



LETTER

Interactive life-history traits predict sensitivity of plants and animals to temporal autocorrelation

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Abstract

Temporal autocorrelation in demographic processes is an important aspect of population dynamics, but a comprehensive examination of its effects on different life-history strategies is lacking. We use matrix population models from 454 plant and animal populations to simulate stochastic population growth rates ($\log \lambda_s$) under different temporal autocorrelations in demographic rates, using simulated and observed covariation among rates. We then test for differences in sensitivities, or changes of $\log \lambda_s$ to changes in autocorrelation among two major axes of life-history strategies, obtained from phylogenetically informed principal component analysis: the fast-slow and reproductive-strategy continua. Fast life histories exhibit highest sensitivities to simulated autocorrelation in demographic rates across reproductive strategies. Slow life histories are less sensitive to temporal autocorrelation, but their sensitivities increase among highly iteroparous species. We provide cross-taxonomic evidence that changes in the autocorrelation of environmental variation may affect a wide range of species, depending on complex interactions of life-history strategies.

Keywords

Environmental variation, fast–slow continuum, life-history strategy, matrix population model, multivariate analysis, population projections, reproductive strategy, seed dormancy, vital rates.

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INTRODUCTION

Most natural populations are exerted to environmental fluctuations (Tuljapurkar 1990; Boyce *et al.* 2006; Morris *et al.* 2008). Such fluctuations typically cause temporal variation in vital rates of individuals (i.e. survival, growth and reproduction), and the effects of such variation on population dynamics have been assessed in a number of theoretical and empirical studies using population models (Tuljapurkar 1990; reviewed in Ehrlén *et al.* 2016). However, temporal variation in environmental conditions has most often been modelled as independent and identically distributed (*i.i.d.*; e.g. Cohen 1979), which assumes an unrealistic lack of temporal autocorrelation in environmental fluctuations (Halley & Inchausti 2004; Ruokolainen *et al.* 2009; Buckley *et al.* 2010).

A better understanding of how autocorrelated environmental perturbations of vital rates may determine current population dynamics is critical for eco-evolutionary questions and conservation management (Tuljapurkar 1982; Metcalf & Koons 2007; Ruokolainen *et al.* 2009; Smallegange *et al.* 2014). Temporally autocorrelated vital-rate variation may be an adaptation to non-independent phases of the environment, for example, high recruitment cued to extreme weather or disturbances (Morris *et al.* 2006; Stige *et al.* 2007), favouring a selection of trait polymorphism (Orzack 1985; Uller 2008) and

therefore increasing viability under a more variable climate (Nadeau *et al.* 2016). In particular, positive autocorrelation, which increases the likelihood of an environment remaining in one particular state (e.g. drought), may benefit species tolerant of climatic extremes and has been shown to increase invasiveness of aliens (Fey & Wiczyński 2016). Therefore, trait selection and population dynamics in taxa sensitive to temporal autocorrelation may directly influence their potential to respond or adapt to environmental change (Morris *et al.* 2006; Koons *et al.* 2009; Engen *et al.* 2013; Nadeau *et al.* 2016).

Population-level responses to temporal autocorrelation are mediated by vital rates with the relatively strongest effect on the population growth rate (Franco & Silvertown 1996, 2004; Tuljapurkar & Haridas 2006). The relative effects of survival, growth and reproduction on population dynamics meanwhile determine differences in life-history traits such as generation time and strategies, that is, the combination of traits (Stearns 1992; Gaillard *et al.* 2016; Salguero-Gómez *et al.* 2016b). These interactions between vital rates, life histories and environmental variation have been studied extensively in stochastic population analyses assuming *i.i.d.* Various studies have shown that long-lived species with *slow* life histories are generally buffered from increased environmental variation (Morris *et al.* 2008, 2011; Sæther *et al.* 2013; but see Jongejans *et al.*

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2010; McDonald *et al.* 2017). Unlike long-lived species, short-lived species with *fast* life histories, where reproduction contributes greatly to population dynamics, are expected to show increasing fluctuations in population sizes with increasing environmental variation (Morris *et al.* 2008; McDonald *et al.* 2017).

Although changes in the patterning of environmental noise may have significant implications for population viability worldwide (Heino & Sabadell 2003; Ruokolainen *et al.* 2009; Fey & Wiczynski 2016), it remains largely unknown whether the proposed link between life-history strategies and resilience to environmental fluctuations (Morris *et al.* 2008) holds in autocorrelated environments. Theory predicts that populations that recover slowly from past perturbations should be more sensitive to temporal autocorrelation than those that are more resilient (Tuljapurkar & Haridas 2006). Low resilience has been shown for iteroparous, long-lived plant species in constant environments (Salguero-Gómez *et al.* 2016b). However, the limited empirical evidence shows weak support for increased sensitivities, or greater absolute changes, of population growth rates of long-lived species to changes in temporal autocorrelation. For example, studies on 11 terrestrial mammal species (Morris *et al.* 2011; Engen *et al.* 2013) did not detect significant patterns, while others have concluded that long-lived species in autocorrelated environments, both animals and plants, are buffered from environmental variation regardless of reproductive strategy (e.g. Metcalf & Koons 2007; Morris *et al.* 2008).

To determine which life-history strategies are most sensitive to temporal autocorrelation, we carried out stochastic simulations using matrix population models (MPMs hereafter) from 327 plant, 3 algae and 124 animal populations. We ask whether species with slow life histories show low sensitivities of stochastic population growth rates ($\log \lambda_s$) to temporal autocorrelation because of a low effect of environmental variation on these life histories (Morris *et al.* 2008). We also explore whether sensitivities differ between habitat types, as expected from the selection to distinct environmental noise patterns associated with them (e.g. Steele 1985; Vasseur & Yodzis 2004; Ruokolainen *et al.* 2009). We classify species according to main life-history strategies by performing a phylogenetically informed principal component analysis (PCA) on life-history traits derived from MPMs. We then simulate stochastic population dynamics of each species' population by perturbing vital rates that define the MPMs based on serial autocorrelation in presumed environmental states. Our approach allows us to assess whether the effects of temporal autocorrelation on a given species' demography can be predicted from its life-history strategy, habitat type or both.

