Including unsexed individuals in sex-specific growth models

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Sexually dimorphic growth models are typically estimated by fitting growth curves to individuals of known sex. Yet, macroscopically ascribing sex can be difficult, particularly for immature animals. As a result, sex-specific growth curves are often fit to known-sex individuals only, omitting unclassified immature individuals occupying an important region of the age-length space. We propose an alternative whereby the sex of the unclassified individuals is treated as a missing data problem to be estimated simultaneously with the sex-specific growth models. The mixture model we develop includes the biological processes of growth and sexual dimorphism. Simulations show that where the assumed growth model holds, the method improves precision and bias of all parameters relative to the data omission. Ability to chose the correct combination of sex-specific and sex-generic parameters is also improved. Application of the method to two shark species -where sex can be ascribed from birth- indicates improvements in the fit but also highlights the importance of the assumed model forms. The proposed method avoids discarding unclassified observations thus improving our understanding of dimorphic growth.

Key words: Dimorphism; EM algorithm; missing data; non-linear clustering; partial classification.
Introduction

Growth is a central process in the life history of an organism (Beverton and Holt 1957; Werner 1986; Starck and Ricklefs 1998; Nentwig 2012) and describing growth is a regular task across many taxa (Lacointe 2000; Krebs and Cowan 1962; Starck and Ricklefs 1998; Jones et al. 2011). Reflecting the importance of understanding growth, modelling growth has a long and productive history in ecology (Robertson 1923; Brody 1926; von Bertalanffy 1938). A commonly applied parametric functional form for the relationship between length and age of given life history stages is the von Bertalanffy model (von Bertalanffy 1938; Chen et al. 1992). Early methods to fit this non-linear model used the linear differenced relationship between mean lengths separated by a unit of age (Ford 1933; Walford 1946). Maximum likelihood fitting under the assumption of normally distributed errors was introduced by Kimura (1980). A sophisticated range of fitting methods now exists, including: non-normal error distributions, more stable re-parameterizations (Gallucci and Quinn 1979; Schnute 1981; Francis 1988), hierarchical methods (Helser and Lai 2004; Brunel and Dickey-Collas 2010; Weisberg et al. 2010), and error-in-variables approaches (Cope and Punt 2007).

Sexually dimorphic growth occurs in many taxa (Fairbairn et al. 2007). Differential selection pressures, mortality schedules and the degree of reproductive investment affect the ultimate differences in size (Roff 1982; Parker 2006). Where it occurs, sexually dimorphic growth has important implications for management. Natural mortality can be size-dependent (Pauly 1980). Harvesting methods are also often size-based, such that the probability of removal changes with size, e.g., small fish might escape through the mesh whereas larger fish are too big to escape (Myers and Hoenig 1997). Where differences exist in the growth of the sexes, selective size-based removal will enact differential mortality between the sexes (Kendall and Quinn 2012). To understand and mitigate for size-selective removals, it is essential that sex-specific growth characteristics are well described.

On first inspection, fitting sex-specific growth models to length-at-age data requires that the sex of an individual is known. For maturing and mature animals, sex can often be assigned visually from primary or secondary sexual characteristics, but immature fish, amphibians, insects and reptiles often require histological methods to distinguish between the sexes (e.g., Vitale et al. 2012).
Histological methods are often not feasible for large-scale continuous sampling programs. As a result, many sex-specific growth datasets contain the sex-designations: Female, Male and Unclassified (FMU).

Extant approaches to fitting sex-specific growth curves to FMU data proceed by first removing unsexed (immature or otherwise unclassified) individuals and then fitting sex-specific curves to the remaining known-sex individuals (e.g., using the methods of Kimura (1980)). Sex-specific growth models of this type are thus fit to a subset of the observed age-length data; we term these FMU fits. FMU fits introduce an extrapolation of the model over an omitted space (typically young immature animals) that may be very informative for certain parameters of the model, particularly the growth rate and intercept. As an extreme illustration, consider a species with determinate growth (somatic growth ceases post-maturation) with all individuals undergoing maturation at a given age (when the sex can be determined). Known-sex lengths will be constant over age and therefore only informative for sex-specific asymptotic size. Without additional assumptions, sex-specific growth rates and intercepts will be confounded. In reality, many species such as many molluscs, fish and reptiles have indeterminate growth (growth continues post-maturation) but we contend that fitting sex-specific growth curves to FMU data in the extant fashion amounts to extrapolation, the severity of which will depend on: ability to determine sex; duration of the immature stage; degree of post-maturation growth; and how protracted the maturation process is across individuals.

