1	Life in fluctuating environments
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#### 34 Abstract

35 Variability in the environment defines the structure and dynamics of all living systems.

36 Organisms have evolved traits and strategies that allow them to detect, exploit and predict the 37 changing environment. Organisms maintain steady internal conditions required for physiological 38 functioning through feedback mechanisms that allow internal conditions to remain at or near a 39 set point despite a fluctuating environment. In addition to feedback, many organisms use 40 feedforward processes, which allow them to adjust in anticipation of an expected future state of 41 the environment. Daily, seasonal and multi-year cycles provide cues that organisms may use to 42 anticipate regular changes in physiologically-relevant environmental conditions. Using 43 feedforward mechanisms, organisms can exploit correlations in environmental variables and act, 44 based on information acquired in the present, to prepare for anticipated future changes. All living 45 systems exploit the information in variable, auto-correlated, and cross-correlated environmental 46 conditions to reduce their uncertainty about the environment. Strategies to obtain, store and act 47 on information about the conditional nature of future events are advantageous and are evidenced 48 in widespread phenotypes such as circadian clocks, social behaviour, diapause, and migrations. 49 Here we provide a framework describing how feedback and feedforward mechanisms operate 50 across scales of organization, and how they allow living systems to persist in fluctuating 51 environments. Humans are altering the ways in which the environment fluctuates, causing 52 correlations between environmental variables to become decoupled, decreasing the reliability of 53 cues. Human-induced environmental change is also altering sensory environments and the ability 54 of organisms to detect cues. Recognizing that living systems combine feedback and feedforward 55 processes is essential to understanding their responses to current and future regimes of 56 environmental fluctuations.

#### 57 Introduction

58 Global change is characterized by trends, cycles and variability in the environment on land and 59 in the oceans. Rates of change in climate [1], habitat loss and fragmentation [2], contamination 60 [3,4], nutrient deposition, and biocide application are high, raising concern among scientists 61 about the capacity of living systems to adapt and persist in the face of these changes [5–7]. While 62 on average, conditions are changing, so too are the patterns of variability around the trends [8.9]. 63 Long-term changes in the variance and autocorrelation of environmental fluctuations can affect 64 biodiversity and ecosystem processes [10-14]. We address here the task of developing an 65 integrated understanding of how individuals, populations, and communities respond to 66 environmental fluctuations.

67

68 Perhaps the simplest way for variation in the environment to affect living systems is for 69 biological systems to track their environment as it varies (Figure 1A, B). Considering 70 temperature variation as an example, biological rates such as photosynthesis or reproduction may 71 increase as temperatures increase and decline as temperatures cool. Fluctuations in food or water 72 availability may directly affect demographic rates and therefore population dynamics. Many 73 examples of biological variation have been explained this way - from insect population cycles 74 responding with a time lag under varying weather conditions [15,16] to population cycles in lynx 75 and hares [17] to the abundance of commercially valuable fish [18,19]. Similar explanations 76 have been invoked for the persistence of diverse communities [20,21]. In this view, living 77 systems react to the fluctuating environment through a range of *feedback* mechanisms -78 responding to the environment when their internal conditions have changed enough to illicit a 79 response in physiological, demographic or other ecological rates (Figure 1C, Figure 2A).

However, relying only feedback mechanisms can be problematic. When populations or
organisms respond directly to their internal states, they are vulnerable to large deviations and
poor conditions that could cause death or extinction. Fluctuating environments can bring
regularly occurring stressful or otherwise poor conditions (Box 1), and organisms or populations
may perform better and be more likely to persist if they can minimize their exposure to these
conditions or be phenotypically prepared for expected changes in advance.

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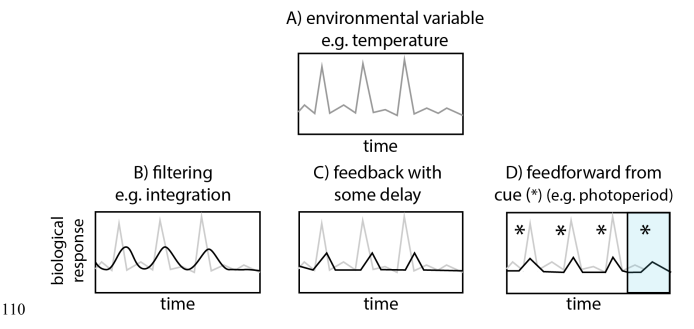
87 Ecological systems can minimize their experience of stressful conditions in varying 88 environments in a variety of ways. Many persist in fluctuating environments by anticipating 89 change through a variety of ecological and evolutionary cue and signal-based mechanisms 90 (Figure 1D, Figure 2B). These mechanisms convey information about correlations between the 91 state of the environment now and its likely state in the near future. Organisms use this 92 information in many ways including circadian rhythms, phenology, behavioral plasticity [22] and 93 transgenerational parental effects. These *feedforward* mechanisms, in contrast to *feedback* 94 mechanisms, allow organisms to prepare (or prime) themselves and/or their offspring for 95 environmental change so that they can adaptively buffer or exploit expected environmental 96 change (Box 2). Ecological and evolutionary models point to the long-term fitness benefits of 97 these processes [23]. Here we review feedforward processes and convey their essential role in 98 the adaptive responses of ecological systems in which many organisms are responding to 99 variable and uncertain environments.

100

101 The challenge for global change ecology is to understand how living systems - organisms,

102 populations, communities - respond and adapt to shifts in the structure of environmental

- 103 fluctuations. We combine knowledge of how the biophysical environment is changing and how
- 104 organisms, populations and communities can respond and adapt to change at different temporal
- scales. We consider feedback and feedforward mechanisms at the level of individuals [24,25],
- 106 populations [10,26], and communities [20]. We argue that a framework combining feedback and
- 107 feedforward mechanisms is required to achieve a robust understanding of how ecological
- 108 systems persist and maintain functioning in fluctuating environments.
- 109



111 Figure 1. Biological systems filter, integrate, respond to and anticipate environmental variation. 112 A) Environments are characterized by regular fluctuations in environmental variables (e.g. 113 temperature, light, precipitation, oxygen). B) Biological systems (individuals, populations, 114 communities) filter or integrate environmental fluctuations (grey line), thereby smoothing 115 environmental time series (black line). As a result, time series of biological or ecological 116 processes that respond to environmental variation tend to have less high-frequency noise 117 compared to the environmental variable itself (i.e. they become 'redder' [see Box 1]) as they are 118 translated through biological systems. C) Feedback mechanisms (i.e. those that respond to their

119 own internal state) allow organisms to respond to environmental fluctuations, but only after the 120 fluctuation has occurred. Therefore, there is an inevitable time lag in the response. D) 121 Feedforward mechanisms are signal or cue-based and use the state of the environment to 122 anticipate environmental change. The advantage of this approach is that organisms are prepared 123 for the expected change in environmental state, which reduces the lag that is inherent in B) and 124 C). The disadvantage with feedforward mechanisms is that if the cue (\*) becomes uncorrelated 125 with the future environmental state (i.e. the cue becomes an inaccurate indicator of the future 126 state) then organisms may initiate an anticipatory behaviour that is no longer beneficial in the 127 later selective environment (blue shaded area in D).

128

129 Characterizing correlations in fluctuating environments at different biological scales 130 Fluctuations in physical, resource, and biological conditions are a core feature of most ecological 131 environments. Here we address two features of this variability: 1) the correlation in time within 132 single variables (i.e., autocorrelation) and 2) the correlations that exist among multiple variables 133 (i.e., cross correlations). Both auto- and cross-correlation patterns occur at the full range of scales 134 and resolutions of space and time, and ecological systems reflect these features of temporal 135 structure at more than one - but not all - scales. In Box 1, we summarize methods we can use to 136 quantify relevant scales of correlation and we address types of correlations that allow organisms 137 to time life events and behaviours that have consequences for fitness.

138

139 Autocorrelation and predictability

140 Periodic, or repeating, temporal fluctuations occur at multiple scales and include diurnal and

141 seasonal cycles of light and temperature, quasi-periodic variation in climates at multiannual (e.g.