MATERIALS AND METHODS

Matrix population models and vital rates

We used the COMPADRE Plant Matrix Database (Salguero-Gómez *et al.* 2015) and COMADRE Animal Matrix Database (Salguero-Gómez *et al.* 2016a) to obtain demographic data. These two databases provide high-quality MPMs representing a wide range of life histories, growth

forms and habitat types (Jones *et al.* 2014; Salguero-Gómez *et al.* 2016b). We limited our study to species that were studied in natural (i.e. unmanipulated) environments as we were interested in naturally occurring patterns of life histories. For studies with > 2 years of data and/or > 1 population, we obtained the arithmetic element-by-element mean of all MPM entries to represent the average MPM across all studied years (Tuljapurkar & Haridas 2006). If several studies described the demography of one species, we kept MPMs of both studies only if they described distinct population dynamics from different ecoregions, for example, species in the invaded vs. natural range. Otherwise, we chose the study with greater temporal or spatial resolution. Using these and other selection criteria (Appendix S1), we retained MPMs for 449 species and a total of 454 populations, including 330 plant/red algae and 124 animal populations (Appendix S1). All MPMs contained information on stage- or age-specific survival, transitions (progression/retrogression) and reproduction (Fig. 1).

Life-history patterns

To characterise variation in life-history strategies among the populations studied, that is, variation in the pace of life (fast vs. slow) and reproductive strategy (spread of reproduction across the lifespan), we derived five life-history traits from each MPM (Appendix S2). The chosen traits are commonly used in comparative demographic studies for plants and animals (e.g. Franco & Silvertown 1996; Tuljapurkar *et al.* 2009; Salguero-Gómez *et al.* 2016b): generation time (T), age at sexual maturity (L_x), annual sexual reproduction (ϕ), degree of iteroparity (S) and net reproductive rate (R_o). Details on the calculation of the traits can be found in Appendix S2 (see also Caswell 2001).

To define main life-history strategies and relate them to sensitivities of $\log \lambda_s$ to temporal autocorrelation in vital rates, we performed a varimax-rotated, phylogenetically informed PCA on the derived life-history traits (Revell 2012; Salguero-Gómez *et al.* 2016b). The traits were log-transformed and scaled to $\mu = 0$ and $SD = 1$ to agree with PCA assumptions. To correct for phylogenetic relatedness among species in the PCA, we constructed a species-level phylogenetic tree coercing different populations of the same species as dichotomous branches at the five species' tips. In the tree, branch length informed about phylogenetic relatedness (Appendix S2). The phylogenetically informed PCA then linked the phylogeny to life-history traits via a modified covariance matrix and estimated Pagel's λ , a scaling parameter for the phylogenetic correlation between species (Freckleton *et al.* 2002; for details see Appendix S2).

Defining environmental states

To incorporate temporal autocorrelation into stochastic simulations of population dynamics of the 454 populations, we defined a discrete Markov chain consisting of two states, (1) favourable or good environmental conditions and (2) unfavourable or bad conditions. The transitions between the two states were defined using the transition matrix:

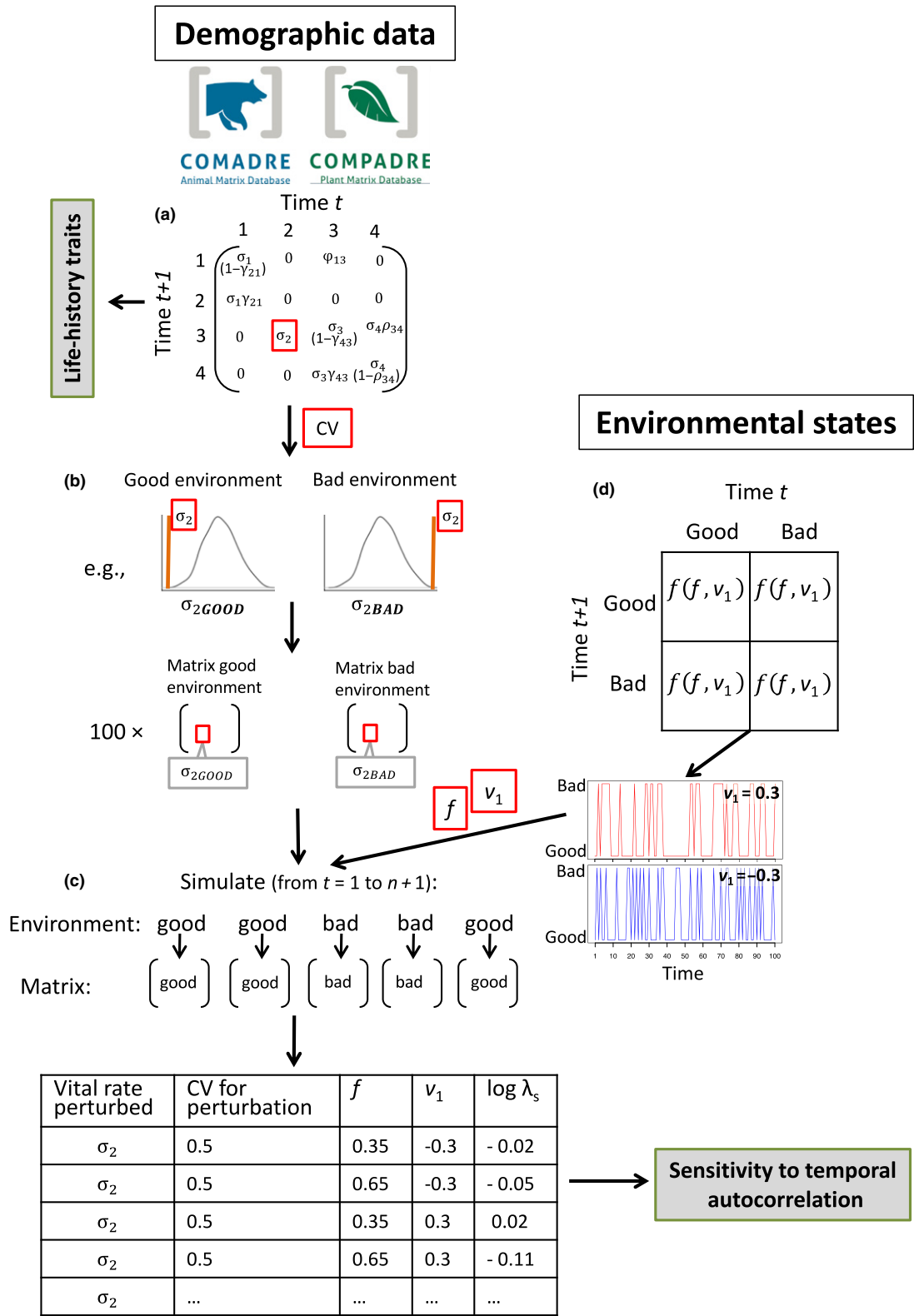


Figure 1 Simulations to test patterns of sensitivity of the stochastic growth rate, $\log \lambda_s$, to temporal autocorrelation across life histories. From each matrix population model (MPM), obtained from COMADRE/COMPADRE, we extracted stage/age-specific vital rates as shown in (a) for survival (σ_2) of the second stage of an example MPM. We then perturbed each vital rate as a function of different coefficients of variation (CV) to obtain distributions of a vital rate above and below its average value (orange) and assembled MPMs for good and bad environmental states (b). Lastly, we used a Markov chain defined by the frequency of the good state (f) and autocorrelation coefficient (v_1) to create environmental sequences (n time steps) as shown by two examples in (c). We then simulated stochastic population dynamics where at each iteration t the environment was associated with one representative MPM (d). Grey boxes show analysis outputs; $\gamma/\rho/\varphi$ – stage-specific progression/retrogression/reproduction respectively.