Instead of omitting unclassified individuals when fitting sex-specific growth models to FMU data, we propose that the sex of the unclassified individuals be treated as a missing data problem to be estimated simultaneously with the growth curves. We develop a mixture model for this purpose where few restrictions are made on the form of the growth curves or on the between- or within-individual variability but we do focus on the von Bertalanffy model, as a commonly applied functional form (Pardo et al. 2013).

Our objectives are to: (1) develop a mixture model for including unclassified individuals in sex-specific growth curves; (2) develop an accompanying expectation-maximisation (EM) fitting algorithm (Dempster et al. 1977); (3) test the performance of the method relative to the extant FMU approach via simulation; and (4) fit to two species of shark where known sexes of
immature and mature individuals provide a useful real data test of performance.

Materials and Methods

We first motivate the unsexed individuals as a missing data problem, then propose a mixture model solution with an accompanying EM-algorithm estimation routine; simulation tests and real data applications follow.

A partially classified two-component mixture model

The observed data consist of a sample of \( i = 1, \ldots, n \) individuals with recorded variables: length \( l_i \), age \( a_i \), and observed sex \( s_{\text{obs},i} \) where \( s_{\text{obs},i} \in \{F, M, U\} \) (Female, Male, Unclassified). For the overall population, following the notation of McLachlan and Peel (2000), we define a two-component (female \( F \) and male \( M \)) mixture model with a length probability density function of:

\[
f(l|a, \Psi) = \pi_F f_F(l|a, \theta_F) + \pi_M f_M(l|a, \theta_M),
\]

where: \( \Psi = \{\pi_F, \theta_F, \theta_M\} \) is a vector of all free parameters; \( \pi_F \) is the mixing proportion, which is the overall probability that the sex is female, \( \pi_F = P(S = F) \), and the complement \( \pi_M = 1 - \pi_F \) is the overall probability that the sex is male; \( f_S \) is the sex-specific (female or male) component density, e.g., lognormal for a single observation (presented without bias correction):

\[
f_S(l_i|a_i, \theta_S) = \frac{1}{l_i \sigma_S \sqrt{2\pi}} \exp \left( -\frac{(\ln(l_i) - \ln(g(a_i, \Omega_S)))^2}{2\sigma_S^2} \right),
\]

where \( \theta_S = \{\sigma_S, \Omega_S\} \) is the sex-specific density parameter vector; \( g(a_i, \Omega_S) \) is the sex-specific growth function, e.g., von Bertalanffy

\[
g(a_i, \Omega_S) = L_{\infty,S} \left(1 - e^{-K_S(a_i-t_{0,S})}\right),
\]
where $\Omega_S = \{L_{\infty,S}, K_S, t_{0,S}\}$ is the sex-specific growth curve parameter vector consisting here of the mean asymptotic length, Brody growth rate (rate at which asymptote is approached) and age at length zero, respectively. Alternative forms for $g$ include other asymptotic, segmented, and additive curves.

To estimate the parameters of the mixture model, we introduce the partially observed indicator variable $Z$ that denotes the true sex of observation $i$

$$z_i = \begin{cases} 
1, & \text{if observation } i \text{ is female, } s_i = F, \\
0, & \text{if observation } i \text{ is male, } s_i = M.
\end{cases} \quad (4)$$

Letting $i = 1, \ldots, m$ index the classified individuals for which the sex is known, we propose that for the remaining $j = m + 1, \ldots, n$ individuals, $z_j$ can be treated as missing data to be imputed from the posterior probability of an unclassified observation being female. Next, we introduce how classified and unclassified individuals are treated within the estimation.

**Classified individuals**

For each $i$, the true value of $z_i \in \{0, 1\}$ and is known. The conditional density of the classified observation can therefore be written

$$f(l_i|a_i, z_i, \Psi) = f_F(l_i|a_i, \theta_F)^{z_i}f_M(l_i|a_i, \theta_M)^{1-z_i}, \quad (5)$$

and the density of the group (female or male) membership

$$f(z_i|\Psi) = \pi_F^{z_i}\pi_M^{1-z_i}. \quad (6)$$

Assuming independence, the joint density of the classified observation and the group membership can thus be written

$$f(l_i, z_i|a_i, \Psi) = [\pi_F f_F(l_i|a_i, \theta_F)]^{z_i}[\pi_M f_M(l_i|a_i, \theta_M)]^{1-z_i}. \quad (7)$$
The joint density of all the classified observations is

\[
f(l, z | a, \Psi) = \prod_{i=1}^{m} [\pi_F f_F(l_i | a_i, \theta_F)]^{z_i} [\pi_M f_M(l_i | a_i, \theta_M)]^{1-z_i},
\]

(8)

the right-hand side of which is the likelihood of the classified data for a given set of growth model parameters.