142 North Atlantic Oscillation, El Nino Southern Oscillation), decadal and millennial time scales 143 (e.g., Milankovich cycles). Aperiodic fluctuations also characterize variation in biotic conditions 144 that link to niche relations, such as resource availability or predation pressure (Figure 1A). 145 Characteristic features of fluctuations (i.e. predictability of environmental changes and 146 periodicity of cycles) hold information that may be used by organisms to time important life 147 history activities to align with conditions best for survival, reproduction and growth. Box 1 148 Section 1 describes how we can characterize *predictability* of temporal dynamics in a single 149 environmental variable, such as temperature. Predictability emerges when environmental 150 variables are temporally or spatially autocorrelated, reflecting the increased likelihood that 151 current conditions predict near-future conditions, such as long runs of above- or below-average 152 conditions. Environments that are dominated by variation at low frequencies (i.e. cycles with 153 long periods, high temporal autocorrelation; Box Figure 1) are more predictable to organisms 154 living in them because current conditions are likely to be accurate predictors of near-term future 155 conditions.

156

## 157 Correlations among different environmental variables

Changes in environmental variables such as light intensity, photoperiod or rainfall that are correlated with some later selective environment can be used as '**cues**' (Table 1). Organisms use the information represented by cues in fitness-defining ways (i.e. timing of growth and reproduction). For example, a cue early in the season can allow organisms to anticipate future favorable conditions for reproduction, migration or development, and initiate the biological processes that will allow these life history events to occur at the time of favorable conditions. In this way, organisms can match their phenotype to expected environmental conditions, increasing their fitness [23]. The value of a cue is related to the correlation, or mutual information [27],

166 between a cue and a later environmental state. In other words, the benefit of the cue to an

167 organism increases as the cue reduces uncertainty about the future environmental state.

168

169 Organisms experience the same environment differently

170 Species interact with the environment over certain ranges of variation and not others, and this 171 influences how they respond to and exploit temporal variation. Species may only detect and 172 respond to fluctuations and cues at a given scale (Box 1, Figure 1C). Species with life spans on 173 the order of a few years have typically evolved to coordinate key life events such as reproduction 174 or hibernation with seasonal shifts in food, mate, or predator availability. More generally, if 175 organisms' generation times and lifespans are longer relative to the period of fluctuations, and 176 individuals experience predictable environmental conditions, then feedforward mechanisms are 177 likely to evolve [28,29]. In contrast, organisms occurring in environments that exhibit little 178 variation within their lifetime, or highly unpredictable variation, are unlikely to rely on cues and 179 anticipatory mechanisms (e.g. [30]).

180

An environmental event or change in state that is used as a cue for one species may be noise for another species. For example, a bacterium moving through surface waters in the ocean may experience strong covariation in temperature, light intensity and oxygen availability. In this environment, an increase in temperature might be correlated with a decrease in oxygen minutes later. If an organism is capable of detecting that correlation, it may adjust its metabolism to prepare for the subsequent low oxygen state [31–33] (Box 1 Figure 2). The same change in temperature and oxygen that was vital to the bacterium may be considered 'noise' to a fish

swimming by. In this way, organisms respond to and integrate fluctuations in their environmentsaccording to their life histories (Box 1 Figure 1).

190

205

# 191 Integrating concepts from systems biology to classify strategies for dealing with fluctuating 192 environments

193 Living systems are characterised by their capacity for homeostatic control, which is their ability 194 to maintain a viable state despite variability in their environment. A homeostat is any set of 195 processes or mechanisms that results in a system property or process being maintained at a 196 (quasi) constant level, within a fluctuating environment. Variables held under homeostatic 197 control remain within a narrower range of values than if they were not regulated, and the 198 regulated value is maintained within a range that is consistent with the viability of the organism. 199 Here we describe how homeostasis is achieved via feedback and feedforward control 200 mechanisms (Box 2), and their combination in the form of general adaptive systems. This 201 classification expands on an earlier framework proposed by Rosen [34,35]. These three classes -202 feedback, feedforward, and general adaptive systems - group different strategies for maintaining 203 viability and fitness in variable environments (Figure 1). Strategies within each class share 204

204 common attributes for how biological systems persist in fluctuating conditions, this may mean

206 the fitness of the organism or the population. Some of these classes, such as feedback and

minimizing the effects of detrimental fluctuations, while exploiting conditions that may increase

207 feedforward (including model-predictive control), have been the focus of a great deal of research

in complex system science, engineering and theoretical biology (e.g., [36,37]). Like any

209 classification it is just one way of understanding how eco-evolutionary systems respond to

210 fluctuating environments and it is meant as a framework to locate the focus of analysis, to guide

211 inquiry about change in ecological systems and to facilitate comparisons among systems.

212

### 213 Class 1: Feedback homeostats

Homeostasis by negative feedback is the most familiar form of adaptation in physiology [38].

215 The mechanisms, such as regulatory pathways, leading to homeostasis in body temperature,

216 water content, energy levels, nutrients and essential cofactors (e.g. iron) are well studied in a host

of model and non-model organisms [39]. For example, in one third of the oceans the

218 bioavailability of iron limits primary production, and phytoplankton have evolved strategies to

219 acquire and recycle iron even when it is extremely limiting. For example, the picoalga

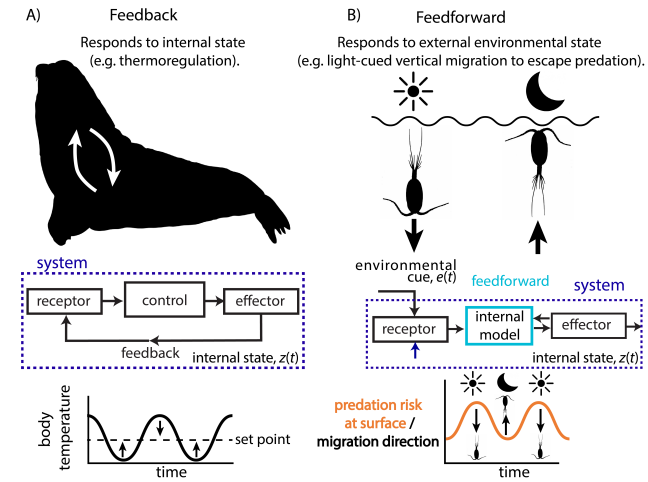
220 Ostreococcus uses the protein ferritin to regulate iron uptake and recycling, and this iron

homeostasis is essential for cell survival under iron limitation [40].

222

223 In abstract terms, any homeostatic system can usually be decomposed into a controlled system or 224 process (some aspect of an organism's physiological system) and a controller (e.g. a regulatory 225 pathway) (Figure 2A). In reality, there may be no simple dichotomy into these subsystems, but in 226 many cases one can identify processes that fall into each. The key property of feedback 227 homeostatic systems is that the controller only measures the internal state of the controlled 228 system, z(t), and not the environment (Figure 2A). Deviations of the state of the controlled 229 system away from the homeostatic state results in a response modifying the dynamics of the 230 controlled system so as to diminish the deviation from the set-point (negative feedback). In the 231 simplest case, homeostats have no memory of past states. An example of this is the thermostat 232 controlling the temperature of your room, which functions by controlling the actions of a heating

system based on deviations in temperature from the given set-point. In endotherms,
thermoregulation occurs when the cooling of the blood stimulates centers in the brain which
'turn on' heat producing mechanisms of the body and the body temperature is adjusted back to
the set point so that temperature is maintained at a constant level [41] (Figure 2A).



238

Figure 2. A) Feedback processes, such as thermoregulation in endotherms, occur as organisms respond to changes in their internal state, z(t), such as deviations from some internal set point (e.g. internal body temperature becoming too warm or cold). B) Feedforward processes, such as diel vertical migrations, occur as organisms respond to some external environmental cue, e(t)

(e.g. light intensity). An internal model allows organisms to 'pull the future into the present' [42]by acting, in the present, on some cue that is correlated to a future environmental state.

245

246 Organisms use feedback mechanisms to adapt to fluctuations in their environment, reducing the 247 variation in their internal physiological state. In addition to feedback mechanisms that operate by 248 adjusting physiological conditions internally, organisms may also use behaviours that allow them 249 to avoid high-frequency and potentially damaging environmental states [43,44]. For example, 250 intertidal organisms exposed to high temperatures and desiccation stress at low tide can buffer 251 their exposure to thermal fluctuations by becoming inactive during exposure extremes (many 252 animals cycle between activity and inactivity on a daily basis). Organisms with a broader range 253 of thermal microenvironments have greater opportunities to thermoregulate, and access to these 254 microenvironments depends on motility, body size and features of the environment. The 255 combination of behavioral thermoregulation and controlling activity patterns allows organisms to 256 avoid variation in body temperature, especially at daily and annual frequencies [43]. Notably, 257 these filtering mechanisms do not require internal models that relate events separated temporally, 258 just the ability to sense internal state and respond as feedback homeostats.