$$\begin{pmatrix} p_g & p_b \\ 1-p_g & 1-p_b \end{pmatrix}$$

Here, p_g and p_b are the probabilities of transitioning to the good environment at time $t + 1$ when the environment was good or bad at t respectively. These transitions can be derived from the long-term frequency of the good environment (f) and temporal autocorrelation, v_1 (Tuljapurkar & Haridas 2006), where $p_b = f(1 - v_1)$ and $p_g = (v_1 + p_b)$.

Linking vital rates to environmental states

For most populations (85%) used in this study, < 5 annual transitions were recorded (Appendix S1). This may limit the reliability of estimates of vital-rate variability and, in particular, autocorrelation – both requiring long-term population time series (Jongejans *et al.* 2010; Engen *et al.* 2013; Metcalf *et al.* 2015). Therefore, while using real life histories described by various combinations of mean vital rates, we perturbed vital rates away from their means for all 454 MPMs based on biologically relevant assumptions of environmental variation and patterning (but see Empirical analyses for additional simulations using observed vital-rate covariation). At the same time, our approach assumes that (a) transition probabilities between environmental states remain constant through time and (b) all vital rates potentially respond to changes in environmental conditions (i.e. incomplete buffering), although these responses are constrained as detailed below.

To link vital rates to the good and bad environment, we derived above- and below-average distributions of each vital rate across populations using three different coefficients of variation (CV; Fig. 1). We determined a new average vital-rate value for good (bad) environments by increasing (decreasing) a given vital-rate value from the mean MPM as a function of the CV (Fig. 1; Appendix S3; Koons *et al.* 2008). For vital rates describing reproduction, the three raw CV values were used in simulations (see Appendix S3 for additional simulation using higher CV). Vital rates describing survival or stage/age transitions, however, were assumed to have a binomial distribution and therefore a maximum bound on their variance and CV (CV_{\max} ; Morris & Doak 2004). We obtained new average values of these vital rates for good and bad states as a function of $CV \times CV_{\max}$. This constraint prevented vital rates with high average values (e.g. adult survival in long-lived species) to vary more extremely between good and bad environmental states than vital rates with relatively lower average values, making simulations more biologically realistic (Koons *et al.* 2008; see Fig. S3.1 in Appendix S3 for distribution of vital rates across environmental states). Such extreme variation would otherwise occur when the CV used in simulations approached the maximum variance of binomial vital rates.

Using the new average vital-rate values for good and bad environments, we simulated 100 vital rates from the beta (for survival/transitions) and gamma (for reproduction) distributions for each environmental state, keeping the non-perturbed vital rates at their mean and therefore assuming no vital-rate covariation. We then assembled 100 MPMs per species and vital rate for good and bad environments. Each of these

100 MPMs was picked at random during stochastic simulations of population dynamics (Fig. 1).

Stochastic simulations of population dynamics

We performed one simulation run for each vital rate per MPM using two long-term frequencies of the good environment ($f = 0.35$ and 0.65) \times 3 autocorrelations ($v_1 = -0.3, 0$ and 0.3) \times 3 degrees of vital-rate variation between good and bad environments (CV = 0.2, 0.5 and 0.8). For each parameter combination, we obtained the stochastic population growth rate, $\log \lambda_s$, by projecting population dynamics for 500 000 time steps after discarding the initial 20 000 iterations (see Tuljapurkar *et al.* 2003). For each species, we repeated 50 simulation runs to account for the stochastic process from which MPMs for good and bad environmental states were assembled (sampling from beta and gamma distributions; Appendix S3). From all simulations, we obtained the sensitivity of $\log \lambda_s$ to autocorrelation, $S^{v_1} = \frac{|\partial \log \lambda_s|}{|\partial v_1|}$ the absolute changes in the stochastic population growth rate as autocorrelation changed from 0 to positive (0.3) or negative (-0.3) (Fig. 1). We quantified changes in absolute terms to focus on the magnitude of S^{v_1} .

Sensitivities to temporal autocorrelation across life histories

We used generalised additive models (GAMs) to correlate S^{v_1} , averaged across the 50 simulation runs, to simulation parameters (f and CV) and life histories (PCA axes) while accounting for nonlinear trends. We assumed a log-normal distribution of S^{v_1} , which provided a better fit to the data than a normal distribution (Appendix S3), and chose the most parsimonious models based on AIC scores (Akaike 1971). First, we quantified differences in S^{v_1} across f and CV. For each combination of f and CV, we then modelled S^{v_1} across distinct vital rates perturbed in simulations as a function of life-history strategies, that is, scores along the main PCA axes. To increase statistical power and facilitate comparisons across species, we grouped perturbed vital rates prior to modelling by

- (1) assigning each stage/age class of an MPM to either the propagule, pre-reproductive, reproductive or non-reproductive class (Appendix S1) and
- (2) summing S^{v_1} for survival, stage/age transitions and reproduction within each class (Appendix S3).

When fitting GAMs, we excluded vital-rate classes (mostly non-reproductive class) that were present in $< 20\%$ of populations (Appendix S3).

Lastly, to explore global trends in S^{v_1} , we modelled S^{v_1} across different vital-rate classes as a function of major habitat type. To allow for statistical comparisons, we summed S^{v_1} for survival, transitions (progression/retrogression) and reproduction for all stages/ages. Information on major habitat types was obtained from COMADRE and COMPADRE. We collapsed habitat types from the original 39 types to five: temperate, tropical & subtropical, arid, alpine & arctic and aquatic (see Appendix S1). We accounted for the effect of MPM dimension and population on the variation in S^{v_1} by fitting the

latter two as random effects in the GAMs (see Appendix S3 for discussion on the relevance of MPM dimension).

