# Identifying direct and indirect associations among traits by merging phylogenetic comparative methods and structural equation models 

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#### Abstract

1. Traits underlie organismal responses to their environment and are essential to predict community responses to environmental conditions under global change. Species differ in life-history traits, morphometrics, diet type, reproductive characteristics and habitat utilization. 2. Trait associations are widely analysed using phylogenetic comparative methods (PCM) to account for correlations among related species. Similarly, traits are measured for some but not all species, and missing continuous traits (e.g. growth rate) can be imputed using 'phylogenetic trait imputation' (PTI), based on evolutionary relatedness and trait covariance. However, PTI has not been available for categorical traits, and estimating covariance among traits without ecological constraints risks inferring implausible evolutionary mechanisms. 3. Here, we extend previous PCM and PTI methods by (1) specifying covariance among traits as a structural equation model (SEM), and (2) incorporating associations among both continuous and categorical traits. Fitting a SEM replaces the covariance among traits with a set of linear path coefficients specifying potential evolutionary mechanisms. Estimated parameters then represent regression slopes (i.e. the average change in trait Y given an exogenous change in trait X ) that can be used to calculate both direct effects ( X impacts Y ) and indirect effects ( X impacts Z and Z impacts Y ).


[^0]4. We demonstrate phylogenetic structural-equation mixed-trait imputation using 33 variables representing life history, reproductive, morphological, and behavioural traits for all $>32,000$ described fishes worldwide. SEM coefficients suggest that one degree Celsius increase in habitat is associated with an average $3.5 \%$ increase in natural mortality (including a $1.4 \%$ indirect impact that acts via temperature effects on the growth coefficient), and an average 3.0\% decrease in fecundity (via indirect impacts on maximum age and length). Cross-validation indicates that the model explains 54\%-89\% of variance for withheld measurements of continuous traits and has an area under the receiver-operator-characteristics curve of 0.86-0.99 for categorical traits.
5. We use imputed traits to classify all fishes into life-history types, and confirm a phylogenetic signal in three dominant life-history strategies in fishes. PTI using phylogenetic SEMs ensures that estimated parameters are interpretable as regression slopes, such that the inferred evolutionary relationships can be compared with long-term evolutionary and rearing experiments.

## KEY-WORDS

evolutionary mechanisms, life history strategies, phylogenetic trait imputation, population and community ecology, structural equation model, trait-based approach, phylogenetic comparative methods

## 1 | INTRODUCTION

Trait-based approaches are essential for improving our understanding of ecological and evolutionary processes. For example, they are used to identify population and community responses to global change (Pacifici et al., 2017), community assembly rules (Gross et al., 2021; Legras et al., 2019), and predict how changes in community diversity affect ecosystem functioning (Díaz et al., 2013) and ecosystem services (Hevia et al., 2017). They can also be used to test theory regarding evolutionary mechanisms (Baker et al., 2020) and support biodiversity conservation (Cardillo et al., 2008). Traits of floristic and faunal species can be quantitative (discrete or continuous) and/or qualitative (binary, nominal, or ordinal variables). For instance, continuous traits include growth rates, body or leaf size, and age at maturity, while categorical traits encompass behaviours (e.g. solitary or gregarious species), diet (autotroph, heterotroph, mixotroph) or reproduction (dispersal modes, guarding vs. nonguarding young) (Hadj-Hammou et al., 2021; Violle et al., 2007).

Trait values are not available for every species of interest, both due to limited scientific resources and ongoing difficulties in collecting and/or sharing trait information across taxa and systems (although see Gallagher et al., 2020). Consequently, there are many potential methods available to impute these missing trait values (Azur et al., 2011; Goolsby et al., 2017; Schrodt et al., 2015). Comparisons of phylogenetic trait imputation (PTI) methods generally show that performance is improved by including phylogenetic information (Debastiani et al., 2021; Penone et al., 2014; Taugourdeau et al., 2014), or even using taxonomy as a proxy for
phylogeny (Johnson et al., 2021) wherein related taxa are more likely to share similar traits than unrelated taxa.

PTI generally involves specifying a statistical process for how trait values change along a phylogenetic (Goolsby et al., 2017) or taxonomic tree (Schrodt et al., 2015; Thorson, 2020; Thorson et al., 2017). This involves estimating parameters to represent correlations $\mathbf{R}$ among $n_{g}$ taxa for a given trait, as well as covariance $\mathbf{\Sigma}$ among $n_{j}$ traits. For example, the function phylopars in R -package RPHYLOPARS is a common implementation for PTI but it cannot be implemented for categorical traits (Johnson et al., 2021; Penone et al., 2014), while such traits are generally easier to assess and collect than continuous ones. Additionally, estimating $\mathbf{\Sigma}$ without constraints (beyond the requirement that it is symmetric and positive definite) has three main limitations (Grace, 2006): (1) results cannot be compared easily with slopes estimated in conventional regression models, such that results are difficult to interpret or validate using experimental data; (2) existing methods cannot use evolutionary theory and experiments to specify the structure of covariance among traits; and (3) the number of parameters in $\boldsymbol{\Sigma}$ without other constraints is $n_{j}\left(n_{j}+1\right) / 2$, which becomes computationally challenging to fit when interpolating a large number of traits.

As an alternative to estimating the covariance among traits directly, we propose to use structural equation modelling (SEM) to specify a parsimonious structure for this trait-covariance $\boldsymbol{\Sigma}$. Given a set of $n_{j}$ traits $\left\{Y_{1}, Y_{2}, \ldots, Y_{n_{j}}\right\}$ with measurements $\left\{y_{1}, y_{2}, \ldots, y_{n_{j}}\right\}$, SEM allows the user to specify a set of dependencies linking these, where each dependency is represented by a path coefficient. These links can be interpreted as a graph wherein