259

Feedback homeostats function as a result of variation in their environment. Organisms are selected to maintain steady state conditions in a range of vital processes and the aggregate response of many individuals forming a population reveals variation among individuals in their capacity to maintain homeostasis in a dynamic biotic and abiotic environment. At the population level these feedbacks take the form of density-dependent variations around a carrying capacity. The performance of feedback mechanisms vary in their stability to deviations from steady state,

which is arguably why this topic has been the focus of so much theoretical research in ecologyand evolution.

268

The set-point or long-term steady state around which feedback regulation occurs is often variable and may be under selection in large populations. Species vary in their capacity to achieve homeostasis under limiting or stressful conditions, so competition among genotypes within and among species is key to understanding the diversity of homeostatic strategies, and the overall functioning of populations and communities under novel patterns of environmental change.

274

# 275 Class 2: Feedforward homeostats

276 Feedforward homeostats add the capacity of the controller to measure the state of the 277 environment. Here the controller can sense an environmental quantity whose present value e(t) is 278 correlated with a future value of the internal state  $(z_{t+\tau})$  of the controlled system (Figure 2B). The 279 correlation between e(t) and  $z_{t+\tau}$  is taken into account by the controller. In feedforward 280 homeostats the controller can modify the state of the controlled system in accordance with the 281 present value of e and z, so as to keep constant some required function of z. Feedforward 282 mechanisms differ fundamentally from feedback mechanisms because the system is using 283 information about the environment (e.g., cues) to predict and prepare for a later state. The 284 correlation between e(t) and  $z(t+\tau)$  represents a model of the external world that has evolved in a 285 system in which environments at one time and internal states at another have been correlated. For 286 this reason, they are sometimes classified as **anticipatory systems** [35] (Table 1). The internal 287 model must encode the range of environmental conditions to which the controlled system has 288 historically (evolutionarily) been exposed and is expected to encounter. In cases where the

feedforward system's model does not accurately predict *z*, perhaps because the environment now includes new states, then it will no longer be adaptive under some range of conditions. If this situation becomes common and is chronic then we say it is maladaptive, and it will threaten the viability of the organism and the population if the maladaptive state occurs for several generations.

294

295 Feedforward mechanisms have some advantages over feedback mechanisms. The controller 296 response is no longer purely deviation or error-driven, meaning that the state need not degrade 297 before it responds. Any purely feedback homeostat has an intrinsic time delay (constant) so this 298 risks failure before a corrective response can be activated. In environments that fluctuate rapidly, 299 or in novel ways, a feedback control system will track the fluctuations rather than exhibit steady 300 state, or homeostasis. Feedforward control systems are founded on regularities in the 301 environment (the correlation between e(t) and  $z(t+\tau)$ ) rather than off the deviations around the 302 set-point, or steady-state of z, that the feedback mechanisms uses. By adjusting ahead of the 303 environmental change, feedforward mechanisms avoid the costs of constant error correction. 304

Examples of feedforward control are very common in biology (Table 2). Any behaviour or activity that uses a cue to prompt its timing is predictive and model-based. Many organisms, ranging from single-celled algae to mammals, use circadian clocks (a type of internal model) to anticipate regular environmental changes and coordinate internal biological processes [45]. For example, plants upregulate photosynthetic machinery before dawn, allowing an immediate response to light when the sun rises [46]. The importance of these anticipatory mechanisms is demonstrated by the fact that when circadian clocks are disrupted, fitness decreases [47,48].

312 Plants and animals prepare life histories in spring and winter on the basis of day length rather 313 than internal temperatures. For example, trees stop growing and shed their leaves in autumn 314 based on day length cues in anticipation of impending winter [49]. Anticipatory developmental 315 switches between alternative phenotypes (i.e. direct development and diapause), are often cued 316 by photoperiod and have evolved independently in a wide variety of taxa [28,50,51]. These 317 switches are often established and maintained if cues are reliable (i.e. they are accurately 318 correlated with later fitness-defining environmental conditions) and available to the organism at 319 the appropriate time to influence development.

320

321 Feedforward mechanisms can also operate across generations. Parents can modify the phenotype 322 of their offspring in response to changes in the environment that act to increase parental fitness 323 by also increasing offspring fitness [52], using a set of mechanisms called anticipatory parental 324 effects, which are a type of transgenerational **phenotypic plasticity**. Anticipatory parental 325 effects are expected to occur in situations where parents can detect and identify current 326 environments, parental environments accurately predict offspring environments (i.e. the cues are 327 reliable), and parents can accurately transmit information to offspring so that it can be integrated 328 into offspring phenotypes [53]. In contrast, populations that experience completely unpredictable 329 and variable environments are unlikely to evolve anticipatory parental effects [27,54]. If 330 environments are variable and unpredictable, then diversified bet hedging, in which parents 331 produce offspring with a variety of phenotypes, are a better strategy [26].

332

333 As stated above, the risk with feedforward mechanisms is that some properties of the

environmental fluctuations may change so that the system's internal model is no longer accurate

or predictive of the future internal state. In this case, in evolutionary terms, the model is
maladapted, and fitness of the population may decline. But the advantage is that, under known
conditions, organisms are prepared for their likely future. If the population is composed of many
individuals with maladapted feedforward control, then population fitness will decline and may
result in extirpation. Examples of this include trophic mismatches due to climate change driven
shifts in the correlation between day length and temperature [55].

341

#### 342 Class 3: General adaptive systems

343 General adaptive systems (GAS) are characterised by an ability to combine feedback and 344 feedforward mechanisms. GAS measure multiple environmental states and can develop multiple 345 models linking their internal state to different e(t). They can also modify their internal models 346 and features of their environments in order to achieve desired future states. GAS can acquire the 347 ability to measure and integrate different sensory modalities about the environment's states; these can include a mix of visual, audible and olfactory states of the environment. Over extended 348 349 periods of time these multi-modal models of the environment may improve an organism's 350 expectation of its fitness and therefore allow a more adaptive short and long-term response to 351 fluctuating conditions. This feature boils down to an individual being able to acquire new 352 sources of information from the environment to reduce uncertainty in the measurement of its 353 state (epistemic uncertainty), and so more reliably anticipate its performance under fluctuating 354 conditions.

355

The ultimate adaptive ability of a GAS is the development of a set of behaviours to modify and manipulate the state of the environment, *e*, using some sort of effector. Modification of the

environment is directed so that the environmental conditions permit the system's future state  $z(t+\tau)$  to more closely match its physiological requirements. This capacity to control the state of the environment falls into definitions of ecosystem engineering [56].

361

In the context of global environmental change where anomalous patterns of environmental variation are occurring with increasing frequency it is not clear whether adequate evolutionary potential exists in existing feedback and feedforward mechanisms to ensure long-term persistence. The relative benefits of each class will be related to the type of variability organisms experience, which we discuss below, and may be altered under global environmental change.

367

368 Ultimately, there are always limits to predictability in ecologically complex ecosystems. All 369 model-based control is limited by the quality of the model *and* the fundamental limits to 370 predictability (ontological uncertainty). The degree to which organisms use feedback and 371 feedforward mechanisms depends on the predictability of the environmental fluctuations and the 372 relative costs and benefits of anticipating vs. reacting to environmental changes. Maintaining an 373 internal model that is required to anticipate environmental changes can be costly, and the degree 374 to which organisms use feedforward mechanisms depends on the costs and benefits of 375 anticipatory behaviours [32,33]. For example, sensing mechanisms involved in chemotaxis have 376 a metabolic cost, and presumably the cost increases as the accuracy of sensing increases [57]. 377 Unavoidable delays between measurement and response involved in feedback strategies also induce a metabolic or fitness cost. The fitness benefits of feedforward mechanisms are related to 378 379 the degree to which the ability to detect and act on cues improves expected fitness of the 380 offspring [27,58]. Ultimately, there may be a fitness trade-off between responding late (i.e.

simply reacting and not anticipating) and the fitness cost of maintaining highly accurate sensingmechanisms.