Empirical analyses

To test the robustness of simulating different CV against natural vital-rate variation, we extracted vital rates for a subset of 109 populations with at least three annual MPMs and used the observed vital-rate correlation to estimate population dynamics in good and bad environmental states (Appendix S4). That is, we assembled 1000 MPMs for good and bad environments from a multivariate vital-rate distribution, using copulas to combine marginal gamma (for reproduction) and beta (for survival/transitions) distributions based on vital-rate correlations (Koons *et al.* 2008; Jongejans *et al.* 2010; Appendix S4). Simulation and statistical analyses of S^v_i then followed the steps outlined above, although we collapsed the habitats to two categories, temperate ($n = 63$) and other ($n = 46$) to allow for statistical comparisons.

We used two approaches to test the robustness of simulating temporal autocorrelation. First, we compared our S^v_i obtained from simulations of temporal autocorrelation (for the subset of 109 species modelling observed vital-rate CV) with empirical evidence for the contribution of temporal autocorrelation to vital-rate variation. To obtain the latter, we identified 13 species where MPMs were available for ≥ 10 consecutive annual transitions within a given site. We then regressed all vital rates at time t against vital-rate values at $t-1$, fitting first-order autoregressive models using a quasi-binomial (for survival/transitions) and gamma (for reproduction) error distributions. Lastly, we correlated average deviance explained by the models across vital rates, a measure of the importance of temporal autocorrelation in explaining vital-rate variation, to S^v_i . Second, we compared our simulated S^v_i using the full set of 454 populations with published information, noting whether autocorrelated environmental variation was explicitly examined in previous studies, and whether it significantly affected stochastic population dynamics (Appendix S4).

RESULTS

Life-history patterns

The life histories of the 449 studied species (454 populations) were adequately captured by two main PCA axes that together explained 67% of variation in key life-history traits (Fig. 2). Following the Kaiser criterion (Appendix S2), we kept these first two PCA axes for subsequent analyses, since their associated eigenvalues were > 1 . PCA axis 1 largely captured the position of the species' populations along the fast-slow continuum, with greatest positive loadings for age at sexual maturity (L_s) and generation time (T), and negative loadings of annual sexual reproduction (ϕ), depicting a trade-off between survival and reproduction (Gaillard *et al.* 2016; Salguero-Gómez *et al.* 2016b). PCA 2 largely captured variation in reproductive strategies, with largest positive loadings corresponding to highly iteroparous (S) species with a high net reproductive rate (R_o) and ϕ (Fig. 2). The positive loading of ϕ onto PCA 2 was explained by a strong association between annual reproduction and

iteroparity in plants (Salguero-Gómez *et al.* 2016b). When considering plants/algae and animals separately, ϕ loadings onto PCA 2 were negligible for the latter (Figure S2.1), which is consistent with previous studies on animals (reviewed in Gaillard *et al.* 2016). Similarly, R_o loaded onto PCA 1 because of the presence of long-lived plants with high R_o in the data (Fig. 2; Fig. S2.1). Therefore, the fast-slow and reproductive-strategies axes were not entirely represented by unique traits when considering both plants and animals. The phylogenetic relationships of the examined species moderately shaped their relative positioning along the life-history strategy space with Pagel's $\lambda = 0.61 (\pm 0.1 \text{ SD})$.

Sensitivity to temporal autocorrelation across life histories

The stochastic population growth rate, $\log \lambda_s$, did not change significantly in magnitude when positive (from 0 to 0.3) vs. negative (from 0 to -0.3) changes in temporal autocorrelation were simulated (Fig. S3.5). Thus, here we present results of S^v_i , the sensitivity of $\log \lambda_s$ to positive perturbations of autocorrelation. S^v_i increased > 10 -fold as the coefficients of variation (CV) used for perturbing vital rates increased from 0.2 to 0.8, but differed little among the two frequencies of the good environmental state, f (Table 1; but see Appendix S3 for the strong effect of f on $\log \lambda_s$).

For the examined CV and f , a significant interaction between the fast-slow (PCA 1) and reproductive-strategies (PCA 2) continua and vital-rate classes perturbed described S^v_i patterns (Table 1; Fig. 3a). For most vital-rate classes, faster life histories showed high S^v_i across a wide range of reproductive strategies. Highest S^v_i were found for pre-reproductive survival and transitions (Table 1; Fig. 3a). Populations with slower life histories, on the other hand, showed increased S^v_i only when they also had a high reproductive output (Fig. 3a). When including the effect of habitat, populations were most sensitive to temporal autocorrelation when the perturbations happened in survival (S) and transitions (T) rather than in reproduction (R) across all habitats considered (Fig. 3b). Perturbations of the latter vital-rate class had the highest effect in aquatic habitats. Removing outliers both in simulated S^v_i and PCA scores did not significantly affect GAM relationships (Appendix S3).

Empirical analyses

The additional empirical analyses validated our simulations. S^v_i based on naturally observed vital-rate covariation for 109 populations showed very similar patterns of variation along life-history strategies and habitat types as S^v_i based on simulated CV (Table 1; Fig. 4a; Fig. S4.1). In addition, S^v_i were significantly correlated (using a log scale; $R^2 = 0.63$; $P < 0.05$) with the contribution of autocorrelation to observed vital-rate variance from 13 populations (Fig. 4b). Lastly, we obtained significantly higher S^v_i for populations where previous studies included temporal autocorrelation in stochastic population models than for populations modelled assuming only i.i.d. environments ($t_{65} = 2.3$, $P = 0.01$; Fig. 4c; Appendix S4). However, the effects of temporal autocorrelation on population dynamics have been assessed for only 8% of the 454 populations simulated here and have been omitted for many