FIGURE 1 Conceptual diagram to illustrate trait correlations using two hypothetical examples involving fish or avian responses to temperature, assuming that temperature affects body size, which in turn affects one continuous and one categorical trait in each example. Analyses start by assembling trait measurements, where values are available for some but not all of six taxa. These conceptual models are then formalized by specifying a text file listing associations, and this in turn can generates the matrix $\boldsymbol{\Gamma}$ (for illustration we assume $\gamma=0.5$ for all associations), and then are used to compute the covariance among traits $\boldsymbol{\Sigma}=\mathbf{L L}^{t}$, where $\mathbf{L}=(\mathbf{I}-\boldsymbol{\Gamma})^{-1} \mathbf{V}^{0.5}$ and $\mathbf{V}$ represents exogenous covariance (evolutionary drift). For illustration we specify $\operatorname{diag}(\mathbf{V})=1$ and convert the covariance to a correlation matrix, shown for each taxon. In practice, associations $\boldsymbol{\gamma}$ (used to form $\boldsymbol{\Gamma}$ ) and exogenous variances $\mathbf{V}$ are estimated from the fit to data (rather than specified as shown here). The covariance $\boldsymbol{\Sigma}$ is then used to generate a probabilistic prediction of missing trait values for each taxon.
each trait is a node and linkages are a directed edge, such that e.g., $Y_{1} \rightarrow Y_{2}$ indicates that a change in trait $Y_{1}$ will cause a subsequent change in $Y_{2}$. The value of path coefficients can then be estimated as fixed effects by identifying their values that maximize the likelihood of data. This use of SEM then allows a user to replace the $n_{j}\left(n_{j}+1\right) / 2$ parameters in a covariance matrix with any set of parameters (from 1, up to the maximum of $n_{j}\left(n_{j}+1\right) / 2$ when not
using Bayesian priors). For example, a trait-imputation model with $n_{j}=30$ traits would require estimating 465 parameters for covariance $\boldsymbol{\Sigma}$ without other constraints, but could be restricted to fewer important parameters using SEM. Furthermore, SEM can be used to estimate the correlation between two traits that are connected by a directed edge ('direct pathways') or mediated by a third trait (called 'indirect pathways'). In this way, SEM decomposes the
correlation between two traits into the contribution from both direct and indirect trait effects.

In this study, we extend PTI to (1) incorporate both continuous and categorical traits, and (2) represent the trait covariance matrix using SEM, while using a Brownian motion model for simplicity of presentation. The approach can be implemented for any traits of floristic or faunistic species, and either using phylogeny for evolutionary distance or using taxonomy as a proxy for relatedness. To demonstrate the benefits of our extensions of PTI, we applied the approach to fishes which have evolutionary trade-offs that are highly structured by temperature and individual length, while also having extensive information about a variety of behavioural, reproductive, and life-history traits (Barnett et al., 2019). We specifically use data for 34 traits for $>32,000$ described fishes, obtained by combining existing in situ trait data (FishBase; Froese, 1990) and morphometric trait data from National Museum of Natural History fish specimens (Price et al., 2019, 2022). We interpret results by computing the direct and indirect impacts of temperature and maximum body length on other traits, and using traits to classify fishes into life-history strategies. Finally, we discuss how phylogenetic imputation of mixed traits using SEMs can help to unify experimental (micro-evolutionary) and comparative (macro-evolutionary) studies of life-history trade-offs.

## 2 | MATERIALS AND METHODS

We extend existing PTI methods in the following two ways:

1. Structural equation modelling: We model the covariance $\boldsymbol{\Sigma}$ among multiple traits using methods derived from SEM. This allows us to specify a small set of path coefficients, despite conducting multivariate trait imputation on many traits.
2. Including categorical traits: We fit our phylogenetic model to a mixture of continuous and categorical traits. Fitting to a categorical trait with $M$ levels involves estimating $M$ - 1 latent variables, and we transform these to the probability of each level using a multivariate logistic transformation given the constraint that these probabilities sum to one. We then model the association between these M-1 latent variables and other continuous traits in a way that permits efficient statistical inference.

We provide further details below (see Supporting Information A for summary of all notation), and implement the approach in the package FIsHLIFE release 3.0.1 (Thorson, 2023) in the R statistical environment (R Core Team, 2021).

## 2.1 | Overview of phylogenetic structural equation modelling

We seek to estimate a vector of traits $\boldsymbol{\beta}_{g}$ for each taxon $g$ in a rooted and additive tree (i.e., including ultrametric phylogenies),
including trait-values for both tips (species) and ancestral nodes as well as the trait-vector $\boldsymbol{\beta}_{0}$ for the root of the tree. We assume that evolution follows a standard model (e.g., Brownian motion, Pagel's lambda, etc) that can be expressed using a multivariate normal distribution (Paradis, 2012). This model allows calculating a correlation matrix $\mathbf{R}$ with dimension $n_{g} \times n_{g}$, where $n_{g}$ is the total number of taxa (tips and ancestral nodes), representing the correlation for a single trait along the phylogeny. We similarly construct the covariance $\boldsymbol{\Sigma}$ among $n_{j}$ traits using methods drawn from structural equation modelling.

This then results in a separable covariance for $\mathbf{B}$ containing latent trait $\beta_{g, j}$ all taxa $g$ and traits $j$ :

$$
\begin{equation*}
\operatorname{vec}(\mathbf{B})^{\sim} M V N\left(\operatorname{vec}\left(\mathbf{1} \otimes \boldsymbol{\beta}_{0}\right), \mathbf{R} \otimes \boldsymbol{\Sigma}\right) \tag{1}
\end{equation*}
$$

where $\mathbf{R} \otimes \boldsymbol{\Sigma}$ is the Kroenecker ('outer') product of the correlation among taxa and covariance among traits, 1 is a vector of 1 s with length $n_{i}$ such that $\mathbf{1} \otimes \boldsymbol{\beta}_{0}$ forms the intercept for every taxon and trait, and MVN is a multivariate normal distribution with these moments. This separable covariance $\mathbf{R} \otimes \boldsymbol{\Sigma}$ can often be implemented more efficiently in some software as a conditional or simultaneous autoregressive model (Ver Hoef et al., 2018), although we present the separable covariance here to agree with standard notation in phylogenetic comparative methods (e.g., Paradis, 2012). In the following we only explore a Brownian motion (a.k.a. random-walk) process for $\mathbf{R}$, although future software developments could easily generalize this to OrnsteinUhlenbeck, Pagel's delta, or other evolutionary models (see Supporting Information B).

We next introduce how to construct trait covariance $\boldsymbol{\Sigma}$ using methods drawn from structural equation modelling. We assume that the user specifies:

1. the structure of a path matrix $\boldsymbol{\Gamma}$ with dimension $n_{j} \times n_{j}$. The user specifies a priori which elements of this matrix are fixed at zero or are instead freely estimated as fixed effects (including cases when multiple path coefficients are constrained to the same estimated value). For example, specifying that $\gamma_{j, j^{*}}=0$ involves assuming that trait $j$ has no direct impact on trait $j^{*}$.
2. a Cholesky matrix $\mathbf{S}$ where $\mathbf{S S}^{t}$ represents the covariance in exogenous variation with dimension $n_{j} \times n_{j}$. At a minimum, this covariance $\mathbf{S S}^{t}$ involves estimating diagonal entries, $\operatorname{diag}(\mathbf{S})=\left(\sigma_{1}, \sigma_{2}, \ldots, \sigma_{n_{j}}\right)$ resulting in an independent exogenous variance $\sigma_{j}^{2}$ for each trait $j$ (where these can again be constrained to the same estimated value). However, traits can also have exogenous covariance by estimating lower-triangle elements of $\mathbf{S}$, which then results in off-diagonal elements for exogenous covariance V. Nonzero elements of $\mathbf{S}$ are then freely estimated as fixed effects.