383

Evidence for feedback and feedforward strategies in ecological systems at higher levels of
 organization

386

387 Populations and communities are composed of individuals that differ in their homeostatic or 388 preferred internal state, and also differ in the range of environmental cues to which they are 389 adapted to detect and exploit to maintain short and long-term fitness. This diversity results in a 390 range of synchronous and asynchronous species' responses to the same environmental variables 391 that mediate when and whether they interact with other individuals to determine population and 392 community dynamics [59]. The collective dynamics of populations and entire assemblages of 393 species can be analysed and understood from the point of view that all living systems are 394 exploiting the information in variable, auto-correlated, and cross-correlated environmental 395 conditions.

396

#### **397 Population level**

A major challenge is to understand how information used by individuals to adaptively adjust their behaviour, movement, aggregation and reproductive investment through feedback and feedforward mechanisms scales up to mediate population fitness and dynamics when environments vary in their quality and predictability over time [60]. Population models differ greatly in the way they formalise fine-scale variation about individuals (e.g. phenotypic traits, life histories and behaviours), and at what level conditional information about the state of the environment is used by individuals. Decisions about how to model these features can have strong 405 effects on resulting population dynamics and the predictions these models make about the effects406 of changes in environmental variation [60–63].

407

Autocorrelated environmental fluctuations are currently understood to have large qualitative effects on the mean and variance of population dynamics and on the probability of extinction and colonization. The evidence stems from a large body of theory for unstructured [64–68], and stage-structured population models [69,70]. This has been supported in laboratory experiments [10,71,72] and analyses of large databases of population time series [73]. Temporal autocorrelation in environmental conditions is expected to have interactive effects with population size when density-dependent processes are at play, such as resource-limited growth,

416 such that time-integration of the environment is not simply additive [12,67,74]. In the case of

417 multiple equilibria, autocorrelated variation mediates the timing of switches between population

418 equilibria [65,75], which defines their resilience. Thus, the population model is a linear or

419 nonlinear filter of the environment where the modeled feedbacks (e.g. density dependence, or

420 switches in equilibria) either dampen or amplify the stochastic environmental signal, with

421 impacts on modelled population extinctions risks [75–77]. Here, feedbacks arising from density

422 dependence (e.g. under or overcompensation) and how they are modelled has a strong effect on

423 the variance, spectral colour and extinction risk of the populations [75].

424

425 Simple population models often involve no time delay between the environment and the

426 population response. Relaxing the constraint that all events happen instantaneously can greatly

427 alter how density dependence is expressed in population dynamics [63]. An important class of

428	models exists that integrates time delays in the model to reflect how vital rates observed in the
429	population arise from previous historical environmental states (e.g., because of changing
430	seasons). These carry-over effects can create quite complex dynamics [63,78]. Among these are
431	feedforward response systems, such as when the environments experienced by parents can
432	mediate the phenotypes and fitness of their offspring [78-80]. These maternal effects evolve
433	when mothers can gain reliable information from cues about the future state of the environment
434	their offspring will face [81,82]. In general, encoding these intergenerational effects into
435	population models can produce quite distinct and complex dynamics. For example, maternal
436	effects generally increase population variability in these models [78].
437	
438	In general, when there is temporal autocorrelation, current conditions not only determine the
438 439	In general, when there is temporal autocorrelation, current conditions not only determine the consequences of current decisions individuals make but are also informative of future conditions
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439 440 441	consequences of current decisions individuals make but are also informative of future conditions [83]. Population theory suggests that accounting for anticipatory parental effects and phenotypic plasticity is important and is improving our understanding of population level outcomes of
<ul><li>439</li><li>440</li><li>441</li><li>442</li></ul>	consequences of current decisions individuals make but are also informative of future conditions [83]. Population theory suggests that accounting for anticipatory parental effects and phenotypic plasticity is important and is improving our understanding of population level outcomes of changing environmental conditions. Changes to environmental predictability of any form (see
<ul> <li>439</li> <li>440</li> <li>441</li> <li>442</li> <li>443</li> </ul>	consequences of current decisions individuals make but are also informative of future conditions [83]. Population theory suggests that accounting for anticipatory parental effects and phenotypic plasticity is important and is improving our understanding of population level outcomes of changing environmental conditions. Changes to environmental predictability of any form (see Box 1) in either the abiotic and biotic environment may lead to maladapted cues. The
<ul> <li>439</li> <li>440</li> <li>441</li> <li>442</li> <li>443</li> <li>444</li> </ul>	consequences of current decisions individuals make but are also informative of future conditions [83]. Population theory suggests that accounting for anticipatory parental effects and phenotypic plasticity is important and is improving our understanding of population level outcomes of changing environmental conditions. Changes to environmental predictability of any form (see Box 1) in either the abiotic and biotic environment may lead to maladapted cues. The demographic consequences of these fitness declines, and the extent to which evolutionary or

448 Feedforward mechanisms influence species interactions. When species strongly interact,

449 fluctuations in abundance of one species can cause the other to respond, hence a varying

450 environment may be both abiotic and biotic. The abundances of Canada lynx and snowshoe hare

451 fluctuate in iconic predator-prey population cycles, out of phase such that peak lynx abundance 452 is followed by very low hare population sizes. These cycles have persisted for centuries, well 453 documented by fur trapping records [17]. Initially, resource limitation was thought to be the 454 primary driver of hare population cycling, which then was assumed to cause declines in lynx 455 abundance, reflecting food limitation. However, resource limitation could never fully explain the 456 cycles. Now, the explanation includes processes based on feedforward mechanisms in the form 457 of maternal effects. The first of these is that hares experience physiological stress when lynx 458 abundances are high and predation rates are high. Stressed mother hares are less successful at 459 reproduction and pass on symptoms of stress to their offspring. It has been hypothesized that 460 maternal stress and subsequent risk-sensitive behaviour in young hares may be a form of 461 maternal adaptive programming [84]. Juvenile hares with higher stress hormone levels spend 462 more time under cover and are less active during field trials, highlighting a potential mechanistic 463 route to allow individuals to cope with a changing environmental risk of predation [85]. A 464 second possible feedforward process occurs in lynx. Research on lynx in Newfoundland, 465 Canada, suggests that when prey is scarce, daughter lynx remain in their mother's territory, 466 repressing their own reproduction during times of hare shortages. Repression of reproduction 467 would keep densities low and allow individuals to reproduce in a subsequent year when hares 468 may be more abundant [86]. This picture of the role of feedforward mechanisms at the individual 469 levels augments significantly our understanding of how density-dependence responses at the 470 population and community levels maintain diversity.

471

472 *Cue-based synchrony in reproduction and species persistence* 

473 In environments that do not experience large environmental fluctuations in light or temperature, 474 some species have evolved to use complex combinations of multiple cues to time life history 475 strategies and synchronize reproduction events. On coral reefs, the high biodiversity and low 476 frequency of many species presents challenges for reproduction and mate finding. Some species 477 have evolved to use a combination of light, temperature, lunar and diurnal cues to reproduce at 478 specific times of year - only once, and within the same hour. Conspecifics use the same lunar 479 cues, so that gametes from the same species encounter each other in the water column and 480 fertilization occurs [87]. These spawning events not only allow rare species to reproduce, but 481 they provide food for predators. The diverse, biological system associated with coral reefs in a 482 relatively stable abiotic environment has generated its own fluctuations in the environment that 483 have in turn become a selective environment for the timing of releasing gametes.

484

485 Co-occurring species perceive and respond to varying environments differently and these 486 differences underpin explanations for the maintenance of diversity. For example, primary 487 producers have evolved to use different cues in the same environments; some species begin 488 budburst and leaf-out earlier than others in the same locations [88]. These differences may reflect 489 evolved partitioning of the temporal niche by primary producers. Phenological tradeoffs between 490 timing and productivity are at the heart of plant coexistence mechanisms [89]. Temporal storage 491 effects are another important mechanism for coexistence of species in fluctuating environments 492 [90]. For example, in aquatic systems, resting stages can be stored in sediments to emerge later, 493 allowing species to 'recolonize' their environment [91]. Similarly, the long-term coexistence of 494 winter annual plants in the Sonoran desert is based on functional tradeoffs in growth [92]. The 495 separation of species along a tradeoff between growth capacity and low-resource tolerance

496 corresponds to differences in demographic responses to environmental variation across years, in
497 particular growing season precipitation. Trade-offs in how species grow in fluctuating
498 environments are increasingly understood to mediate community-level climate change responses
499 [88,93] and biodiversity changes via the establishment of non-native species [94].