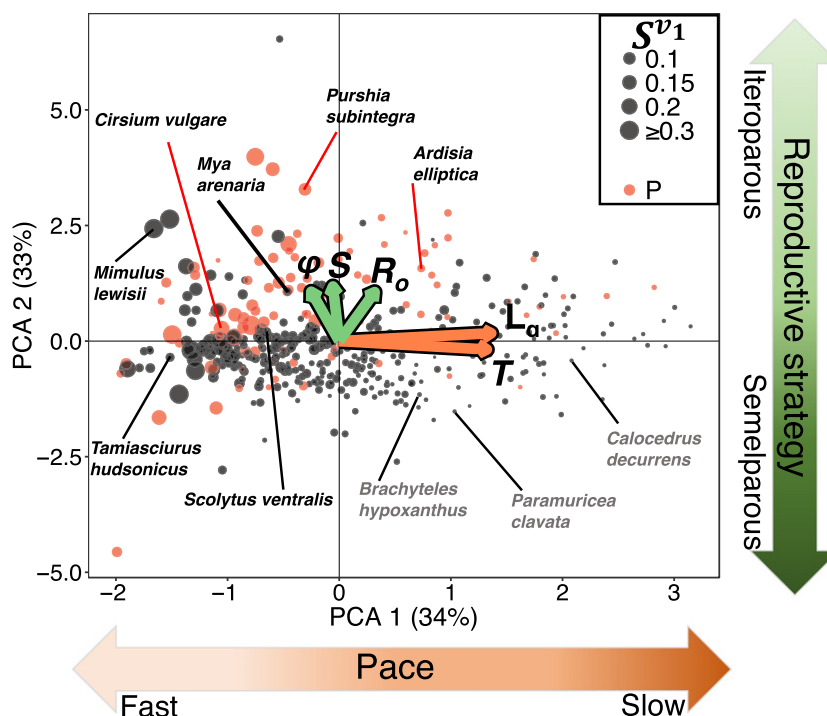


Figure 2 Life histories of study species' populations (points) are characterised by life-history traits representing the pace of life (fast–slow continuum) and lifetime reproduction (reproductive strategies). To characterise life histories, a phylogenetically informed PCA was performed on five traits: generation time (T), age at sexual maturity (L_a), sexual reproduction (ϕ), iteroparity (S) and net reproductive rate (R_o). Arrow lengths are proportional to the loadings of each trait onto the two axes. Red points show populations with propagule (P) stages. Point sizes are proportional to the sensitivity of the stochastic population growth rate to temporal autocorrelation, S^{v1} , summed across all vital rates (for simulations at $CV = 0.5$ and $f = 0.65$). Locations along the axes of some populations for which the effect of temporal autocorrelation on population dynamics has been assessed are shown; populations are coloured based on relatively low (< 25 th percentile; grey) and high (> 50 th percentile; black) S^{v1} .

species with high S^{v1} (Appendix S4). Our simulations indicate that *c.* 44% of populations affected by environmental variation may also be affected by changes in the patterning of environmental states (Fig. 4c; Appendix S4).

DISCUSSION

Identifying life-history strategies associated with high sensitivities of demographic processes to temporal autocorrelation may allow for a much needed inference on species affected by predicted changes in the patterning of environmental noise (Heino & Sabadell 2003; Laakso *et al.* 2003; Engen *et al.* 2013). Using data on 454 naturally occurring plant/algae and animal populations (449 species), we provide global empirical evidence that the fast–slow continuum of life-history variation, interacting with a species' reproductive strategy, help predict the demographic vulnerability to temporal autocorrelation (Table S2.2). In addition, our results highlight that life-history responses to temporal autocorrelation may be strongly mediated by vital rates and habitat types affected by changes in environmental patterning.

Although, generally, long-lived species have been predicted to be more sensitive to perturbations in population structure than short-lived species (Franco & Silvertown 2004; Salguero-Gómez *et al.* 2016b; but see Gamelon *et al.* 2014) and hence potentially to temporal autocorrelation (Tuljapurkar & Haridas 2006), our simulations indicate that most long-lived

organisms are buffered not only against increases in vital-rate variation (e.g. Morris *et al.* 2008; Tuljapurkar *et al.* 2009; McDonald *et al.* 2017) but also against changes in its temporal patterning. Our results agree with theory showing a negative correlation between generation time and magnitude of the effect of serial vital-rate correlations on fitness in age-structured populations (Tuljapurkar *et al.* 2009; Sæther *et al.* 2013). In empirical studies, low sensitivities to both interannual vital-rate variation and temporal autocorrelation have been found for several mammal species, for example, *Brachyteles hypoxanthus* and *Gorilla beringei* (Morris *et al.* 2011; Engen *et al.* 2013). Most of these species were located at the slow end of life-history strategies in our analyses and exhibited, along with most long-lived aquatic (e.g. *Paramuricea clavata*) and tree and herbaceous species (e.g. *Calocedrus decurrens*) low simulated sensitivities of their growth rates to temporal autocorrelation (Fig. 2).

The low sensitivities of population growth rates to temporal autocorrelation in long-lived species may be explained by vital-rate specific responses to changes in environmental patterning (Ezard & Coulson 2010). Species with slow life histories are characterised by high juvenile and adult survival (Franco & Silvertown 2004; Morris *et al.* 2008, 2011), which are typically less affected by environmental variation (Morris & Doak 2004; McDonald *et al.* 2017) or autocorrelation (Morris *et al.* 2008, 2011). Our simulations replicated such buffering by constraining the variation in binomial vital rates

Table 1 Across simulations of vital-rate variation (*i*), sensitivity of the stochastic growth rate, $\log \lambda_s$, to temporal autocorrelation (S^{vi}) is affected by simulation parameters (CV and *f*) and differs between vital rates, life histories (defined by PCA scores in Figure 2; see also Table S2.1) and major habitat types. S^{vi} also differ among habitat types when observed vital-rate covariation is modelled (simulating only *f*; *ii*). Parameter estimates from the most parsimonious GAM models are shown (see Appendix S3 for all models considered and details on parameters). Models fit at CV = 0.5 and *f* = 0.65 are indicated by arrows

