the claims of some anglers, and we believe this discrepancy has important implications for fisheries management in Alberta. Some anglers report catching more fish than they did in the 1990s and that many of the fish they do catch are below the harvestable size. Numerous conversations with anglers during public consultation meetings in January and February 2020 revealed that anglers often capture many sublegal size walleye before capturing a single harvestable-size fish and that many interpret this as evidence for slow growth rates and (or) stunting; this is then used to advocate for more liberal harvest regulations (C.L. Cahill, personal observation). However, we explicitly tested the density-dependent growth hypothesis and showed that growth rate suppression was not supported. We suggest that it is important to remember that fishing mortality caused many accessible walleye populations in Alberta to decline and collapse by the mid-1990s (Post et al. 2002) and that stringent regulations such as minimum length limits, catch and release, and a novel harvest lottery tool have since been used to improve walleye abundance in many of the lakes (Sullivan 2003; Spencer 2010). Additionally, survey data in some lakes clearly show that walleye grow until they reach harvestable size and that beyond this size, length frequency histograms appear truncated. This truncation often coincides with the minimum length limit for a given lake (see examples in Spencer 2010). We note that while harvest regulations vary by lake and year across the province, this size truncation is most apparent in several lakeyears in $\mathrm{S5}^{1}$ where 50 cm minimum length limits have been in place for a minimum of 5 years ( $\mathrm{S}^{1}$; see Buck Lake 2005-2010, 2017; Iosegun Lake 2005; Round Lake 2017; Smoke Lake 2005, 2013; Sturgeon Lake 2017). This information coupled with our findings suggests that the reason anglers catch many small fish is not because those fish grow slowly or are stunted, but rather because large fish may be removed from these populations by harvest. Ironically, if harvest was increased in an attempt to improve growth rates, fish may be exposed to unsustainable levels of exploitation and decline toward population levels observed in the 1990s (see Post et al. 2002; Sullivan 2003; Spencer 2010).

Differences between the by-lake model and top ranked ar1-st model illustrate an important finding of our work; failure to account for complex correlation structures in real-world datasets can have unexpected consequences for an analysis and thus change ecological inference (see also Fig. 6; Zuur et al. 2010, 2017; Cressie and Wikle 2015). This is relevant given many landscapescale inland fisheries studies often consider some variant of the by-lake model as their most complex hierarchical model (Ward et al. 2016; Rypel et al. 2018; Höhne et al. 2020). Not only were error
bars from the by-lake model typically narrower than those in the ar1-st model, but point estimates for certain parameters also changed from negative to positive values and were typically larger in magnitude (Fig. 6). Additionally, the by-lake and ar1-st models ultimately resulted in different inferences regarding the SGWH in Alberta. While we do not know the underlying "truth" in the case study, we believe it is likely that the ar1-st model provides a better appraisal of the inherent complexity and uncertainty present in the Alberta walleye dataset, particularly when considered in conjunction with our simulation findings and our case study model comparisons using AIC and cross-validation.

From an applied perspective, disparities between the by-lake and ar1-st models could result in different advice being provided to biologists and policy makers managing walleye in Alberta. For example, the commonly used by-lake model indicated that sizebased fisheries objectives may require reductions in walleye densities through more liberal harvest regulations to improve growing conditions (see arguments in Walters and Post 1993; Wilson et al. 2016). In contrast, the better-supported ar1-st model showed that reducing density was unlikely to improve walleye growth rates. Thus, naïve adoption of the simpler hierarchical model's results could result in harvest policies not supported by the data and which could lead to overfishing. This finding is noteworthy given the prevalence of datasets originating from surveillancestyle monitoring programs in both fisheries and applied ecological settings more generally and because such datasets are often used to inform resource management across broad spatial-temporal scales (see examples in Lester et al. 2003; Nichols and Williams 2006; Cain et al. 2019). While decision analysis would ideally be used to guide management actions such as these, we note that such analyses are rarely employed across landscapes in inland recreational fisheries (see Punt and Hilborn 1997; van Poorten and MacKenzie 2020). As a result, we suggest that coupling spatialtemporal methods such as those used here with decision analysis and value of information modeling may represent a fruitful area for future research in inland recreational fisheries (see Hansen and Jones 2008).

The top-ranked model as judged by AIC and cross-validation indicated that walleye growth rates were correlated to a spatial range of about 52 km in Alberta and that there was high temporal persistence in this spatial-temporal field. This distance is remarkably similar to results presented in Myers et al. (1997), who showed that recruitment in freshwater fishes was spatially correlated to distances of about 50 km . These authors hypothesized that the correlation they detected in recruitment was primarily due to the effects of "planktonic patchiness" and (or) predation in otherwise isolated lakes. Similarly, growth rates in our study were spatially correlated among lakes at short distances after we accounted for sex, intraspecific and interspecific effective density, and temperature, even though lakes were not physically connected. The processes we modeled suggest that this residual correlation may be due in part to similarities in primary productivity and (or) in how this food is partitioned among communities in nearby lakes, as both factors could govern the food supply available to walleye within a given lake and hence local growth rates. The residual spatial-temporal correlation we documented may represent a fundamental limitation of this dataset, and experimentation may be necessary to further resolve our collective understanding of the remarkable variation in walleye growth rates across this large inland recreational fishery landscape (see Figs. 4 and 5; see also Walters 1986; Walters et al. 1988).

Many of the key issues in inland recreational fisheries science and management inherently require spatial-temporal thinking (e.g., see Arlinghaus et al. 2017), and yet spatial-temporal statistical techniques remain underused in the literature (however see Myers et al. 1997; Isaak et al. 2014; Hocking et al. 2018). When compared with a suite of commonly used mixed-effects von Bertalanffy growth models that assumed lake and year indepen-
dence, the spatial-temporal growth model introduced here provided more accurate and less biased growth rate estimates in datasets similar to those found in many applied ecological settings. Furthermore, the spatial-temporal growth model improved fits to real-world data when compared with the simpler mixedeffects models, and differences in key predictions between the model types could have resulted in misleading biological advice to resource managers. While this study focused on nonlinear spatial-temporal growth modeling, we see several potential avenues for future research. For example, questions involving spatial exploitation patterns near urban areas (Wilson et al. 2019b), the effective design of landscape-scale monitoring programs or adaptive management experiments (Walters 1986; Williams et al. 2018), and the examination of fish-habitat relationships to inform fisheries management practices (Grüss et al. 2019) can all be confronted within the modeling framework presented here. Thus, we hope that embracing the spatial-temporal complexity inherent to inland recreational fisheries will result in richer ecological inferences and improve policy recommendations in the future.

## Acknowledgements

We thank the biologists and technicians involved in the FWIN program. CC acknowledges critical conversations with M. Faust, G. Courtice, K. Wilson, and J. Thorson and is supported by a Vanier Canada Graduate Scholarship and a research grant from the Alberta Conservation Association. D. Goethel and three anonymous reviewers improved an earlier version of this manuscript.

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[^0]:    ${ }^{1}$ Supplementary data are available with the article through the journal Web site at http://nrcresearchpress.com/doi/suppl/10.1139/cjfas-2019-0434.