500

501 Differences among species in their internal models of the environment can explain the 502 maintenance of diverse food webs. In aquatic systems, many plankton populations in temperate 503 systems shift from stationary overwintering growth phases to fast-growing phases when 504 photoperiod becomes suitable and temperatures warm [95,96]. The spring bloom is the most 505 intensely productive time of the year in many pelagic systems, in which much of the annual 506 carbon is fixed before resource limitation sets in. The timing and magnitude of the spring bloom 507 influences ecosystem structure and function for the following year. Shortly following the spring 508 phytoplankton bloom and sudden resource availability, zooplankton populations grow rapidly, 509 grazing down fast-growing phytoplankton populations. Young of year fish consume 510 zooplankton, allowing fish to grow and spawn. The timing and magnitude of the bloom, and its 511 importance, exist because of temporal (annual) fluctuations in light and temperature. The variety 512 of biological processes that respond to this regular environmental fluctuation including 513 phenological cues on daylength and temperature, physiological cues as temperature warms, 514 triggering the end of diapause for some zooplankton, and temporal storage of populations and 515 biomass [96,97], enable the maintenance of diversity in these communities.

516

517 Diversity and homeostasis at the community level

518 Homeostasis at the community level has been considered to underlie the finding that species 519 richness at the community level is relatively stable even while environmental conditions and the 520 composition of species can vary substantially over time [98–100]. Compensatory dynamics 521 describe the negative correlation among species' abundances within the community - suggesting 522 one compensates ecologically for the other, in a negative density-dependent manner [101]. When 523 one species increases in abundance, others decline such that total diversity or energy flux remain 524 consistent throughout the change. Hence, community functions may remain within certain 525 bounds characteristic of homeostasis. When coexisting, competing species exhibit negative 526 covariances in population dynamics, such that the total resource use at the community level 527 remains more stable than would be expected by chance or by independent population changes 528 not connected temporally through the interaction [99]. Compensatory dynamics might thus be an 529 example of feedback homeostasis at the community level in a varying environment. However, 530 we are not arguing for a fixed set-point value for species' diversity, rather that feedback and 531 feedforward processes arising from interspecific interactions for limiting resources tend to 532 balance extinction and colonization, keeping diversity within bounds. Over the very long-term 533 (i.e. paleoecological scales) variation in environmental constraints linked to climate and resource 534 availability will mediate non-stationary variation in biodiversity [102].

535

#### 536 Anthropogenic influences on environmental fluctuations

There is growing evidence that humans are changing the way the environment fluctuates
[103,104]. Several key statistics, such as the variance, autocorrelation and periodicity of
environmental fluctuations are predicted to change over the coming century [8,9,105]. Humans
are also altering the reliability of the correlations underlying many environmental cues as their

541 timing and phases shift over time, within and across years. There is also evidence that humans 542 are modifying the ability of organisms to detect cues [104,106,107]. Changes to the sensory 543 environment, such as changes in light and acoustic conditions, visual properties of water, or 544 additions of chemical compounds may distort the production, transmission and perception of 545 signals and cues. For example, metal and chemical pollutants influence the development and 546 production of signals by influencing endocrine function and other cellular processes involved in 547 signal production [108]. We now assess the evidence for human-induced changes in 1) cue 548 reliability and detectability and 2) the statistical structure of environmental variability.

549

550 Changes in cue detectability

551 Human impacts on ecosystems are distorting or altering auditory, visual and chemical cues and 552 hampering their ability to be detected by focal organisms [106,109,110]. Acoustic pollution from 553 human sources interferes with the detection and discrimination of acoustic signals. For example, 554 low frequency, human-generated, noises in aquatic ecosystems, such as noise from boat traffic, 555 often overlap in frequency with the hearing range of most animals, and the frequencies of the 556 calls of many species, including marine mammals [111]. By masking acoustic signals, humans 557 are effectively decreasing the distance from which an individual is able to detect a conspecific's 558 call and making auditory cues more difficult to detect. Human impacts are also altering the visual 559 environment. Eutrophication and run-off are altering the availability of light in aquatic 560 environments, and changes to the bandwidth of available light can have severe consequences for 561 the detectability of cues among aquatic species. Eutrophication in Lake Victoria has altered the 562 light environment such that two species of cichlid fish have hybridized because females are 563 unable to distinguish red males from blue males [112]. High turbidity levels reduce the distance

564 from which predators can see their prey, which reduces foraging efficiency and food intake in 565 brown trout [113] and Eurasian perch [114]. Artificial light sources associated with human 566 settlements and ships on the ocean are altering lightscapes. For example, when artificial lights 567 are brighter than the horizon over the ocean, sea turtle hatchlings move towards human 568 settlements instead of the ocean [115]. Together, human-induced changes in the sensory 569 environment influence organismal fitness by altering individuals' ability to find food, avoid 570 predation, acquire mates, provide parental care and interact with various aspects of the biotic and 571 abiotic environment.

572

573 *Changes in cue reliability* 

574 Many feedforward mechanisms rely on light as an information source, and artificial light 575 pollution can cause adaptive feedforward mechanisms that rely on light as cue to become 576 maladaptive. Many organisms use lightscapes as cues for directional movement [115], and changing lightscapes can result in disruptions to movement patterns. For example, nighttime 577 578 light can alter nocturnal downstream migrations in Atlantic salmon [116]. Artificial light 579 pollution influences the orientation of individuals that rely on visual cues for daily movement 580 [117] and may disrupt light-cued diel vertical migrations in zooplankton [118,119]. Artificial 581 light after dusk or before dawn can cause phase shifts in circadian rhythms, either by delaying or 582 advancing the cycle relative to natural diurnal day-night cycles and thus cause physiological 583 functions to become out of phase with relevant ecological conditions. Persistent levels of low 584 light or short pulses of bright light from ships or cars can be enough to entrain circadian rhythms 585 [120,121]. In addition, artificial light can lead to mistiming of events which require photoperiod 586 cues. For example, some species of deciduous trees maintain their leaves for longer in autumn in

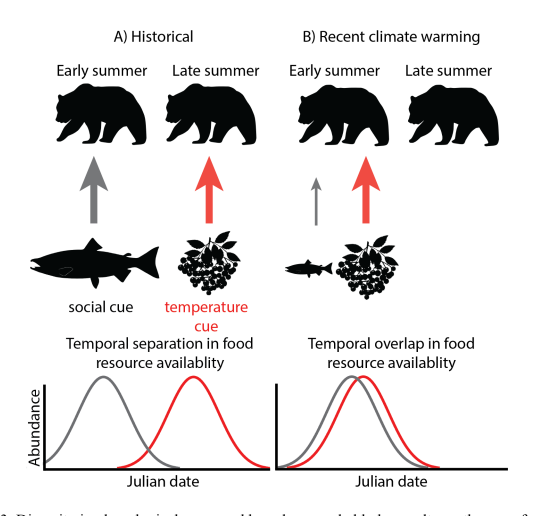
the vicinity of street lights [122,123], potentially leaving them exposed to higher rates of frostdamage.

589

590 Disruptions in relationships between historically related conditions (i.e. cross-correlations 591 between temperature and daylength) may alter the outcome of species interactions. If individuals 592 evolved to rely heavily on one correlated environmental cue, and that cue is no longer a good 593 indicator of some physiologically relevant condition at a later time, then this may result in the 594 mistiming of important life history events and lead to phenological shifts [55,124–126]. In a 595 community context, different organisms use different cues for their phenologies (i.e. 596 temperature, rainfall, photoperiod). Phenological mismatches may occur across trophic levels 597 when the cue used by one trophic level changes at a different rate than the cue used by a higher 598 trophic level [125,127–129] (Figure 3). Consumers generally have lower sensitivity to 599 environmental cues than their resources and, as a result, they generally have weaker responses to 600 changes in the cue than their resources, leading to potential mismatches in consumer-resource 601 interactions [130–132]. Even if both interacting species use same type of cue (e.g. temperature), 602 these cues may occur at different times of the year or have different dimensions (e.g. duration, 603 frequency, mean, extreme), and since temperatures at different times of the year have been 604 shifting at different rates, phenological mismatch may occur [130]. Similarly, even the same cue, 605 at the same time of year, can elicit different responses in co-occurring species [88]. 606 607 In a food web context, differences in the cues used by different species to time life history events 608 can cause shifts in major energy channels as the climate changes. In coastal Alaskan ecosystems,

brown bears feeding on salmon are a critical link between marine and terrestrial ecosystems.