**Unclassified individuals: Expectation step**

Unclassified individuals are typically immature but could also consist of a sample of unsexed mature animals. The density of the unclassified observations is a mixture and written, as in Equation (1), as:

\[
f(l_j | a_j, \Psi) = \pi_F f_F(l_j | a_j, \theta_F) + \pi_M f_M(l_j | a_j, \theta_M).
\]

(9)

For each \(j\), the true value of \(z_j \in \{0, 1\}\) but is now unknown. Similar to Dean et al. (2006) and McLachlan and Krishnan (2008), a key quantity is the expected value of the group membership indicator for unclassified individuals given the observed data and a given set of parameters (denoted by an asterisk)

\[
E(Z_j | a_j, l_j, \Psi^*) = Pr(Z_j = 1 | a_j, l_j, \Psi^*).
\]

(10)

This pivotal probability is available via Bayes’ theorem (Bayes 1764), as:

\[
Pr(Z_j = 1 | a_j, l_j, \Psi^*) = \frac{Pr(Z_j = 1) Pr(l_j | z_j = 1, a_j, \Psi^*)}{Pr(l_j | a_j, \Psi^*)},
\]

(11)

\[
= \frac{\pi_F^* f_F(l_j | a_j, \theta_F^*)}{\pi_F^* f_F(l_j | a_j, \theta_F^*) + \pi_M^* f_M(l_j | a_j, \theta_M^*)}.
\]

(12)

The estimated component membership probabilities are denoted by \(z_j^*\). Equation (12) thus provides a means for assigning the probability of group (i.e., female or male) membership to unclassified individuals for a given growth model and set of parameters. We can thus proceed, as for classified individuals, to an expected value for the log-likelihood, conditional on the
expected values of the group membership (termed the “complete data log-likelihood”)

\[
E[\ln L(\Psi | l_j, z^*_j, a_j)] = \sum_{j=m+1}^{n} z^*_j \ln(\pi_F f_F(l_j | a_j, \theta_F)) + (1 - z^*_j) \ln(\pi_M f_M(l_j | a_j, \theta_M)).
\] (13)

As an aside, the simplicity here is due to the log-likelihood based on a representation, as in Equation (8), that is linear in \(z^*_j\). Combining the classified and unclassified individuals, the expected complete data log-likelihood of all observations is then given by

\[
E[\ln L(\Psi | l, z, a)] = \sum_{i=1}^{m} [z_i \ln(\pi_F f_F(l_i | a_i, \theta_F)) + (1 - z_i) \ln(\pi_M f_M(l_i | a_i, \theta_M))] + \sum_{j=m+1}^{n} [z^*_j \ln(\pi_F f_F(l_j | a_j, \theta_F)) + (1 - z^*_j) \ln(\pi_M f_M(l_j | a_j, \theta_M))].
\] (14)

**EM algorithm**

Estimation of the expected log-likelihood (Equation 14) is carried out by the following steps:

1. **E-step**

   Starting with a given set of parameters \(\Psi^{(0)} = \{\theta^{(0)}, \pi_F^{(0)}\}\), calculate the expected value of the log-likelihood (Equation 14), which amounts to finding \(z^*_j\) in Equation (12).

2. **M-step**

   a. Maximise the expected value of the complete data log-likelihood with respect to the growth parameters

   \[
   \theta^{(1)} = \arg\max_{\theta} E[\ln L(\theta | l, z^{(0)}, a, \pi_F^{(0)})].
   \] (15)

   Non-linear growth models require numerical optimisation for this step.

   b. Update the unconditional probability of being female (sex ratio), if not assumed fixed, via

   \[
   \pi_F^{1} = \frac{\sum_{i=1}^{m} z_i + \sum_{j=m+1}^{n} z^*_j^{(0)}}{n},
   \] (16)
which assumes the same overall sex ratio for the unclassified and classified; but this can be relaxed to classified or unclassified only, fixed at a given value, or modelled with covariates where data allow.

3. Replace $\Psi^{(0)}$ with $\Psi^{(1)} = \{\theta^{(1)}, \pi^{(1)}_F\}$ and repeat steps 1 and 2 until convergence, as assessed by a stopping criteria based on a relative change in the observed log-likelihood.