	Response variable	Explanatory variables	Best model	% S^{vi} variation explained
(i) Simulated CV	$\log(S^{vi})$	<i>f</i> , CV	$-8.30_{(0.08)} + 1.77_{(0.05)}CV_{0.5} + 2.85_{(0.05)}CV_{0.8}$	41.8
	$\log(S^{vi})$ → CV _{0.5} ; <i>f</i> _{0.65}	vital rate, PCA 1, PCA 2	$-0.21_{(0.04)}f_{0.65} + s(\text{MatDim}_{df:6.3}) + s(\text{population}_{df:391.1})$ $-6.89_{(0.27)} + 1.15_{(0.28)}\text{Survival PR} + 0.5_{(0.29)}\text{Survival R} + 1.1_{(0.29)}\text{Transitions PR} + 0.11_{(0.30)}\text{Transitions R} - 2.17_{(0.77)}\text{Reproduction P} - 1.79_{(0.29)}\text{Reproduction PR} + te(\text{PCA 1, PCA 2, Survival P}_{df:3.0}) + te(\text{PCA 1, PCA 2, Survival R}_{df:3.0}) + te(\text{PCA 1, PCA 2, Transitions PR}_{df:6.0}) + te(\text{PCA 1, PCA 2, Transitions R}_{df:6.4}) + te(\text{PCA 1, PCA 2, Reproduction P}_{df:6.4}) + te(\text{PCA 1, PCA 2, Reproduction PR}_{df:6.1}) + s(\text{MatDim}_{df:6.3}) + s(\text{population}_{df:50.8})$	56.3
	$\log(S^{vi})$ → CV _{0.5} ; <i>f</i> _{0.65}	vital rate, habitat, PCA 1, PCA 2	$-4.55_{(0.24)} - 0.30_{(0.09)}\text{Transitions} - 3.62_{(0.09)}\text{Reproduction} - 0.19_{(0.23)}\text{Temperate} - 0.02_{(0.24)}\text{Tropical\&Subtropical} - 0.30_{(0.27)}\text{Arid} + 0.04_{(0.5)}\text{Aquatic} + te(\text{PCA 1, PCA 2, Survival}_{df:6.1}) + te(\text{PCA 1, PCA 2, Transitions}_{df:4.5}) + te(\text{PCA 1, PCA 2, Reproduction}_{df:13.2}) + te(\text{PCA 1, PCA 2, Alpine\&Arctic}_{df:2.6}) + te(\text{PCA 1, PCA 2, Temperate}_{df:12.1}) + te(\text{PCA 1, PCA 2, Tropical\&Subtropical}_{df:2.6}) + te(\text{PCA 1, PCA 2, Arid}_{df:5.5}) + te(\text{PCA 1, PCA 2, Aquatic}_{df:4.8}) + s(\text{MatDim}_{df:3.0})$	72.8
	$\log(S^{vi})$ → <i>f</i> _{0.65}	<i>f</i>	$s(\text{MatDim}_{df:0.3})$	39.4
(ii) Observed CV	$\log(S^{vi})$	habitat, PCA 1, PCA 2	$-6.44_{(0.19)} - 0.76_{(0.31)}\text{other} + te(\text{PCA 1, PCA 2, Temperate}_{df:1.9}) + te(\text{PCA 1, PCA 2, other}_{df:1.0})$	51.7
	$\log(S^{vi})$ → <i>f</i> _{0.65}			

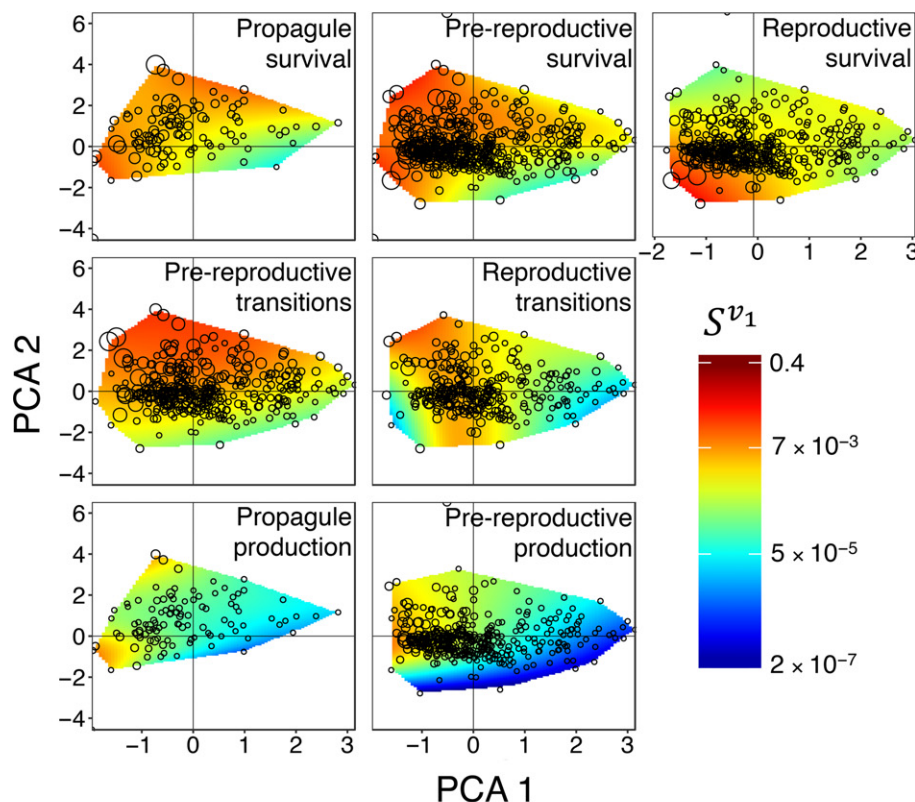
CV (categorical covariate with values 0.2, 0.5 or 0.8) – coefficient of variation used to perturb vital-rate classes (P)propagules, Pre-reproductive (PR), and (R)eproductive; *f* – frequency of the good environmental state; MatDim/populations – dimension of the matrix population model and populations used in the study respectively (fitted as nested random effects); Functions $te(x_{df})$ and $s(x_{df})$ are the tensor products and spline smoothing functions of *x*, respectively, with the given degrees of freedom *df*. Significant smoothing terms are in bold.

by the maximum coefficients of variation (CV_{\max} ; Morris & Doak 2004; Koons *et al.* 2008). Therefore, slow life histories vulnerable largely to changes in adult survival, which varied little if the average value was high (Appendix S3), responded overall weakly to changes in temporal autocorrelation. On the other hand, perturbing adult survival resulted in highest sensitivities for populations where this vital rate showed more variation but remained of relatively high importance for life-cycle dynamics, that is, for shorter lived semelparous species (Metcalf & Koons 2007). Similarly, some short-lived species, such as the invasive plant *Brassica napus*, where annual reproduction dominates life-cycle dynamics (Garnier & Lecomte 2006), may be little affected by environmental patterning in growth or survival (Fig. 4b). These patterns of vital-rate specific sensitivities remained when omitting a maximum bound on vital-rate variation to allow for extreme life histories (Fig. S3.4 in Appendix S3; Jongejans *et al.* 2010; McDonald *et al.* 2017) or when increasing the CV in reproduction perturbations (Fig. S3.5 in Appendix S3). This indicates that our results were not an artefact of the CV used. Most importantly, our simulations highlight that interpreting a population's

responses to temporal autocorrelation requires detailed knowledge of the variation in underlying vital rates (Morris & Doak 2004; Ezard & Coulson 2010).