610 Typically, brown bears feed on stream-spawning salmon early in the summer, and then switch to 611 feeding on elderberries later in the summer [133] (Figure 3). The sequential timing of the arrival 612 of stream-spawning salmon, followed by elderberries ripening creates a relatively long period of 613 foraging opportunities for bears during the short Alaska growing season. As spring temperatures 614 have warmed, elderberries have shifted to ripening earlier in the summer, overlapping more with 615 the stream-spawning salmon. One potential explanation for the difference in relative phenology 616 shifts is that salmon and elderberries rely on different environmental cues. While elderberry 617 phenology is likely cued by temperature [133,134], the cues salmon use to time their migrations 618 are likely a combination of temperature, stream flow and social information [135–138]. When 619 both resources are available at the same time, bears prefer elderberries, and abandon the salmon. 620 This climate-induced diet-switching by the bears due to synchronized resource availability may 621 fundamentally alter energy flows in stream food webs. Bears feeding on stream-spawning 622 salmon play a large role in modulating energy pathways in the food web and are capable of 623 transferring large amounts of marine-derived nutrients into terrestrial ecosystems and food webs. 624 As a result, changes in the relative phenology, caused by changing cues of prey species, can 625 drive diet switches of generalist consumers and potentially alter major energy pathways in 626 ecosystems.



628 Figure 3. Diversity in phenological cues used by salmon and elderberry alter pathways of energy 629 flow in food webs as the climate warms. A) Historically, brown bears fed on stream-spawning 630 salmon and then switched to feeding on elderberries once they were ripe, later in the summer. 631 This temporal separation in resource availability allowed bears to feed through an extended 632 period of the growing season. B) In recent years, red elderberries have begun ripening earlier in 633 the summer while the salmon have continued spawning at the same time. This means that red 634 elderberries are available to bears at the same time as the stream-spawning salmon. That the 635 elderberries have altered their phenology more than higher trophic levels, including salmon and 636 bears may be common across ecosystems, since primary producers tend to be more sensitive to 637 abiotic environmental cues [55]. The newly established synchrony in resource availability for

bears may fundamentally alter energy pathways in this coastal ecosystem. Based on data from[133,135].

640

#### 641 Changes in temporal variance and autocorrelation

642 Changing variance at different frequencies will alter the environment differently for organisms 643 with different life histories. Environmental variables have become more temporally 644 autocorrelated over the last fifty years, and these trends are expected to continue [105]. While 645 increased temporal autocorrelation may increase environmental predictability, and therefore 646 performance for some organisms, it may also reduce population persistence, because as the 647 duration of poor conditions increases, refugia and rescue effects are diminished and extinction 648 risk increases [139–141]. The effects of increased temporal variance in environmental variables 649 will depend on the frequency at which variance increases and the life history of the organisms 650 affected. For example, if variance increases at annual time scales, organisms with short 651 generation times that are active only during the summer months may experience large changes to 652 growth rates due to multiple successive generations experiencing high summer temperatures. For 653 longer lived organisms whose reproductive cycle encompasses the whole year, if variance 654 increases at the annual time scale, then the increase in warm temperatures may be balanced (or 655 not) by colder winter temperatures [105]. Alternatively, reduced variance at annual time scales, 656 such as reduced differences between summer and winter temperatures in the form of milder 657 winters, can substantially alter ecosystem structure and function. Changing community and 658 ecosystem responses to milder winters are enhancing productivity and expanding growing 659 seasons as climate changes in temperate and polar regions. This outcome is reducing the effects

of extreme seasonal conditions and the life history traits that allow organisms to reduce activityin winter.

662

#### 663 Looking forward

664 Organisms in naturally variable environments exploit fluctuations and correlations among 665 environmental variables to survive and persist. The ways in which they sense, anticipate and 666 respond to environmental fluctuations determines patterns of biodiversity. Humans are changing 667 patterns of auto- and cross-correlations upon which cues are based. The extent to which these 668 anthropogenic influences will alter the structure and function of ecosystems will depend on the 669 mechanisms by which individuals respond to and anticipate fluctuations and adapt to changing 670 fluctuation regimes. Here we have provided a framework that includes feedback and feedforward 671 as different modalities of response and described how these mechanisms operate at multiple 672 scales of biological organization. Recognizing that organisms employ a range of feedback and 673 feedforward systems to mediate fitness suggests we must study the internal models they use to 674 predict future ecological outcomes and how those models result in effects at population and 675 community levels. An understanding of community responses to environmental change will 676 require the study of the diversity of cues and internal models used by community members. 677

There is a high cost to ignoring the manner by which organisms and systems have adapted to fluctuating environments when considering global change. Although it is a tall order to empirically measure yet another aspect of biotic responses to environmental change, we suspect that similarities and generalities in response types will be revealed, allowing them to be predicted. If a feedforward mechanism exists, then an experiment that lacks appropriate cues

683 may grossly mis-estimate the effects of environmental change (e.g. the loss of  $CO_2$  responses in 684 fish when parental effects were allowed [107]). One way to probe the internal model of an 685 organism would be to expose it to different types of cues in a controlled way, so as to identify the 686 relevant cue. Manipulating the correlations between different environmental variables (e.g. 687 temperature and oxygen, or light wavelength and depth) would reveal which signal and cues are 688 important, and to what extent organisms can update their internal models when cues are no 689 longer reliable. Manipulating the colour of environmental noise by adding variation (power) at 690 different frequencies and studying responses at the individual, population and community levels 691 would allow us to understand how changes in fluctuations are amplified or absorbed across 692 trophic levels and how organisms with different life histories are influenced by fluctuations at 693 different frequencies.

694

#### 695 Conclusion

696 An outstanding challenge is to understand the degree to which feedback and feedforward 697 mechanisms generate the dynamics of living systems. Explicitly considering the processes by 698 which organisms respond to uncertainty about the future state of the environment may 699 dramatically change our predictions of how living systems will respond to global environmental 700 change. The task for ecologists is to discover the internal models which organisms use to 701 anticipate environmental fluctuations, and how diversity in these models among individuals and 702 species governs responses to environmental change in the context of populations and 703 communities.

704

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710	Boxes
/10	Boxes

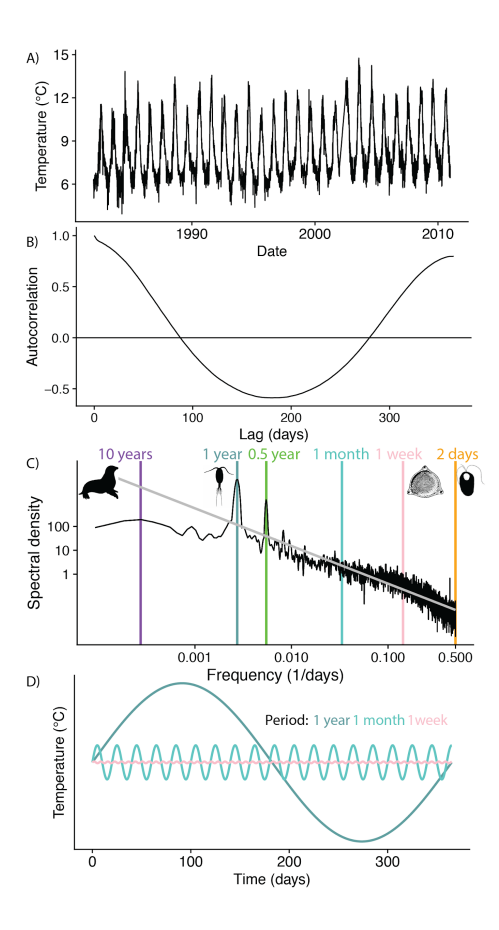
711	
712	Box 1. Quantifying the predictability of environmental fluctuations from an organism's eye
713	view
714	Here we consider the predictability of an environment from the perspective of organisms living
715	in fluctuating environments. We consider two types of predictability: 1) predictability that
716	emerges from temporal autocorrelation in a single environmental variable (e.g. how similar
717	today's temperature is to tomorrow's temperature); 2) predictability that emerges from
718	correlations between two distinct environmental variables (e.g. temperature and oxygen, or
719	photoperiod and temperature).
720	
721	1) Temporal autocorrelation increases predictability
722	
723	Regular variation in a time series lends itself to prediction (Box Figure 1A). The most
724	straightforward case is temporal autocorrelation without a time lag, in which the conditions at
725	any time point are very similar to the conditions in the previous time point. From the perspective
726	of an organism, the greater the temporal autocorrelation, the greater predictability of the
727	environment, because there is an increased probability of having long runs above or below
728	average conditions. Autocorrelation can be visualized using a correlogram, which quantifies the
729	dependence of values in a time series on values preceding them (at a distance of $k$ lags) (Box
730	Figure 1B).
731	

Time-series can present predictable variation through periodic variation, where conditions at a given time are most similar to conditions at some time in the past - perhaps in the previous year.
Environmental variation can incorporate multiple periods of variation (Box Figure 1C), and different biological processes or different organisms may cue on or focus on one or a few aspects of a complex temporal structure. Temporal autocorrelation increases as the dominance of variation at low frequencies increases.