The observed log-likelihood is given by

$$\ln L(\Psi^*|l, z, a) = \sum_{i=1}^{m} (z_i \ln (\pi^*_F f_F(l_i|a_i, \theta^*_F)) + (1 - z_i) \ln (\pi^*_M f_M(l_i|a_i, \theta^*_M))) + \sum_{j=m+1}^{n} \ln (\pi^*_F f_F(l_j|a_j, \theta^*_F) + \pi^*_M f_M(l_j|a_j, \theta^*_M)).$$

(17)

Note that it may be possible to maximise this partially classified observed likelihood directly without the EM algorithm, but direct maximisation does not in general perform well, except in the vicinity of the maximum or via a grid search.

Parameter inference and standard errors

In the same manner as Kimura (1980), parameters may be bound or free across the sexes. We fit all combinations of bound and free parameters and choose the best fitting model by Bayesian Information Criterion (BIC using the: observed data log-likelihood (Equation 17); log of the number of observations; and number of parameters of the component distribution and the mixing proportion). A preliminary investigation of the performance of the Akaike Information Criterion showed it to select overly complex models compared to BIC. Various options are available to approximate the standard errors of the parameter estimates from an EM algorithm (McLachlan and Peel 2000). A key difficulty is obtaining the observed information matrix. The method we implement is to fit the model to convergence via the EM algorithm as above and then optimise the partially-classified mixture log-likelihood (Equation 17) directly, including the mixing proportion. The EM algorithm finds the maximum likelihood which would be difficult to do via direct optimisation. Once at the maximum, direct optimisation is relatively straightforward and can be used to estimate the curvature of the observed log-likelihood and
hence provide standard errors.

**Method testing**

*Simulations*

Our simulation framework study focusses on some of the primary factors influencing the estimation of sex-specific von Bertalanffy growth models for non-hermaphroditic species. Two overall functional types were tested: asymptotic ($L_{\infty,F} = 40\text{cm}; k_F = 0.3\text{year}^{-1} t_{0,F} = -1\text{years}; \text{and } \sigma_F = 0.1 \text{ lognormal error standard deviation}$); and a more ‘linear’ form ($L_{\infty,F} = 40\text{cm}; k_F = 0.1\text{year}^{-1} t_{0,F} = -1\text{years}$) (Figure 1). The distribution of ages (50 female and 50 male per simulation) was drawn from a negative binomial distribution with a mean age of 5 and a shape parameter of one giving an age distribution with 5th and 95th quantiles at 0 and 16 years, respectively. Dimorphic differences (20%) for males were included by parameter settings with: $L_{\infty,M} = 0.8L_{\infty,F}; k_M = 0.8k_F$ and $t_{0,M} = 0.8t_{0,F}$. Early and late maturation were included by setting the age at which 50% of the population are mature to ages two and five, respectively; a maturation range between 25% and 75% mature was set to one year. To emulate the scenario where immature animals cannot be sexed, we set all immature animals to unclassified sex (see Discussion). These values provided a realistic but pragmatically limited set of sexually-dimorphic differences (Figure 1) to illustrate the main aspects of performance without over-loading with simulation output. All combinations of sex-specific and sex-generic parameters (16 models in total) were fit to each simulation dataset and the best fit for each method chosen on the basis of the lowest BIC. Simulation fits were started at the true value for all methods. Each simulation scenario was run 1,000 times with different random draws to provide stable sampling distributions.

![Figure 1 about here.](image)

The performance of the methods on the simulated data was appraised using: 1) parameter root-median-square error (RMSE); 2) classification rates (based on maximum a posteriori / binary classification); 3) ability to chose the correct model form via BIC; and 4) visually via boxplots of the relative error ($\frac{\hat{\theta} - \theta_{true}}{\theta_{true}}$) of the parameter estimates.
Real data

Elasmobranchs (sharks and rays), represent a useful taxon to test the performance of the method given that the sex can be distinguished from birth, owing to two extensions of the pelvic fins (claspers) on the males. We use two datasets on deepwater lantern shark species: velvet belly, *Etmopterus spinax*, and smooth lanternshark, *Etmopterus pusillus*, which are found off the south and southwestern coasts of Portugal (Coelho et al. 2005). The datasets consist of 733 (*E. spinax*) and 518 (*E. spinax*) age-length readings. The data are cross-sectional in that there is a single data point per individual. Females represented approximately 60% (*E. spinax*) and 43% (*E. pusillus*) of the samples, respectively. High proportions of the individuals were classified as immature (*E. spinax*: 60%; *E. spinax*: 77%) based on macroscopic investigation of the reproductive organs (Coelho and Erzini 2007, 2008).