This path matrix (and resulting path diagram) is central to structural equation modelling, which has been reviewed elsewhere for describing interaction networks and physiological performance (Frauendorf et al., 2021; Garrido et al., 2022). However, structural equation models have not to our knowledge been fitted
simultaneously with phylogenetic covariance. Previous studies have either adjusted data or estimated residual covariance based on phylogeny and then fitted a SEM to those residuals (Mason et al., 2016; Santos, 2012) or have fitted a series of phylogenetic linear models to represent dependencies in a path diagram (van der Bijl, 2018). Specifying a path diagram requires enumerating the set of variables (graph vertices) and dependencies (directed edges), where these dependencies can be interpreted as mechanisms for causal inference (Pearl, 2009). The reliability of causal inference requires correct specification of the path diagram (Grace \& Irvine, 2020), and we recommend further simulation and case-study evaluation of causal inference within phylogenetic comparative methods.

These two matrices are then used to solve a simultaneous equation for $\mathbf{x}^{\sim} \operatorname{MVN}(\mathbf{0}, \boldsymbol{\Sigma})$, i.e., a hypothetical draw from covariance among traits $\mathbf{\Sigma}$ (Kaplan, 2001):

$$
\begin{gather*}
\mathbf{x}=\boldsymbol{\Gamma} \mathbf{x}+\boldsymbol{\varepsilon} \\
\varepsilon^{\sim} \operatorname{MVN}\left(0, S^{t}\right) \tag{2}
\end{gather*}
$$

where $\boldsymbol{\Gamma}$ represents endogenous mechanisms linking variables and $\boldsymbol{\varepsilon}$ represents exogenous variation with variance $\operatorname{Var}(\boldsymbol{\varepsilon})=\mathbf{S S}^{t}$. We then solve for the Cholesky of trait covariance as:

$$
\begin{equation*}
\mathrm{L}=(\mathbf{I}-\boldsymbol{\Gamma})^{-1} \mathbf{S}, \tag{3}
\end{equation*}
$$

where trait covariance $\mathbf{\Sigma}=\mathbf{L L}^{\mathbf{t}}=\operatorname{Var}(\mathbf{x})$.
Constructing trait covariance $\mathbf{\Sigma}=(\mathbf{I}-\boldsymbol{\Gamma})^{-1} \mathbf{S} \mathbf{S}^{t}\left(\mathbf{I}-\mathbf{\Gamma}^{t}\right)^{-1}$ in this way generalizes several existing models:

1. Brownian motion: The analyst might specify $\boldsymbol{\Gamma}=\mathbf{0}$ and $\mathbf{S}$ as a diagonal matrix, and this then reduces to a standard Brownian motion model.
2. Phylogenetic path analysis: In some cases, variables can be reordered such that $\boldsymbol{\Gamma}$ is lower-triangular. In these cases, the model can be estimated using phylogenetic path analysis, for example fitted using piecewise SEM or d-separation methods (van der Bijl, 2018; von Hardenberg \& Gonzalez-Voyer, 2013). However, $\boldsymbol{\Gamma}$ might also include loops, where for example, trait $j_{1}$ affects $j_{2}, j_{2}$ affects $j_{3}$, and $j_{3}$ affects $j_{1}$. This cannot be represented using standard phylogenetic path analysis but can be using SEM (Equations 2 and 3).
3. Phylogenetic factor analysis: In other cases, the analyst might specify $\boldsymbol{\Gamma}=\mathbf{0}$ and $\mathbf{S}$ having lower-diagonal entries that are nonzero for only a few columns. In this case, $\mathbf{\Sigma}=\mathbf{S S}^{t}$ where the nonzero columns of $\mathbf{S}$ represent 'factors loadings' in a phylogenetic factor analysis (Hassler et al., 2022; Thorson et al., 2017).
In general, covariance $\boldsymbol{\Sigma}$ among $n_{j}$ traits involve $n_{j}\left(n_{j}+1\right) / 2$ moments, and the analyst can specify anywhere from one to $n_{j}\left(n_{j}+1\right) / 2$ parameters within the two matrices $\boldsymbol{\Gamma}$ and $\boldsymbol{S}$. To simplify the user-interface, we require the user to specify linkages as a text file following the format of R-package sem (Fox et al., 2020), and then parse this text file to construct $\boldsymbol{\Gamma}$ and $\mathbf{S}$ from a vector of estimated parameters.

## 2.2 | Fitting both continuous and categorical variables

We next outline how this model is fitted to a set of $n_{c}$ continuous and $n_{d}$ categorical traits, for a total of $n_{t}=n_{c}+n_{d}$ traits. This has been done previously using a 'threshold model' to combine categorical and continuous traits (e.g., Cybis et al., 2015; Felsenstein, 2012; Tolkoff et al., 2018), although we instead fit categorical traits using a Categorical distribution based on estimated probabilities for each categorical level (similar to Hadfield \& Nakagawa, 2010). These traits are assembled in a matrix $\mathbf{Y}$ with dimension $n_{i} \times n_{t}$, where missing values are recorded as NAs and are excluded when computing the likelihood across available data. We also record the number of levels $m_{t}$ for each trait $t$, where categorical traits have $m_{t} \geq 2$ by definition and we adopt the convention that $m_{t}=1$ for continuous traits. Categorical traits are modelled via a probability vector that is constrained to sum to 1 , so it requires $m_{t}-1$ variables to describe a categorical trait with $m_{t}$ levels. For trait-matrix $\mathbf{Y}$ with $m_{t}$ levels for each trait $t$, we therefore must estimate latent trait matrix $\mathbf{B}$ with $n_{j}=n_{c}+\sum_{t=1}^{n_{t}}\left(m_{t}-1\right)$ columns and $n_{g}$ rows (where $n_{g}$ is the total number of taxa in the tree). We also define a vector $h$ with length $n_{j}$, where $h_{j} \in\left\{1,2, \ldots, n_{t}\right\}$; this vector associates each column of $\mathbf{B}$ with a corresponding column of $\mathbf{Y}$. If trait $t$ is continuous then only one value of $h_{j}=t$. Alternatively, if trait $t$ is categorical then $h_{j}=t$ for $m_{t}-1$ elements. Finally, we associate $n_{i}$ rows of $\mathbf{Y}$ with $n_{g}$ rows of $\mathbf{B}$ by defining a vector $\mathbf{g}$ with length $n_{i}$ where $g_{i}$ provides the taxon associated with sample $i$. The process of fitting latent traits $\mathbf{B}$ to trait measurements $\mathbf{Y}$ differs somewhat between continuous and categorical traits, as we explain next.