In addition to its location along the fast–slow continuum, a species' reproductive strategy (Salguero-Gómez *et al.* 2016b; Salguero-Gómez 2017) was also critical to capture its sensitivity to temporal autocorrelation. In fact, some long-lived species showed relatively high sensitivities to temporal autocorrelation (Fig. 2). Most of these species were plants, for example, the invasive big-sage (*Lantana camara*) or the tropical, seed-bank-producing tree, *Ardisia elliptica*, and were characterised by relatively large net reproductive rates (R_0) and degree of iteroparity. These species explain the positive loading of R_0 onto the first life-history strategy axis, otherwise dominated by life span and total annual reproduction. They show that slow life histories may be highly susceptible to environmental patterning if they also spread their reproduction over many years (McDonald *et al.* 2017). Examples may include species that “track” changes in the environment by opportunistically recruiting in gap openings in forests or coral reefs. For such species, predictable, high-recruitment events,

(a) Vital rates



(b) Habitat types

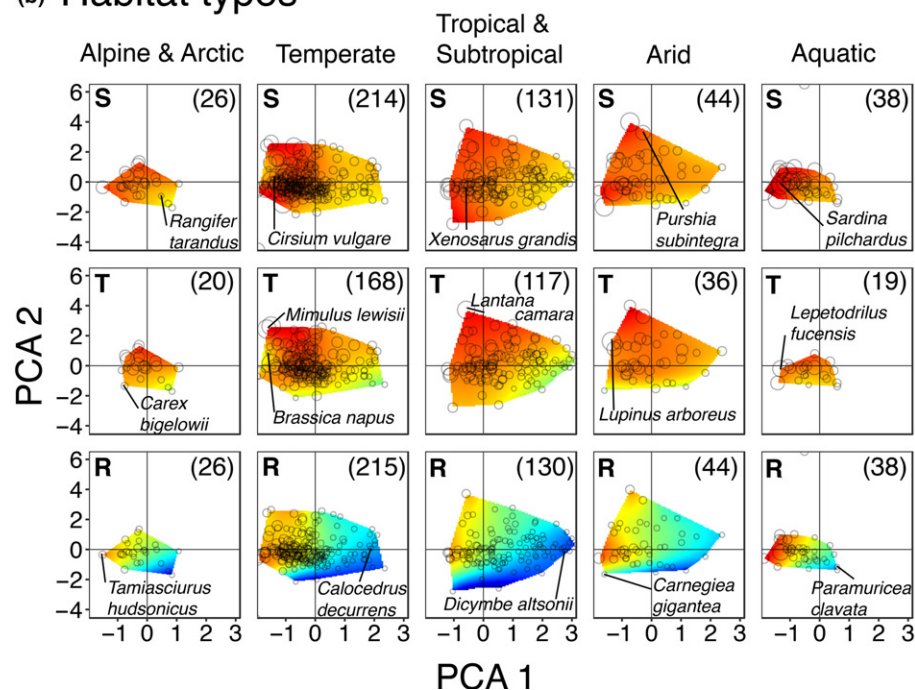


Figure 3 Fast life histories (along PCA 1) across reproductive strategies (along PCA 2) generally show highest sensitivities of the stochastic growth rate, $\log \lambda_{ss}$, to temporal autocorrelation (S^{v1}). Raster plots show predictions of S^{v1} across the two PCA axes after perturbing various classes of vital rates across all habitats (a) and for five different habitat types (b). In (b), vital-rate classes include survival (S), stage/age transitions (T) and reproduction (R). Predictions were limited to the range of observed PCA scores. Number of vital-rate samples in each habitat are shown in parentheses. Points are proportional to raw S^{v1} obtained from simulations. The results of vital-rate perturbations at $CV = 0.5$ and $f = 0.65$ are shown here (see Appendix S1 for all results and Table 1 for significance of predictions). Locations along the axes of example populations discussed in the main text are shown.

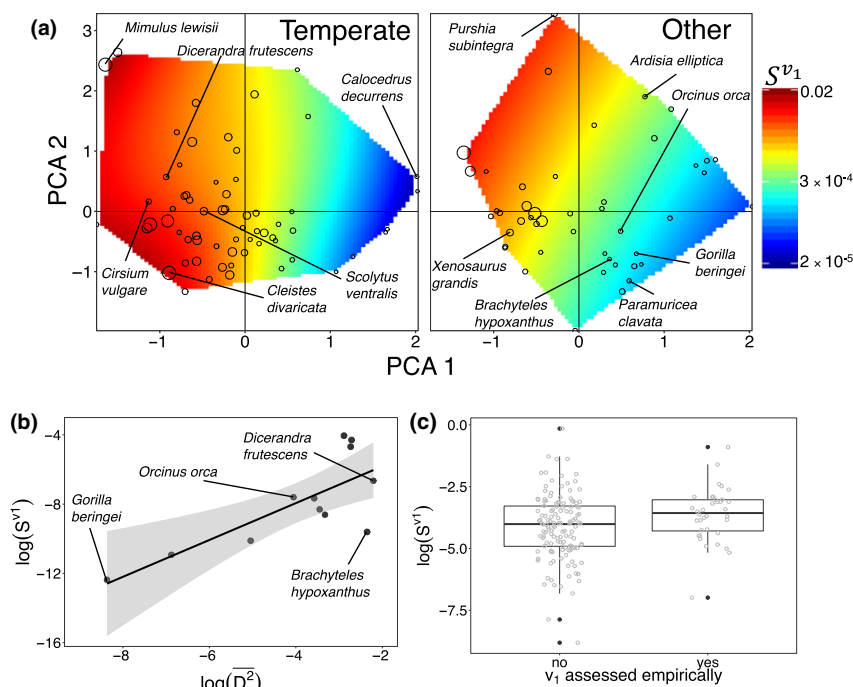


Figure 4 Empirical patterns of the sensitivities of the stochastic growth rate, $\log \lambda_s$, to temporal autocorrelation (S^v_1) validate simulations. (a) Among 109 population for which observed vital-rate covariance was used to define good and bad environmental states, fast life histories (along PCA 1) across reproductive strategies (along PCA 2) show highest S^v_1 in temperate and other habitat types. (b) S^v_1 obtained from simulations (a) for 13 species with the longest time-series data (≥ 10 annual matrix population models) are significantly correlated, on a log scale, ($R^2 = 0.63$; $P < 0.05$) with relative deviance explained by first-order autoregressive vital-rate models for the same populations (deviances, \bar{D}^2 , shown are averaged over vital-rate models). (c) For the 454 populations used to simulate S^v_1 $\log(S^v_1$; grey points) differ among published studies that determined v_1 to be an important component of population dynamics (yes) and ones that did not (no).

accompanied by relatively low survival of offspring, define population dynamics (Connell 1978; Metcalf *et al.* 2009). On the other hand, slow life histories with semelparous reproductive strategies have been shown to be buffered from changes in the patterning of environmental states (Metcalf & Koons 2007) and also exhibited lowest sensitivities in our simulations (Fig. 3).