To test the performance of the method, we compare sex-specific von Bertalanffy fits from: (1) all individuals, using mature and immature sex determinations; (2) only the mature animals - to mimic the FMU situation where sex cannot be distinguished prior to the onset of maturation; and (3) all individuals with the sex of the immature animals treated as an unknown classification problem and fit via the mixture model developed here. A grid of plausible starting values was used to initialise fitting for each method.

Software

We developed the R package *lhmixr* (*life history mixture models in R*: available on CRAN or at https://github.com/mintoc/lhmixr/) to fit sex-specific growth models with missing classifications. von Bertalanffy models are implemented with analytical gradients for all parameters in the complete-data log-likelihood and a generic binding formulation to allow parameters to be bound equal or free between the sexes. Normal and log-normal error distribution assumptions are included. Both real datasets are included and documented in the *lhmixr* package.
Results

Simulation performance

Example simulation fits show that, where fish can be classified early (Figure 2(A)), the FMU approach can perform well in recovering the full data curve, though with decreased precision (Table 1, Figure 3). In contrast, for late maturing animals the example FMU fit performed poorly for males and females (Figure 2(B)) and in general had increased bias and decreased precision for the growth rate and intercept terms, particularly for strongly asymptotic growth curves (Table 1, Figure 3 (C, E)).

[Table 1 about here.]

Overall, as judged by RMSE, the MIX model (mixture model developed here) outperforms the FMU approach for all parameters and both curve typologies (Table 1). The differences in RMSE are most pronounced in the asymptotic fits where the MIX-estimated intercept $t_0$ often displays an order of magnitude difference in RMSE over the FMU fits (Table 1). For early maturing animals with linear growth, the RMSE is reduced by half or more using the MIX method relative to the FMU approach (Table 1). The MIX method consistently had RMSE slightly larger than the full knowledge fit, reflecting the additional uncertainty when classification is needed (Table 1).

[Figure 2 about here.]

All methods had similar ability to detect sex-specific difference in the asymptotic size, irrespective of curve typology or maturation timing (Table 2, $L_{\infty}$ scenarios). Ability to detect differences in the growth rate is considerably reduced in the FMU fits compared to the full data fits, particularly for late maturing animals with more asymptotic growth where the correct model was identified in only 14% of simulations (Table 2, asymptotic growth form and late maturing FMU $K_s$ scenario). This was improved to 46% chosen correct using the MIX model. The performance is not as poor for the more linear growth form (33% FMU chosen correctly), though the MIX model still improved the performance (69% chosen correctly). When $t_0$ differs by sex and the growth curve typology is asymptotic, the FMU approach performed poorly.
in choosing the correct model (7% and 2% for early and late maturing animals, respectively) (Table 2, FMU $t_{0,s}$ scenarios); this is somewhat improved for early maturing animals using the MIX model (36%) but is low for the late maturing animals (24%) (Table 2, MIX $t_{0,s}$ scenarios). With asymptotic growth and where $t_0$ or $K$ differ by sex and the FMU approach is used, the chosen model is often that of no difference between the sexes (Table 2, FMU None row).

Omitted models from Table (2) often included the correct model but with residual standard deviation differing by sex (e.g., where only $L_\infty$ differs by sex, a model with $L_{\infty,s}$ and $\sigma_s$ is the most commonly chosen model of those not shown).

Working with the best fitting model per simulation, the classification rates where typically lower when using the FMU model; whereas the MIX model recovered similar but consistently lower classification rates to the FULL data scenario (Table 3). Relative to the FMU approach, large gains in the classification rates are seen where growth is asymptotic (Table 3, third row). As expected, classification rates are poor across all methods where no true differences exist among the sexes (Table 3, “None” columns).

Reflecting the RMSE results, differences in the relative error across methods are least for the $L_\infty$ parameter, reflecting the predominant importance of mature animals for estimating that parameter (Figure 3). In many cases the relative error of the MIX model is comparable to that of the full data scenario for these simulations (Figure 3).

Real data performance

The two elasmobranch species investigated spend a relatively long period of their observed lifespan in the immature phase in addition to a relatively large size at maturation (Figure 4). As a result the best fitting FMU $t_0$ estimates for female and male $E.\ spinax$ and $E.\ pusillus$ are considerably more negative than when the full data are used (Table 4, sixth column).
FMU-estimated intercepts also had considerably decreased precision with no evidence of a sex-specific difference. These biases and lack of precision simply reflect that the FMU model excludes those immature animals and thus only fits to the mature animals. In contrast, the MIX approach results in more comparable estimates to the full data scenario.