For a continuous trait $t$, we extract column $\mathbf{y}_{t}$ from $\mathbf{Y}$. We also extract the column from $\mathbf{B}$ for which $h_{j}=t$ and call this submatrix $\mathbf{B}^{(t)}$. We then specify a normal distribution for residual (measurement) variation:

$$
\begin{equation*}
y_{i, t} \sim \operatorname{Normal}\left(\beta_{g_{i}, 1}^{(t)}, \sigma_{j}^{2}\right) \tag{4}
\end{equation*}
$$

where $\sigma_{j}^{2}$ is the magnitude of measurement errors and is estimated as a fixed effect, although we fix $\sigma_{j}=0.01$ (i.e., forcing $\beta_{g_{i, j}}$ to approach $y_{i, j}$ ) for any trait $j$ that does not have replicated measurements and hence cannot estimate $\sigma_{j}^{2}$.

For a categorical trait $t$, we again extract column $\boldsymbol{y}_{t}$ from $\mathbf{Y}$. However, we then expand $\mathbf{y}_{t}$ to an indicator matrix $\mathbf{Z}^{(t)}$ with dimension $n_{i} \times m_{t}$, such that a trait with $m_{t}$ possible levels is converted to a matrix with $m_{t}$ columns where each row $i$ contains a 1 in the column corresponding to level $y_{i, t}$ and zeros otherwise. We also extract the $m_{t}-1$ columns from $\mathbf{B}$ for which $h_{j}=t$ and again call this submatrix $\mathbf{B}^{(t)}$. We calculate the probability $\boldsymbol{\pi}_{g}^{(t)}$ for each level $k \in\left\{1,2, \ldots, m_{t}\right\}$ of categorical trait $t$ via a multivariate logistic transformation of each row $\boldsymbol{\beta}_{g}^{(t)}$ of $\mathbf{B}^{(t)}$ :

$$
\begin{equation*}
\pi_{g, k}^{(t)}=\frac{\frac{e^{f_{3 k}}}{f_{k}^{(t)}}}{1+\sum_{k_{1}=1}^{m_{t}-1} e^{f_{s k^{\prime}}^{(t)}}} \quad \text { if } k \leq m_{t}-1 . \tag{5}
\end{equation*}
$$

This multivariate logistic transformation converts $m_{t}-1$ unbounded values in $\boldsymbol{\beta}_{g}^{(t)}$ to $m_{t}$ probabilities $0<\boldsymbol{\pi}_{g}<1$ where $\sum_{k=1}^{m_{t}} \pi_{g, k}=1$ by construction. We then fit a categorical distribution:

$$
\begin{equation*}
\mathbf{z}_{i}^{(t) \sim} \sim \text { Categorical }\left(\pi_{g(i)}^{(t)}\right) \tag{6}
\end{equation*}
$$

This differs from previous specifications of a 'threshold model' to predict categorical traits, which have typically predicted a response of $z_{i, k}=1$ whenever a 'liability' variable $\beta_{g(i), j}$ exceeds an estimated threshold and zero otherwise (e.g., Felsenstein, 2012). Such a threshold model must integrate across values of $\beta_{g(i), j}$ that fall on the right side of a given threshold for a measurement $y_{i, j}$, typically accomplished using Bayesian hierarchical models and MCMC sampling. By contrast, we specify that latent traits $\boldsymbol{\beta}_{g}^{(t)}$ for each taxon $g$ are transformed to the probability $\boldsymbol{\pi}_{g}^{(t)}$ for each level of a categorical variable.

Parameters of this model remain identifiable given missing data (i.e. entries of $y_{i, t}=N A$ ). In these cases, the model continues to integrate across latent variables $\mathbf{B}$, and simply does not include these missing values of $\mathbf{Y}$ in the likelihood. We note that we assume trait measurements $\mathbf{Y}$ are missing at random. If the probability of having an available trait measurement (termed 'sampling intensity') is correlated with latent traits $\boldsymbol{\beta}_{j}$, then this assumption will result in 'preferential sampling' bias (Diggle et al., 2010). We recommend further research regarding model-based mitigation of this bias (e.g., Conn et al., 2017), but do not explore the topic further here.

## 2.3 | Parameter estimation and interpretation

We identify maximum likelihood estimates for all model parameters (see Supporting Information B for estimation details). This requires calculating an objective function as the product of the likelihood (Equation $4 / 6$ ) and the probability of random effects (Equation 1). We obtain the marginal likelihood by integrating the objective function across random effects $\mathbf{B}$, composed of random effects $\beta_{g, j}$ for all taxa $g$ (including tips and ancestors) and traits $j$. This multivariate integral is approximated using the Laplace approximation and implemented using R -package тмв, and this is computationally efficient because the inverse-covariance $(\mathbf{R} \otimes \boldsymbol{\Sigma})^{-1}$ has a value of 0 for any two taxa that are not adjacent in the specified tree (Kristensen et al., 2016). We then maximize the marginal likelihood with respect to remaining fixed effects ( $\boldsymbol{\Gamma}, \mathbf{S}, \boldsymbol{\beta}_{0}$, and $\boldsymbol{\sigma}^{2}$ ), export the estimate of SEM-coefficients $\boldsymbol{\Gamma}$ and $\mathbf{S}$, extract 'empirical Bayes' predictions for latent traits B (which includes imputed values for missing trait values), and use r-package sem to visualize the estimated path diagram.

Path coefficients $\boldsymbol{\Gamma}$ can be interpreted as a regression slope, but the precise interpretation depends upon the transformation that was chosen by the analyst for connected variables $Y_{1} \rightarrow Y_{2}$. For example, if $Y_{1}$ is untransformed (e.g. temperature in Celsius) and $Y_{2}$ is log-transformed (e.g., log-maximum body length), then e.g., $\gamma_{1,2}=0.1$ indicates that a 1 Celsius increase in $Y_{1}$ is associated on average with a $10 \%$ increase in $Y_{2}$. By contrast, if $Y_{1}$ is log-transformed (e.g. log-maximum body length), and $Y_{2}$ and $Y_{3}$ are two levels of a
categorical variable, then $\gamma_{1,2}=0.1$ and $Y_{1,3}=-0.1$ indicates that a $10 \%$ increase in maximum body length is associated on average with a $e^{0.1(0.1)} / e^{0.1(-0.1)}=2 \%$ increase in the odds of level $Y_{2}$ relative to level $Y_{3}$. We also note that the covariance among traits $\boldsymbol{\Sigma}$ is estimated as being constant across the entire phylogenetic tree (i.e., that $\operatorname{Var}(\mathbf{B})=\mathbf{R} \otimes \mathbf{\Sigma}$ ). In reality, slope and variance parameters may be nonstationary, representing different evolutionary trade-offs and rates resulting from environmental context and ecological traits that are not being modelled. We recommend further research extending the approach to include nonstationarity, and interpret parameters in this study as representing a sample-weighted average across the tree being analysed.