738

739 Observing temporal variation and distinguishing patterns that might lend themselves to 740 prediction by biological systems can be challenging and requires appropriate statistical analyses. 741 Spectral analysis is a method to decompose variation in time series into component frequencies, 742 allowing one to determine how much of the variance in the time series is associated with 743 different frequencies (Box 1 Figure 1B). The Fourier transform [142] can be used to shift 744 between the time domain (i.e. time on the x axis) and the frequency domain (i.e. frequency on the x axis) (Box 1 Figure 1A to C). In this way, any time series can be rewritten as a sum of sine 745 746 waves, each with its own amplitude and phase. The spectrum, a plot of variance vs frequency, 747 provides a standardized map of the relative contributions of the underlying components of a time 748 series (e.g. yearly vs. daily cycles, Box Figure 1C). When there are smaller amplitudes and less 749 variance at high frequencies (short periods) compared to low frequencies (long periods), the 750 environment can be considered as being more predictable based on the current state, because 751 there is an increased probability of having long sequences of above or below the average 752 conditions. In this way, the predictability of the environment can be understood as the slope of 753 the relationship between variance and frequency. Specifically, if variance scales with frequency 754 (f) according to an inverse power law,  $1/f^{\beta}$ , then the predictability of the time series can be

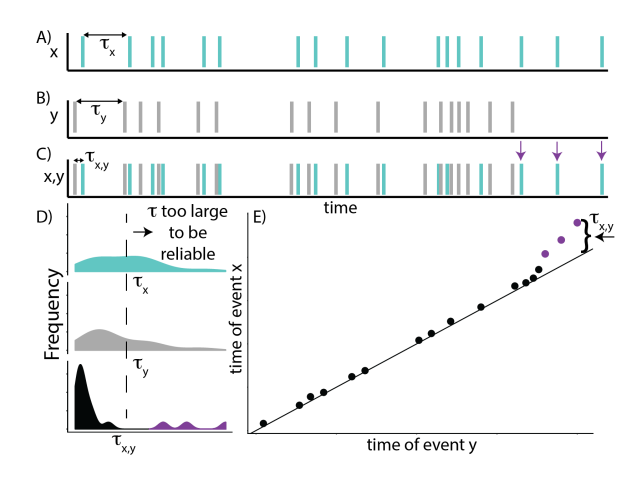
frequency (also called <b>white noise</b> ), and random through time. As the value of $\beta$ increases reveals autocorrelation at longer time scales, which means greater predictability, because t time series is dominated by variation at lower frequencies. By analogy with light we say th	755	quantified by the value of $\beta$ . Where $\beta = 0$ , this indicates that the time series is composed of an
reveals autocorrelation at longer time scales, which means greater predictability, because t time series is dominated by variation at lower frequencies. By analogy with light we say th temporal variation is reddened when it is dominated by low frequency (long period) cycles	756	equal mix of cyclic components at all frequencies, and the variation is random with respect to
time series is dominated by variation at lower frequencies. By analogy with light we say th temporal variation is reddened when it is dominated by low frequency (long period) cycles	757	frequency (also called <b>white noise</b> ), and random through time. As the value of $\beta$ increases it
760 temporal variation is reddened when it is dominated by low frequency (long period) cycles	758	reveals autocorrelation at longer time scales, which means greater predictability, because the
	759	time series is dominated by variation at lower frequencies. By analogy with light we say that
761 $0.5 < \beta < 1.5.$	760	temporal variation is reddened when it is dominated by low frequency (long period) cycles, and
	761	$0.5 < \beta < 1.5.$



764	Box 1 Figure 1. A) Variation in daily sea surface temperatures at a site off the coast of Norway
765	over the time period from 1981 - 2011. B) A correlogram of the time series in panel A, showing
766	the autocorrelation between time points as a function of time lag, $k$ (days). C) A Fourier
767	transform can be used to transform the time series in panel A to a frequency spectrum, which
768	illustrates how the variance (power) is spread across a range of frequencies. The negative slope
769	of this frequency spectrum, $\beta$ , is -1.58 (95% CI -1.60 , -1.56), characteristic of 'reddened' time
770	series in marine environments. Coexisting organisms, including a harbour seal, a copepod, a
771	diatom and a green alga, with different lifespans experience different components of the
772	frequency spectrum. D) The Fourier transform decomposes the time series into a set of sine
773	waves, each with a characteristic frequency and amplitude. Three of these frequencies (1 year
774	(dark green), 1 month (turquoise) and 1 week (pink)) from panel C are illustrated here.
775	
776	Wavelet analysis is an extension of spectral analysis, and is localized [143,144], in the sense that
	Wavelet analysis is an extension of spectral analysis, and is localized [143,144], in the sense that instead of estimating the variance spectrum of the entire time series, it estimates the frequency at
776	
776 777	instead of estimating the variance spectrum of the entire time series, it estimates the frequency at
776 777 778	instead of estimating the variance spectrum of the entire time series, it estimates the frequency at each point in the time series. It reveals changes in the variance spectrum through time and so is
776 777 778 779	instead of estimating the variance spectrum of the entire time series, it estimates the frequency at each point in the time series. It reveals changes in the variance spectrum through time and so is
776 777 778 779 780	instead of estimating the variance spectrum of the entire time series, it estimates the frequency at each point in the time series. It reveals changes in the variance spectrum through time and so is
776 777 778 779 780 781	instead of estimating the variance spectrum of the entire time series, it estimates the frequency at each point in the time series. It reveals changes in the variance spectrum through time and so is particularly useful for examining non-stationary time series in the context of climate change.
776 777 778 779 780 781 782	instead of estimating the variance spectrum of the entire time series, it estimates the frequency at each point in the time series. It reveals changes in the variance spectrum through time and so is particularly useful for examining non-stationary time series in the context of climate change.
776 777 778 779 780 781 782 783	<ul> <li>instead of estimating the variance spectrum of the entire time series, it estimates the frequency at each point in the time series. It reveals changes in the variance spectrum through time and so is particularly useful for examining non-stationary time series in the context of climate change.</li> <li>2. Predictability emerges from temporal context of correlated events.</li> </ul>

787 correlated (Box 1 Figure 2). As illustrated in Box 1 Figure 2, if x is a change in oxygen, and y is 788 a change in temperature, and if organisms are capable of internalizing the correlation between 789 these two variables (i.e. employ an internal model), they can exploit the correlation to anticipate 790 a vital change in the environment. For example, they can use an increase in temperature as a **cue** 791 that is associated with an impending drop in oxygen and adjust their metabolism (i.e. switch 792 from aerobic to anaerobic metabolic pathways) accordingly. In this way, even if a change in 793 oxygen *per se* is relatively unpredictable, as long as organisms can detect a change in 794 temperature, they can initiate a metabolic response in advance of the change in oxygen, thereby 795 increasing their performance relative to individuals who wait to sense and respond to the change 796 in oxygen.