Estimates of $L_\infty$ are similar across methods and both sexes for *E. pusillus*. The FMU and mixture models estimate a lower asymptotic size for female *E. spinax* (Figure 4(C,E)) most likely because they are more influenced by the larger known sex females, which are fit poorly in the full data scenario (Figure 4(A)). The *E. spinax* male asymptotic size is estimated as lower in the FMU fit compared to the full data fit; whereas the mixture model estimated a larger and more uncertain male asymptotic size (Table 4, row 6).

The female *E. pusillus* mixture model estimated growth rate is faster than the full data scenario and results in a higher curve (Figure 4(F)). As a result many of the smaller fish for a given age are classified as male, whereas this is not true from the full data, where there is a mixture of males and females. This may arise from the choice of functional form in that there appears to be a broadly similar growth pattern for until older ages where the male growth slows considerably for both spacies (see Discussion). As the FMU approach has smaller males and the immature animals fall under both curves, the predicted classification is unity for males but very poor for females (Table 4). The mixture model improves on this but goes to a solution that separates the sexes more than is apparent in the true data (Figure 4). Overall the classification rates from all models, including the full data scenario, are poor across both sexes owing to a lack of clear separation.

Discussion

Discarding unclassified data degrades the precision and accuracy of fitted sex-specific growth models. Here, we have proposed a redress that simultaneously attempts to classify unclassified individuals while estimating parameters of a growth curve of interest. To our knowledge this
is the first recognition of the potential to couple classification and life history model parameter estimation in this manner.

**Performance**

Where the fitting assumptions match the data-generating assumptions, as in our simulations, the mixture model offers general improvements on the estimation of sex-specific growth curves from partially-classified data (Table 1, Figure 3). Since a lognormal distribution and the von Bertalanffy model are the most common assumptions for these data we think that the method should also improve fits to real datasets.

Where animals can be classified at early ages, the FMU approach may provide good estimates of the parameters of the growth curve (Figure 2). This does have the cost of reduced precision (Table 1, Figure 3), which will decrease the ability to correctly choose differences between the sexes (Table 2). In reality, the impact of the omission of unclassified individuals will vary by population depending on how early immature animals can be classified. Even where classification can occur at a relatively young age, maturity is often dependent on reaching a threshold size (Stearns and Koella 1986). Within an early age-group, larger fish may mature first and hence be distinguished earlier, which may further bias estimates of FMU-derived sex-specific growth compared to when all individuals are included, as proposed here.

Other solutions in the FMU case may include fixing the intercept at a certain value. Though this may seem a simpler solution, it is not recommended given the resulting bias in the growth rate for single fits (Pardo et al. 2013). In addition, for sex-specific fits, it is difficult to envisage how fixing the intercept term would not bias the other sex-specific parameters where differences exist.

Lantern shark datasets investigated demonstrated some extreme challenges to fitting with the FMU approach (Figures 4). In reality, it may be possible to ascribe sex for some individuals prior to maturity but these fits serve to emphasise some of the issues, as discussed below.
Caveats

A primary concern with the von Bertalanffy model is that the functional form of the growth curve differs between the immature and mature phases. Such valid concerns have given rise to alternative mechanistic growth models such as the biphasic growth model of Roff (1983) where somatic growth is linear to the onset of maturity and thereafter a decreasing function of the gonadosomatic index. The von Bertalanffy model theoretically describes only post-maturity growth (Ricker 1975; Lester et al. 2004). We stress, however, that the form of the growth model \( g \) in Equation 3) we use for our derivation or implementation of the algorithm is not restricted and our use of the von Bertalanffy form only reflects its common application. However, hockey-stick implementations of sex-specific Roff biphasic growth models could be very difficult to fit to asymptotic datasets containing unclassified immature animals. The mixture model developed here may contribute in this regard. Overall, it is important to recognise that the method classifies on the basis of the assumed model, so a functional form mis-match may result in poor classification (e.g., \( E. pusillus \), Figure 4). It is difficult to guard against this, other than by comparing with more local functional forms.

The error distribution assumption will affect the classification - we therefore recommend careful consideration of the error-distribution and the mean-variance relationship thus implied. In the \textit{lhmir} package we provide two error distributions (normal and lognormal) but envisage other density forms could be useful in particular applications (e.g., gamma, heteroscedastic formulations). We envisage many ways of improving and highlight the \( E. pusillus \) fits (Figure 4) as a caution of how the performance may be critiqued in given applications.

Of additional concern is the assumption that the sex of the individual is fixed. Fish and other taxon display a great diversity of sexual life histories (Warner 2012), including sequential hermaphroditism, where the sex of the animal can change. On first inspection, classifications to such families as Labrids or gobies, for what may be a non-constant trait, could be a less than sensible approach but again we propose a possible development via the inclusion of relevant covariates for the mixing proportions (Equation 16).