## 2.4 | Case study: Estimating life-history traits of fishes

To test and apply these methodological advances, we seek to estimate life-history traits for all described fishes (Chondrichthyes and Osteichthyes) included in FishBase in November 2019, where previous research has validated that these data are likely unbiased (Thorson et al., 2014). There is no phylogeny available for all fishes, despite ultrametric phylogenies existing separately for a subset of bony (Rabosky et al., 2018) and cartilaginous fishes (Stein et al., 2018). We therefore follow past research (Johnson et al., 2021; Thorson et al., 2017) in approximating phylogeny via taxonomy, that is, where all taxonomic classes are assumed to have a single common ancestor, and then including ancestral levels for order, family and genus. Package FishLife then converts taxonomy to a tree using Rpackage APE (Paradis \& Schliep, 2019), and when using taxonomy we specify phylogenetic distance $d_{g}=1$ for each level of the taxonomic tree (i.e., for family to genus, genus to species, etc). We later provide a sensitivity analysis with a novel merged phylogeny.

We analyse 17 continuous-valued traits and four categorical traits, where the latter include 16 levels in total. These trait data include at least one measurement for 26,622 fish species. However, life-history data in particular are missing for many species (Figure B1), where $2 \%-27 \%$ of species have at least one measurement of a given trait related to growth, mortality, or body size. These 'inclusion rates' are higher for genera ( $7 \%-24 \%$ ), and family levels (26\%-76\%), suggesting that phylogenetic information is necessary to infer trait-values for many species based on their genus or family.

We classify these 33 variables into six trait categories, expanding upon the list from Hadj-Hammou et al. (2021) where traits are broadly classified into five categories: (1) behaviour, (2) life history, (3) morphology, (4) diet and (5) physiology. The list includes at least one variable in each category (see Table 1 for details). The morphometric traits are composed of continuous measures of body shape traits that describe overall body shape for 5940 extant species of actinopterygian fishes spanning 392 families, taken on specimens at the Smithsonian Museum of Natural History and averaged by species (Price et al., 2019, 2022). These data include eight linear measurements in three dimensions: standard body and jaw length; head,
body, and caudal peduncle depth; and body, jaw, and caudal peduncle width. We standardized specimen morphometrics to account for variation in individual development for museum specimens, by dividing each measurement by the geometric mean of specimen length, width, and height.

We use several design principles to assemble the SEM for fishes, and this in turn defines the structure of SEM coefficients $\boldsymbol{\Gamma}$ and exogenous covariance V. Specifically we specify that:

1. temperature (in Celsius) is the exogenous 'root' of the path diagram. This recognizes that life-history studies typically use temperature as a covariate to predict size and mortality
(Gislason et al., 2010; Palomares et al., 2022; Pauly, 1980), and hence our estimates are comparable to widely reported slopes.
2. von Bertalanffy length $\left(L_{\infty}\right)$ in units $m m$ has the greatest number of impacts on other traits, in recognition of the central role of body size in size-structured evolutionary theory (Andersen, 2019). Von Bertalanffy length is the asymptotic body size of a fish. We include linkages to other measurements of size (in mm or g), growth (in year ${ }^{-1}$ ), and mortality parameters (in units year ${ }^{-1}$ ), as well as to categorical traits representing reproductive behaviour, feeding mode, and habitat (Denéchère et al., 2022; Palomares et al., 2022).

TABLE 1 Life-history traits included in the analysis, listing the variable name, trait category (using five defined by Hadj-Hammou et al. (2021) while also adding 'Reproductive' as a sixth category), whether the trait is continuous or categorical, the transformation applied to continuous variables achieve a close-to-normally distributed process for evolutionary changes, and the levels for factor-valued traits.

|  |  | Continuous <br> (C) or | Transformation <br> (if continuous) | Levels (if <br> factor-valued) |
| :--- | :--- | :--- | :--- | :--- |
| Name | Trait category | categorical (F) | Natural log | - |
| age_max | Life-history | C | Identity | - |
| trophic_level | Diet | C | Natural log | - |
| aspect_ratio | Morphology | C | Natural log | - |
| fecundity | Reproduction | C | Natural log | - |
| growth_ <br> coefficient | Physiology | C | Identity | - |
| temperature | Physiology | C | Natural log | - |
| length_max | Physiology | C | Natural log | - |
| length_infinity | Physiology | C | Natural log | - |
| length_maturity | Physiology | C | Natural log | - |
| age_maturity | Physiology | C | Natural log | - |
| natural_mortality | Physiology | C | Natural log | - |
| weight_infinity | Physiology | C | Natural log | - |
| max_body_depth | Morphology | C | Natural log | - |
| max_body_width | Morphology | C | Natural log | - |
| lower_jaw_length | Morphology | C | Natural log | - |
| min_caudal_ | Morphology | C |  |  |

3. both growth and mortality rates affect age and length at maturity, in recognition that their ratio affects the optimal maturation timing (Holt, 1958).
4. for each categorical trait $t$ (i.e., all columns $j$ of $B$ where $h_{j}=t$ ), the exogenous covariance $\mathbf{V}$ is symmetric and positive definite but otherwise unconstrained (i.e., the body-shape trait has five measured levels and involves estimating $\frac{4 \times 5}{2}=10$ covariance parameters in $\mathbf{S}$ ), while continuous traits have independent exogenous variance (i.e., $\mathbf{S}$ is diagonal for these rows and columns).
Future research could compare fit with alternative assumptions about life-history trade-offs (e.g., Mason et al., 2016).

### 2.4.1 | Sensitivity, validation and performance

We assess the performance of the model, validate results, and explore sensitivity to alternative assumptions using several auxiliary analyses.

First, we compare phylogenetic structural equation modelling with the r-package phylolm (Tung Ho \& Ané, 2014) as widely used example of standard phylogenetic comparative methods (Supporting Information D). We specifically compare model structure, and also using a short simulation experiment with 500 replicates to confirm that the phylogenetic SEM can generate identical estimates of regression coefficients to an existing phylogenetic linear model package. For each replicate, we simulate an additive tree with 100 'tips' and randomized branch lengths and structure. We then simulate two variables under a Brownian motion model from this tree, exploring scenarios either with complete data for each taxon, or $60 \%$ of taxa missing measurements for each trait. We record the estimated slope parameter for these two models.