Apart from some slow life histories with a high reproductive mode, a large number of taxa with a short life span and high annual reproduction exhibited the highest simulated sensitivities to temporal autocorrelation. This was true across semelparous and iteroparous species. Short-lived aquatic, for example, the clam *Mya arenaria*, and terrestrial animals, for example, the rodent *Tamiasciurus hudsonicus*, many insect species, for example, *Scolytus ventralis*, and numerous short-lived plants, particularly invasive ones, for example, *Cirsium vulgare*, are highly sensitive to changes in vital rates other than adult survival and also exhibit high variability in these vital rates depending on the state of the environment (Meijden *et al.* 1992; Morris *et al.* 2008; Koons *et al.* 2009). For such taxa, assessing the patterning of environmental variation would likely greatly improve knowledge on population dynamics (Jongejans *et al.* 2010). However, such assessments are lacking, as quantified in a thorough literature review here (Appendix S4), particularly for aquatic species and insects.

A common life-cycle adaptation to track autocorrelated environmental variation, for example, to recruit only under

favourable conditions, is dormancy (Cáceres 1997; Morris *et al.* 2006). Plant species with dormant propagule stages (seed banks), adapted to disturbance-prone habitats (Doak *et al.* 2002), exhibited the highest sensitivities of their growth rates to temporal autocorrelation (Fig. 2a). For species like *Mimulus lewisii* (Fig. 2b), whose populations are regularly exposed to flooding, changes in disturbance regimes and patterns of post-disturbance habitat succession may be detrimental to viability (Angert 2006; Turner 2010). In addition, propagation of many invasive plant species is aided by seed banks (Gioria *et al.* 2012). Such species, for example, *Ardisia elliptica* or *Lantana camara*, exhibited high sensitivities to temporal autocorrelation, despite being relatively long-lived (Fig. 2). Although environmental patterning may be critical for plant invasion (Fey & Wiczyński 2016), its importance was only considered in previous studies for 1 out of 10 invasive populations included in our analysis (Appendix S1). For animals, MPMs do not typically include dormant propagule stages (Salguero-Gómez *et al.* 2016a,b), such as cysts or larvae in diapause (Cáceres 1997; Schiesari & O'Connor 2013). Such species may be particularly vulnerable to predicted changes in environmental condition or may constitute emerging pests (Cáceres 1997), and must therefore be the focus of more research.

In addition to species with dormant life cycles, habitat-specific population dynamics deserve a particular focus when assessing potential ecological impacts of global changes in

environmental patterning. Our simulations show that sensitivities of stochastic growth rates to temporal autocorrelation differ between major habitat types, particularly when perturbing reproduction. For the latter, sensitivities for fast life histories were highest in aquatic habitats, although populations in aquatic habitats represent a narrow sample of the total life-history PCA space (Fig. 3). Differences in sensitivities to temporal autocorrelation between terrestrial and aquatic systems may be explained by the fact that the latter are more strongly (positively) autocorrelated and therefore predictable (Steele 1985; Cáceres 1997; Vasseur & Yodzis 2004). Therefore, species, such as the fish *Sardina pilchardus*, may exhibit strong responses when environmental patterns change, particularly because of anthropogenic influences (Ruokolainen *et al.* 2009). However, MPMs on aquatic species are still relatively scarce (Fig. 3b), and the effects of temporal autocorrelation on population dynamics in aquatic systems are rarely considered (Appendix S4). The same is true for arctic and alpine habitats, where climate change is predicted to have major effects on population dynamics (Post *et al.* 2009). At the same time, the relationship between the two major life-history axes and the sensitivity to temporal autocorrelation held in all habitats studied, indicating that future changes in the patterning of environmental states may significantly affect species globally (Vasseur & Yodzis 2004; Turner 2010).

Our results are robust to the choice of CV when perturbing vital rates and are in agreement with previous empirical studies regarding the importance of temporal autocorrelation in population dynamics (Fig. 4; Appendix S4). However, five aspects need further attention. First, the MPMs that were available to us represent a biased subset of life histories, given the underrepresentation of certain habitat types and life cycle (Salguero-Gómez *et al.* 2015, 2016a). Our life-history framework must therefore be validated with future additions of little-studied taxa. Second, density dependence was not considered in our study but is known to interact with temporal autocorrelation to regulate population responses (Levine & Rees 2004; Greenman & Benton 2005; Engen *et al.* 2013). A recent study by Koons *et al.* (2016) incorporated population structure and density dependence to assess differences in vital-rate variation among life-history strategies; future studies would benefit from applying this approach to serial vital-rate correlations. Third, our simulations of changes in environmental states may not have reflected complex processes such as habitat succession (e.g. Tuljapurkar & Haridas 2006). Fourth, we simulated autocorrelated changes in environmental states on an annual basis, potentially producing stronger effects for short-lived species simply because of the temporal scale used (Vasseur & Yodzis 2004; Stige *et al.* 2007). Lastly, our simulations of vital-rate variation omitted complex demographic processes such as strong buffering (no variation in vital rates among years; e.g. Morris *et al.* 2011) and nuances in covariation beyond the ones considered for the 109 species (Jongejans *et al.* 2010). Therefore, while we provide a robust general assessment of the role of autocorrelation across life histories and habitats, we acknowledge that a more detailed assessment must rely stronger on empirical data.

CONCLUSIONS

Temporal autocorrelation in demographic processes, historically rarely considered in demographic studies, may have strong effects, across habitats, on fast and slow life histories alike, depending on the reproductive strategy. Importantly, we found that taxa exhibiting highest sensitivities to temporal autocorrelation are also the ones least studied. With predicted global changes in environmental patterning, we argue that future demographic studies will only accurately predict important population processes such as viability or invasiveness if researchers explicitly consider the effects of these changes on key underlying demographic rates like survival, stage/age-specific transitions and reproduction.

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AUTHORSHIP

RSG and MP performed phylogenetic analyses, MP performed modelling work and main analyses, with modelling framework on life-history traits developed by RSG and assistance in modelling provided by AO. MP wrote the first draft of the manuscript, and all authors contributed substantially to revisions.

DATA ACCESSIBILITY STATEMENT

All authors confirm that the data supporting the results are archived on Dryad and the data DOI is included. Matrix population models are available at www.compadre-db.org. The data supporting the results of the study can be found on Dryad <https://doi.org/10.5061/dryad.d851q>.

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