In terms of model choice and parameter inference, it is known that the regularity conditions of the asymptotic distribution of the deviance break down when comparing mixture models with...
differing numbers of components (McLachlan and Peel 2000). The same is true of AIC and BIC, though BIC has been shown to perform well in some situations for identifying the number of components (Wang et al. 1996). We recommend first using BIC to compare a fully saturated two component model (all parameters sex-specific) with a single component model, where all parameters are the same across sexes. Once the number of components is decided, and interest lies in parameter significance, nested models within that number of components can be tested using likelihood ratio tests with a chi-squared distribution. Overall, we propose that the asymptotic distribution of the deviance in this partially classified mixture model requires further work.

Management implications

Given that yields and mortality schedules are directly impacted by growth parameters, having reliable estimates of sex-specific growth parameters is paramount for management of dimorphic species. While the mixture model offers general improvements over the unclassified omission case (Figure 3), we envisage the method assisting most where growth is more strongly asymptotic after maturation (Figure 2). Where faster growing individuals mature earlier and are therefore classifiable earlier, positive biases may exist in previously estimated sex-specific growth curves. This would imply faster growth than when all the data are used. In this setting the method could contribute to management by using all the data to estimate the relevant growth parameters and quantify uncertainty around the sex-specific growth curves. These curves can then be used, for example, in age-length based population assessment models (e.g., Methot 2000; Frøysa et al. 2002).

Future developments

While the approach developed here focusses on partially classified datasets, the mixture model could also be applied to datasets where no sexed animals are recorded, as in common applications to fully unclassified datasets (McLachlan and Peel 2000).

Extending the method longitudinally would allow for the life histories of unclassified individ-
uals to be understood in the context of sex-specific differences. Longitudinal features could be built in by modelling the correlation structure of the errors (e.g., using generalized least squares, estimating equations or random effects (Diggle et al. 2003)).

Other sex-specific life history relationships may also be amenable to the methods developed here. We are currently developing a similar approach for the treatment of maturity. Perceivably these could be coupled into a multivariate partially classified mixture model, where it is explicitly recognised that animals develop along multiple coupled processes.

The estimation time is short (seconds) and \textit{lhmixr} von Bertalanffy complete data log-likelihoods are also accompanied by their analytical gradients to provide more stable fitting. The EM algorithm can, however, be comparatively slow in terms of the number of iterations taken to convergence. This could be improved by implementing the loops in a compiled language (e.g., C++). It would also be of particular interest to start the model at the M-step (e.g., assuming an equal probability of 0.5 for unclassified individuals) and test performance compared to starting at the E-step with a grid of starting values (as done here) or starting at the FMU solution.

Bayesian estimation of finite mixture models is an active area of development (Frühwirth-Schnatter 2006), which would assist straightforward estimation of parameter uncertainty using MCMC, for example, but with associated chain convergence and runtime overheads.

Despite the long history of application of mixture models to ecological problems (Pearson 1895; Bhattacharya 1967; Macdonald and Pitcher 1979) and recent applications (Thorson et al. 2011; Cosgrove et al. 2014), we note in closing that the EM algorithm is at present a considerably under-recognised algorithm in ecological modelling. The EM algorithm contributes to areas as diverse as state space modelling (Shumway and Stoffer 1982), structural equations (Ullman and Bentler 2003), hidden Markov systems (McLachlan and Krishnan 2008), all of which are important topics in modern ecological modelling, yet other routines (MCMC, Laplace, quadrature approximations to the marginal likelihood) are more commonly used when estimating complex ecological models (e.g., Bolker et al. 2013; Kristensen et al. 2015). While recognising the central importance of these methods, we contend that for many users there is a lack of understanding as to how the methods work. In contrast, the steps of the EM algorithm are transparent when estimating unobserved constructs. We believe that the EM algorithm de-
veloped here demonstrates a small amount of the potential of this family of algorithms, which
deserve more recognition in modern ecological modelling.

Acknowledgements

We are grateful to Dr Guillaume Bal, Marine Institute, Ireland for valuable feedback. Two
anonymous reviewers are thanked for greatly improving the manuscript. We thank John Corco-
ran for initial discussions on sex-specific growth models for perch that instigated development
of the method. CM thanks Dr Conor Nolan for encouraging insights on animal growth.

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Table 1: Root-median-square error by: parameter, method and whether maturation occurred early or late in the lifespan of the simulated species. *Asymptotic* and *linear* refer to the form of the simulated curves. Full, FMU and MIX refer to the complete knowledge, mature animals only and mixture model, respectively.