We also assess sensitivity of results to using taxonomic information as a proxy for evolutionary relatedness. To do so, we first merge publicly available chondrichthyan (Stein et al., 2018) and actinopterygian (Rabosky et al., 2018) ultrametric trees, using branch lengths to infer the location of their common ancestor. We then subset our data to the 11,070 species that can be matched between trait data and the merged phylogeny, and repeat the analysis on this subset. Subsetting to these matched species reduces the number of available trait measurements from 246,736 to 152,596 , so we present these estimates using phylogenetic information as a sensitivity analysis.

Next, we validate the predictive performance of the model by conducting a 4 -fold cross-validation experiment. To do so, we randomly partition each row of original data matrix $\mathbf{Y}$ into one of four bins (labelled $\{A, B, C, D\}$ ). For the first experiment, we then fit the model to all data in bins $\{B, C, D\}$ and use the estimated parameters to predict $\boldsymbol{\beta}_{t}$ for continuous traits and level probabilities $\boldsymbol{\pi}_{g}^{(t)}$ for categorical traits corresponding to data in bin $A$. We record these and then repeat this process for the other three bins, comparing predictions with the withheld data. This experiment evaluates performance when predicting new data that are collected via the same process as the original data set (Roberts et al., 2017), and we recommend future research use a blocked cross-validation design to
explore performance when predicting traits for taxa that are systematically under-represented in available data.

We then evaluate performance separately for continuous and categorical traits:

- Continuous traits: for continuous trait $t$, we plot unfitted observations $\boldsymbol{y}_{t}$ against the out-of-bag predictions $\boldsymbol{\beta}_{j}^{(t)}$ (where $\boldsymbol{\beta}_{j}^{(t)}$ is the column of $\mathbf{B}$ for which $h_{j}=t$ ), and also calculate the percent variance explained relative to a null model that predicts $\boldsymbol{y}_{t}$ based on its mean value $\bar{y}_{t}$ :

$$
\begin{equation*}
P V E_{t}=1-\frac{\sum_{i=1}^{n_{i}}\left(y_{i, t}-\beta_{g(i), j}^{(t)}\right)^{2}}{\sum_{i=1}^{n_{i}}\left(y_{i, t}-\bar{y}_{t}\right)^{2}} \tag{7}
\end{equation*}
$$

$P V E_{v}$ predicts the proportion of variance that would be explained for a hypothetical 'new' sample, where a value of 0 indicates no out-of-bag explanatory power (i.e. no improvement relative to predicting new samples as the mean of all data) and a value of 1 implies perfect explanatory power.

- Categorical traits: for latent trait $\boldsymbol{\beta}_{j}^{(t)}$ representing a level of a categorical trait, remember that we expand original data $\mathbf{y}_{t}$ to an indicator matrix $\mathbf{Z}^{(t)}$ where $\mathbf{z}_{k}^{(t)}$ is the column corresponding to level $k$ of latent trait $t$. This indicator column has value 0 when a taxon does not have that level and 1 when it does, while the model estimates the probability $\pi_{k}^{(t)}$ for that level of the categorical trait, and these probabilities sum to one across levels $k \in\left\{1,2, \ldots, m_{t}\right)$. To evaluate model performance, we plot the receiver operator characteristics (ROC) curve for each level, which involves calculating the rate of false-positives and falsenegatives when converting the predicted probability to a predicted indicator using different potential threshold values. We then calculate the area under the ROC (AUC) using $R$ package pROC (Robin et al., 2011), where an AUC of 0.5 indicates no out-of-bag ability to discriminate between 0 and 1 values for an indicator, and an AUC of 1 implies perfect ability to discriminate between these.


### 2.4.2 | Identifying life-history strategies

We illustrate results by identifying a small number of life-history strategies for fishes, defined as an extreme combination of trait values that frequently occur together, such that all fishes can be characterized as some mixture of strategies (i.e., following the usage in Winemiller \& Rose, 1992). Previous studies have applied clustering methods to a smaller subset of species than we have available, e.g., for North American fishes (Winemiller \& Rose, 1992), selected North Pacific marine fishes (King \& McFarlane, 2003), freshwater fishes (Mims et al., 2010), or European marine fishes (Pecuchet et al., 2017). However, our study is the first to predict the life-history strategies for all described fishes worldwide, representing more than 34,000 species.

We specifically follow Winemiller and Rose (1992) in estimating 'archetypes' that represent an extreme combination of lifehistory characteristics. All fish taxa are then described as a finite mixture of these archetypes, and we refer to these archetypes as 'life-history strategies'. This contrasts with other studies that have clustered taxa continuously within the space of life-history traits (King \& McFarlane, 2003). To do so, we extract predictions $\boldsymbol{\beta}^{(t)}$ for continuous traits and level probabilities $\boldsymbol{\pi}_{g}^{(t)}$ for categorical traits for all taxa that have at least one observation (i.e., are not purely drawn from the predictive distribution based on its taxa). We then apply 'archetypal analysis' (Cutler \& Breiman, 1994) following methods from Pecuchet et al. (2017), using R package archetypes (Eugster \& Leisch, 2009). Archetypal analysis involves estimating $n_{b}$ 'archetypes' $\boldsymbol{\alpha}_{b}$ composed of values $\alpha_{b, j}$, representing the value of variable $j$ in archetype $b$. Each taxon $\boldsymbol{\beta}_{g}$ is then predicted as a finite mixture of these archetypes, with mixture coefficients $p_{g, b}$ defined such that $\sum_{k=1}^{n_{k}} p_{g, b}=1$ and $p_{g, b}>0$. Archetypal analysis then estimates the value of $\alpha_{g, b}$ and $p_{b, j}$ to minimize the sum of squared distance (SSD) between predicted and inputted $\mathbf{B}$. We use a scree-plot to visualize how the SSD decreases when using 1-6 archetypes and we select the number by visually identifying when further increases generate little improvement in SSD. We then explore the results in two ways:

- Archetype trait values: We extract trait values for estimated archetypes, $\boldsymbol{\alpha}_{b}$, to interpret which traits are associated with each. We specifically convert $\alpha_{b, j}$ to a percent score $\alpha_{b, j}^{*}$ by calculating the proportion of fishes having a predicted trait $\boldsymbol{\beta}_{j}<\alpha_{b, j}$ :
- Simplex by taxonomy: Similarly, we extract mixture coefficients $\mathbf{p}_{g}$ for each taxon. We then use package archetypes to apply a skew-orthogonal transformation to visualize $p_{g, b}$ in a two dimensional simplex (Seth \& Eugster, 2014). We specifically compare $p_{g, b}$ for major taxa and compare resulting assignments with previous studies (Winemiller \& Rose, 1992).