797





800 Box 1 Figure 2. Organisms can exploit repeated associations between correlated environmental 801 variables with a time lag to anticipate change. In this example, x is an event (i.e. a decrease in 802 oxygen) that occurs in some random temporal sequence (A), as evidenced by the large variation 803 in the time lags between successive x events,  $\tau_x$  (D, top panel). Similarly, y is another event, (i.e. 804 an increase in temperature) which also occurs with a wide distribution of time lags,  $\tau_v$  (B, D, 805 middle panel). In spite of the unpredictability of x and y, x is highly predictable within the 806 temporal context of y, such that the delay between y and x is relatively constrained, as seen in the 807 distribution of time lags between event y and x,  $\tau_{x,y}$  (C, D, bottom panel). If event x no longer 808 occurs shortly after event y, and the two event types become decoupled in time, as illustrated in 809 the purple arrows in panel C, purple dots in panel E and purple peaks in the bottom panel in D,

810	then the predictability of the environment decreases, because the value of $y$ as a cue for $x$
811	decreases. Adapted from [31].
812	
813	
814	Box 2.
815	
816	Box 2. What is feedback vs feedforward, reactive vs. proactive?
817	Whether a process or event studied in ecology is reactive to the system's present state or
818	proactive to an expected state is open to debate, but a clean and operational distinction can be
819	made about what the organism (or any homeostatic system such as a cell or organ) senses and
820	what information it uses to adjust its behaviour, physiology etc. to the present, and likely future,
821	environment.
822	In feedback control systems, the organism responds to a sensed or measured deviation in its <i>own</i>
823	state, $z(t)$ , or performance relative to a desired, or reference, state (Figure 2A). The organism
824	senses a deviation and its distance from the desired state, regardless of what fluctuation causes
825	this difference.
826	In feedforward control, the change/disturbance(s) in the environment, $e(t)$ , are measured (e.g
827	cues, signals) and the organism's response is based on an internal model (Figure 2B). In a strictly
828	feedforward response there is no feedback with self to assess a deviation from the desired state.
829	It is the measured change in the environment, $e(t)$ , that causes the organism's behaviour or
830	physiology to change. It is adaptive if the cue permits a response that maintains positive fitness
831	under expected environmental change.

832 Indeed, in a feedforward system, the organism may simply respond to an external event, and treat 833 that event as a 'cue' that is temporally correlated with other environmental conditions such that 834 there is an order to them; one event can serve as a **cue** for a likely future event. If that future 835 event also presents a selective environment, then organisms that act on the cue to begin an 836 activity such as development or migration may have a fitness advantage over others that do not. 837 Certainly, more complex cognitive behaviours are also examples of feedforward systems, but 838 cognition is not necessary, and there are many examples in which selection acts on responses to 839 proximate cues that are correlated in time to future selective environments.

Feedback control is *reactive* since it reacts to changes in its own state, while feedforward is *proactive* since it acts ahead of the organism's expected change based on the environment's measured state. Feedforward systems also react to deviations, but they are in the measured state of the environment. In feedforward control, the system's output can change without any observable deviation from the desired state.

845

## 847 Tables

848	Table 1.	Definitions	of key terms.

Term	Definition	Examples
Cue	Environmental variable (either abiotic or biotic) that triggers an event or process and is predictive of a future environmental condition [145].	Variable features of the environment such as photoperiod, temperature, rainfall. For example, temperature is an environmental cue for sexual reproduction in many algal species, dispersal in fish, or diapause in invertebrates.
		By sensing cues early in the season, organisms can anticipate the best time to initiate seasonal reproduction, migration, dormancy, etc., or to produce a particular seasonal morph, thereby matching their phenotypes to the expected conditions [23].
Signal	Signals have four components [146]: (1) acts or structures produced by signalers, which (2) evolved for the purpose of conveying information to recipients, such that (3) the information elicits a response in recipients, and (4) the response results in fitness consequences that, on average, are positive for both the signaler and the recipient.	Pheromone trails laid by ants, peacocks' ornamented tail, electric pulses used to electric fish to communicate in water, bird songs.
	In contrast to cues, which may contain information as a by-product of organisms' behaviour, signals have evolved for the specific purpose of conveying information, and influencing others' behaviour.	
Prediction	A probabilistic conditional expectation about the future, informed by past and present events and an internal model. Allows organisms to prepare for impending changes in the environment	Cells can internalize correlations between multiple environmental variables (e.g. temperature and oxygen), which allows them to express an appropriate energy-

	<ul><li>[31].</li><li>"Prediction is not prescience but simply 'output from an anticipatory model""</li><li>[42]</li></ul>	extracting metabolic pathway at the right time. Predictive behaviour is in contrast to stochastic switching, or diversified bet hedging, which allows for diverse phenotypes but doesn't require prediction of any particular future environmental state.
Internal model	A simplified description of a system [42]. In organisms, this may be the physical instantiation of a probabilistic model [31]. We learn something new about a system by studying its internal model.	A model can be encoded in the pathways of a gene or metabolic regulatory network.
Feedback homeostatic control	A process or mechanism whereby a system quantity can be returned to at a constant level (the set point), within a fluctuating environment. A deviation from the controlled set-point is countered by a controller that modifies the dynamics of the controlled system so as to diminish the error [41]. Homeostasis typically involves a negative feedback loop that counteracts the error. This type of control only responds to the state of the controlled system rather than that of the environment.	Thermoregulation in endotherms, food switching to achieve stoichiometric homeostasis (i.e. regulate elemental composition) [147,148].
Feedforward homeostatic control	In a feedforward system, the control variable adjustment is not based on the self-state. Rather, the controller senses an environmental quantity, $e(t)$ , whose value is correlated to a likely future value of the state of the controlled system, $z(t+\tau)$ . This introduces the role of prediction. The controller can modify the dynamics of $z(t)$ according to the present value of $e(t)$ and the state of $z(t)$ , so as to maintain constant the state of z(t). In feedforward control, disturbances are detected and accounted for before they have time to affect the system.	Negative phototropism, autumnal plant cessation of growth, immune priming, heat hardening etc.

Anticipatory system	To anticipate means to expect or predict. Rosen (1985) [35] defined an anticipatory system as a natural system that contains an internal predictive model of itself and of its environment, which allows it to change state in accord with the model's predictions pertaining to a later instant. In contrast to a reactive system, which can only react in the present to changes that have already occurred in the causal chain, an anticipatory system's present behaviour involves aspects of past, present and future.	An individual organism (an <i>E. coli</i> cell, a tree, a copepod), any natural system that contains an internal model. See Table 2.
Phenotypic plasticity	Phenotypic plasticity refers to the ability of a single genotype to produce different phenotypes under different environmental conditions [23]. To do so organisms use cues.	Plastic responses such as changes in development, behaviour and allocation of resources to competing demands can allow individuals to match their phenotypes (or those of their offspring, in the case of plastic maternal effects) to spatial or temporal variations in their abiotic and biotic environments.
		For phenotypic plasticity to be effective organisms must be able to accurately forecast environmental challenges affecting their fitness.
Colour of environment	Refers to the power spectrum of a stochastic environmental signal	Pink or red noise corresponds to variation that has more power at
al noise	estimated by a Fourier analysis of the	low frequencies, <b>white noise</b> is
(spectral	signal. By analogy to light, the colour	temporally uncorrelated and
colour)	refers to the profile of power across the signal's frequency spectrum [140].	variance is spread equally across all frequencies [103,140].

853 854 Table 2. Examples of anticipatory mechanisms and internal models (correlations) on which they

rely.

Example	Internal model
Circadian clocks in microbes, plants, mammals [45,149] allow organisms to time physiological processes.	Correlation between clock time and diurnal day/night cycle. Gene regulatory networks and metabolic pathways link the clock to particular biological processes, ensuring they peak at the appropriate times of day or night.
Toads sense water levels in temporary ponds, allowing them to switch to rapid metamorphosis [150] before ponds dry out.	Correlation between water level and time to pond drying.
Maternal light environment of understory forest herbs influences offspring life history and fitness, an example of anticipatory parental effects [151].	Correlation between maternal light environment and offspring light environment
Reaching a critical short photoperiod is a cue used by boreal and temperate trees to stop growing in the autumn [152–154].	Correlation between photoperiod and impending winter conditions.
Negative phototaxis and daily vertical migration in <i>Daphnia</i> , <i>Artemia</i> [155] and marine invertebrates (e.g. crab larvae, copepods) is cued by a change in light intensity, and allows invertebrates to avoid visual predators by swimming to darker areas. [156] [157]	Correlation between light intensity and predation risk.
Daphnia reared in the presence of a predator produce predator-resistant offspring [158]. Immune priming in plants allows increased resistance to pathogen infection following	Correlation between maternal kairomone environment and offspring predation risk. Correlation between pathogen exposure and likelihood of repeated exposure.

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