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<th>$K$</th>
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<th>$t_0$</th>
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<td>0.076</td>
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</table>
Table 2: Proportion of time correct model chosen. True model refers to the true simulated model (e.g., true $L_{\infty,s}$ means the sexes differ in their asymptotic size), “Chosen” refers to the model chosen via BIC in fitting. Full, FMU and MIX refer to the complete knowledge, mature animals only and mixture model, respectively. Early and late denote maturation age. All combinations of bound and unbound parameters by sex were fitted (such that each setting sub-column sums to unity) but only the four true model results are presented.

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<th>$L_{\infty,s}$</th>
<th>$K_s$</th>
<th>$t_{0,s}$</th>
<th>MIX</th>
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<th>$K_s$</th>
<th>$t_{0,s}$</th>
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</tr>
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Table 3: Median maximum a posteriori classification rate across iterations. *Asymptotic* and *linear* refer to the form of the simulated curves. FULL, FMU and MIX refer to the complete knowledge, mature animals only and mixture model, respectively.

<table>
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<th>$K_s$</th>
<th>$t_{0,s}$</th>
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<td></td>
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<td></td>
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<td>0.69</td>
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<td>0.48</td>
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</table>
Table 4: Best fitting (lowest AIC) von Bertalanffy parameter estimates by method for two deepwater lantern shark species *Etmopterus spinax* and *Etmopterus pusillus*. Full, FMU and MIX refer to the complete knowledge, mature animals only and mixture model, respectively. Standard errors are provided in parentheses.

<table>
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<tr>
<th>Species</th>
<th>Sex</th>
<th>Method</th>
<th>$\ln L_\infty$</th>
<th>$\ln K$</th>
<th>$\ln -t_0$</th>
<th>$\ln \sigma$</th>
<th>Class rate</th>
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<tbody>
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<td>FULL</td>
<td>4.03 (0.04)</td>
<td>-2.16 (0.07)</td>
<td>0.7 (0.05)</td>
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<td></td>
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<td>0.2 (0.04)</td>
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<tr>
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<td>0.41 (0.03)</td>
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</tr>
<tr>
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<td>Male</td>
<td>FULL</td>
<td>4.03 (0.04)</td>
<td>-2.34 (0.06)</td>
<td>0.92 (0.04)</td>
<td>0.44 (0.04)</td>
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<tr>
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<td></td>
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<td>FULL</td>
<td>3.96 (0.01)</td>
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<td>0.42 (0.03)</td>
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</tr>
<tr>
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<td></td>
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<td>4.07 (0.08)</td>
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<td>0 (0.06)</td>
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<tr>
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<td>-1.91 (0.03)</td>
<td>0.88 (0.05)</td>
<td>0.34 (0.04)</td>
<td>0.91</td>
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</table>
Figure 1: Simulation framework illustration. Asymptotic and linear (weakly asymptotic) refer to the form of the simulated curves. Rows denote parameter differences between females and males. Female and male mean curves are shown in black and grey, respectively. Dashed vertical lines denote early and late maturation ages.
Figure 2: Example simulations for an early maturing (left column) and late maturing (right column) population. FMU and MIX refer to the mature animals only and mixture models, respectively. Full data curves are displayed as solid lines (females black, males grey). Estimated FMU and MIX curves are shown as short-dashed and long-dashed curves, respectively. Known-sex individuals are shown as circles and unknown-sex individuals as squares. The fill colour of the points denotes the probability of the individual being female (from the MIX model for unknowns).
Figure 3: Boxplots displaying the distribution of the relative error by: parameter, method, and whether maturation occurred early (white) or late (grey) in the lifespan of the simulated species. Boxes correspond to the 25th and 75th percentiles, whiskers extend to the closest points to one and a half times the interquartile range. Full, FMU and MIX refer to the complete knowledge, mature animals only and mixture model, respectively. Asymptotic and linear refer to the form of the simulated curves (Figure 1). Y-axis range is restricted to facilitate comparison of most distributions.
Figure 4: Best fitting (lowest BIC) von Bertalanffy fits to age-length data from two deepwater lantern sharks species *Etmopterus spinax* and *Etmopterus pusillus*. Full, FMU and MIX refer to the complete knowledge, mature animals only and mixture model, respectively. Estimated curves for females and males are shown as solid and dashed lines, respectively. Known-sex individuals are shown as circles and unknown-sex individuals as squares. The fill colour of the triangles in the MIX model case denote the probability of the individual being female. Age values are jittered for visualization.