## 3 | RESULTS

The phylogenetic structural equation model quantified the direct impact of temperature on size and growth. Specifically, a one degree Celsius increase was associated with a $4 \%$ increase in growth coefficient (with standard error $S E=0.3 \%$ ), $2 \%$ increase ( $\mathrm{SE}=0.2 \%$ ) in mortality rate, and $2 \%$ decrease ( $\mathrm{SE}=0.3 \%$ ) in asymptotic body length (Figure 2; Table E1), where these represent average associations across the wide range of fishes being analysed. In turn, a $10 \%$ increase in asymptotic body length was associated with an $8.2 \%$ ( $\mathrm{SE}=0.3 \%$ ) decrease in natural mortality and a $6.6 \%$ decrease ( $\mathrm{SE}=0.2 \%$ ) in growth coefficient. When both direct and indirect effects are included (Table E2), temperature had a slightly larger impact on the growth coefficient $(0.051)$ than on the mortality rate $(0.035)$. Temperature was estimated to have a minimal effect on reproductive behaviour, feeding mode, or spawning type, while asymptotic length had a larger effect on these traits (Table E2). For example, a $10 \%$ increase in asymptotic length was estimated to decrease the
odds of guarding behaviour relative to non-guarding behaviour by $34 \%$ (Table E2). Finally, the model also captured previously documented life-history trade-offs, including the association between earlier maturation and higher relative mortality (Figure E1).

The simulation experiment confirmed that FishLife and the widely used R -package PHYLolm give essentially identical estimates when fitting continuous traits and data are available for all species (Figure D1, left panel), and that FishLife shows a small improvement in estimation performance when data are missing at random (Figure D1, right panel). Four-fold cross-validation confirmed that the model fitted to real-world data had good performance when predicting records that were randomly dropped from the model fitting (Figure 3). Continuous-valued traits had a percent-variance explained (PVE) ranging from $51 \%$ to $89 \%$. Among these variables, performance was particularly high (>80\% PVE) for traits measuring length, weight, and fecundity, but lower for traits measuring age, growth, maturity, and trophic level. Similarly, levels of categorical traits had an area under the receiver-operator-characteristics curve (ROC) ranging from 0.86 to 0.99 , with lower (but still high) power to discriminate levels for the feeding-mode trait. Comparing the model fitted using taxonomy with one using a subset of data and phylogeny to represent evolutionary distance (Figure E2) shows similar estimates of linkages for life-history parameters (i.e., for mortality, growth, size, and maturity parameters) between analyses. However, the estimated impact of body size on body shape was substantially larger when using phylogeny.

Our approach is further demonstrated by the archetype analysis, which identified three life-history strategies (Figure E2), in agreement with Winemiller and Rose (1992). The first archetype (purple in Figure 4 and top panel in Figure 5) was associated with higher maximum age, trophic level, slow growth, and low temperatures. This suite of traits corresponded to the 'equilibrium' strategy from Winemiller and Rose (1992). The third archetype (yellow in Figure 4 and bottom panel in Figure 5) corresponded to the opportunistic strategy from Winemiller and Rose (1992). It had the lowest maximum age and fecundity, while having high natural mortality and probability of guarding their young. Finally, the second archetype was somewhat intermediate in terms of growth and size, while typically having highest fecundity, being mainly pelagic and having the highest probability of a non-guarding reproductive strategy. As expected, there was strong phylogenetic signal in these life-history strategies, with Elasmobranchii (sharks and rays) representing the equilibrium strategy, Clupeidae (herrings and sardines) largely representing the periodic strategy, and Gobiidae (gobies) largely representing the opportunistic strategy (Figure 6).

## 4 | DISCUSSION

We extended phylogenetic trait-imputation methods to include two additional features: (1) representing the covariance among traits via a structural equation model, and (2) incorporating both continuous and categorical traits. We fit categorical traits using latent variables


FIGURE 2 Path diagram representing specified causal linkages and estimated $\boldsymbol{\Gamma}$ coefficients (see Figure 1 for description) linking fish traits when using taxonomy to represent evolutionary distance, using package sem to generate the plot (Fox et al., 2020), where levels of the categorical variables are abbreviated (H: habitat; FM: feeding mode; BS: body shape; ST: spawning type) and coefficients for categorical variables represent the log-odds relative to a specified base level (H: demersal; FM: generalist; BS: fusiform/normal; ST: nonguarders). Note that evolutionary variance and covariance parameters $\boldsymbol{\Sigma}$ are not shown here for clarity of presentation.
that are then transformed to calculate the probability for each categorical level. Unlike past analyses (e.g. Felsenstein, 2012), however, we use a computational method (the Laplace approximation) that allows rapid inference on large trees. These two developments have wide relevance for applications across life-history databases for any taxonomic group within various ecosystems worldwide such as plants, mammals, fishes, birds, insects, as well as comparing across these taxa (Capdevila et al., 2020). For example, Kattge et al. (2011) documented 52 traits for 69,000 plant species in the TRY global plant database, of which 15 are categorical including Mycorrhiza type, nitrogen fixation capacity, and pollination mode. In addition, GRooT (Guerrero-Ramírez et al., 2021) includes 38 root
traits, from 38,276 species-by-site mean values based on 114,222 trait records, for more than 1000 species, such as root mass fraction, root carbon and nitrogen concentration, lateral spread, root mycorrhizal colonization intensity, mean root diameter, root tissue density, specific root length and maximum rooting depth. Similarly, the bird trait database AvoNET (Tobias et al., 2022) includes continuous morphological traits but also categorical traits like trophic level (three levels), foraging niche (nine levels) and foraging locomotory behaviour (five levels) for 11,009 species. Likewise, the foraging database EltonTraits (Wilman et al., 2014) includes foraging time as a categorical trait for 9993 bird and 5400 mammal species. Clearly there is potential for both phylogenetic signal within these


FIGURE 3 Evaluating predictive performance for all variables based on a four-fold cross-validation experiment. For continuous-valued traits, plots show the held-out value ( $x$-axis) against the predicted value ( $y$-axis), along with the one-to-one line (black line) and list the percent-variance-explained (PVE). A well-performing model will have predictions near the one-to-one line and a PVE approaching 100\%. For discrete-valued traits, we used the held-out factor-level indicator ( 0 or 1 ) and the predicted class probability to calculate the receiveroperator characteristics curve (ROC). A well-performing model will have ROC in the upper-left corner and an AUC approaching 1.0.


FIGURE 4 Frequency distribution ( $y$-axis) for estimated values ( $x$-axis) for each life-history trait (panels) with the trait-value for each of three life-history strategies identified using the 'archetype' analysis (vertical lines; purple: Equilibrium; green: Periodic; yellow: Opportunistic).

FIGURE 5 Illustration of traits associated with each estimated lifehistory strategy (equilibrium, periodic and opportunistic), specifically showing the proportion of species with a trait-value lower than that of a given archetype ( $y$ axis) for each trait ( $x$-axis) and archetype (panel).

categorical traits, and ecologically meaningful relationships between qualitative and quantitative traits. This is clearly illustrated in our case study, which estimates that fishes with smaller adult body sizes (a continuous trait) are more likely to guard their young (a categorical trait) in agreement with recent theoretical prediction (Denéchère et al., 2022). Similarly, categorical traits (e.g. reproductive behaviour in fishes, or nitrogen fixation in plants) can be highly relevant when measuring functional diversity or predicting responses to new competitors or climates. Overall, this underlines the importance of including categorical traits when imputing traits for both regional and macroecological studies.

We also argue that SEM will be increasingly attractive for phylogenetic trait imputation as the number of traits increases. This utility arises because phylogenetic trait imputation with $n_{j}$ traits typically requires on the order of $n_{j}^{2}$ parameters for the covariance among traits (Bruggeman et al., 2009), or $n_{j} n_{f}$ parameters when specifying $n_{f}$ factors that represent major axes of covariance among traits (Hassler et al., 2022; Thorson et al., 2017). These approaches scale rapidly with an increase in the number of traits, which becomes prohibitive when there are many traits to consider,
such as in the TRY database version-5 containing 2100 traits. By contrast, SEM allows customized specification of the number of parameters, ranging from 1 (i.e., identical evolution rate for each trait) to $n_{j}\left(n_{j}+1\right) / 2$. Furthermore, path parameters in $\Gamma$ are interpretable as regression slopes, such that individual parameters can be compared with pre-existing theory about trait linkages, whether from field observations or laboratory experiments. In our study for example, we estimate a nearly isometric (2.96) scaling of asymptotic body length and body mass and a linear scaling of asymptotic and maximum length (0.99), and these parameters are easily corroborated when evaluating model plausibility. Indeed, future SEMs could consider fixing these and other parameters a priori to improve parsimony and the resulting precision for difficult-to-estimate trait linkages. Alternatively, we estimate the total (direct and indirect) impact of $\log$ (length) on $\log$ (natural mortality) of -0.82 , and this differs somewhat from the inverse relationship claimed by Lorenzen et al. (2022), such that in some cases it is helpful to test for differences relative to existing theory. Finally, SEM starts by specifying a graph (where nodes represent variables, and edges represent dependencies), which can


FIGURE 6 Illustration of where species (circles) within each family or class (colours, using legend in top-left) fall among three life-history strategies (Per = Periodic; Oppor = Opportunistic; Equil $=$ Equilibrium), noting (1) that Sebastes within Scorpaenidae is expected to be between Periodic and Equilibrium, (2) Etheostorna within Percidae is expected between Equilibrium and Opportunistic,
(3) Salmonidae are expected between Equilibrium and Periodic, and
(4) Gambusia within Poecilidae are expected to be opportunistic (Winemiller \& Rose, 1992). Similarly, Equilibrium fishes are expected to exploit a stable environment while Periodic exploit large-scale and/or seasonal patches (Winemiller \& Rose, 1992), which we illustrate by showing Elasmobranchii and Clupeidae, respectively. Finally, Pecuchet et al. (2017) discussed Gobiidae as an example of the Opportunistic strategy.
be readily derived from existing conceptual or theoretical models for a given taxonomic group. For example, length-structured models for fish evolution have already derived boldness as a function of exogenous changes in mortality rate (Andersen et al., 2018) or temperature (Neubauer \& Andersen, 2019), and future research could adapt these graphical models within multivariate trait imputation.

We suggest that in the future SEM and causal inference may help unite research at disparate scales. Causal inference is often described using graphical models, and if the path diagram accurately describes the real-world processes then the resulting estimates can be interpreted as causal mechanisms (Laubach et al., 2021). Phylogenetic trait imputation using SEM provides new avenues to combine laboratory and natural experiments (micro-evolution) with comparative studies of life-history trade-offs (macro-evolution). This may be particularly important for studies of global change biology. For example, natural experiments in insects suggest that increased temperatures can lead to longer flight season (e.g., a wider range of days where adults are present) (Merckx et al., 2021). This observed relationship could then be supplied when using SEM to conduct phylogenetic trait imputation of an insect database, such as the Odonate Phenotypic Database (Waller et al., 2019). Similarly,
rearing experiments involving artificial harvest of fishes suggests that changes in mortality will negatively impact age-at-maturity (van Wijk et al., 2013). However, these experimental results have not previously been used in comparative analyses of life-history parameters. Ultimately, we hope that the use of SEM in phylogenetic trait imputation will contribute to the ongoing discussion between experimental and observational studies of trade-offs in floristic or faunal traits. Allowing the explicit recognition of the theoretical assumptions implied in phylogenetic trait imputation will provide a new avenue for ecological theory to be applied in community diversity and macroecology.

## AUTHOR CONTRIBUTIONS

James Thorson, Romain Frelat and Aurore Maureaud conceived the ideas and designed methodology; Sarah Friedman, Maria L. D. Palomares, Samantha Price and Peter Wainwright collected the data; Jennifer Bigman merged the publicly available phylogenies for fishes; James Thorson and Aurore Maureaud analysed the data; and James Thorson led the writing of the manuscript. All authors contributed critically to the drafts and gave final approval for publication.

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## CONFLICT OF INTEREST STATEMENT

We have no conflicts of interest to declare.

## DATA AVAILABILITY STATEMENT

All data inputs, parameter estimates, trait predictions, and software are available publicly in R package FISHLIFE (https://github. com/James-Thorson-NOAA/FishLife) using release number 3.0.1 (Thorson, 2023).

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## SUPPORTING INFORMATION

Additional supporting information can be found online in the Supporting Information section at the end of this article.
Supplementary Materials A: List of all notation.
Supplementary Materials B: Constructing the correlation matrix for an additive tree
Supplementary Materials C: Illustrating missing data.
Supplementary Materials D: Comparison with phylogenetic comparative methods.
Supplementary Materials E: Detailed results.

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