

Size-dependent eco-evolutionary feedbacks in fisheries

Eric Edeline^{1,2*}, Nicolas Loeuille¹

1: Sorbonne Université/UPMC Univ. Paris 06/CNRS/INRA/IRD/Paris Diderot Univ. Paris 07/UPEC/

5 Institut d'Ecologie et des Sciences de l'Environnement – Paris (iEES-Paris), 4 Place Jussieu, FR-75252
Paris, France.

2: ESE Ecology and Ecosystem Health, INRAE, Agocampus Ouest, 35042 Rennes, France.

* Corresponding author: eric.edeline@inrae.fr

Abstract

Harvesting may drive body downsizing along with population declines and decreased harvesting yields. These changes are commonly construed as consequences of direct harvest selection, where small-
15 bodied, early-reproducing individuals are immediately favoured. However, together with directly selecting against a large body size, harvesting and body downsizing alter many ecological features, such as competitive and trophic interactions, and thus also indirectly reshape natural selection acting back on body sizes through eco-evolutionary feedback loops (EEFLs). We sketch plausible scenarios of simple EEFLs in which one-dimensional, density-dependent natural selection acts either
20 antagonistically or synergistically with direct harvest selection on body size. Antagonistic feedbacks favour body-size stasis but erode genetic variability and associated body-size evolvability, and may ultimately impair population persistence and recovery. In contrast, synergistic feedbacks drive fast evolution towards smaller body sizes and favour population resilience, but may have far-reaching bottom-up or top-down effects. We illustrate the further complexities resulting from multiple
25 environmental feedbacks using a co-evolving predator-prey pair, in which case outcomes from EEFLs depend not only on population densities, but also on whether prey sit above or below the optimal predator/prey body-size ratio, and whether prey are more or less evolvable than their predators. EEFLs improve our ability to understand and predict nature's response to harvesting, but their integration into the research agenda will require a full consideration of the effects and dynamics of natural selection.

30

Key words: Body size, Co-evolution, Competition, Density-dependent selection, Eco-evolutionary feedbacks, Global changes, Harvesting, Natural selection, Predation.

Glossary

- 35 **Absolute fitness:** number of offspring reaching the reproductive stage.
- Evolutionary deterioration:** evolutionary change leading to smaller population densities, thereby increasing its probability of extinction (e.g., due to demographic stochasticity).
- Evolutionary rescue:** adaptive evolutionary change that restores positive growth to declining populations and prevents extinction.
- 40 **Evolvability:** trait potential to evolve.
- Fitness landscape:** multidimensional surface depicting fitness as a function of phenotypic traits.
- Relative fitness:** absolute fitness of a given phenotype divided by average absolute fitness of all phenotypes in the population.
- Selection gradient:** Trait-specific slope of the fitness landscape, i.e., holding other traits constant.

45

Introduction

The management of exploited populations is classically based on density-dependent population models in which harvesting, while decreasing population size, also relaxes density-dependent competition so that individual biomass productivity is increased (Verhulst 1838, Schaefer 1954, Hilborn and Walters 50 1992). However, this classical view has been repeatedly challenged by studies showing that individual biomass productivity often tends to decrease, not to increase, with harvesting (Kristiansen and Svåsand 1998, Conover and Munch 2002, Swain et al. 2007, Edeline et al. 2007, 2009, Biro and Post 2008, Heino et al. 2013).

55 This negative relationship between harvest effort and individual biomass production is generally interpreted as a rapid evolutionary response to direct harvest selection against large-bodied individuals by fishers (Trippel 1995, Law 2000, Kuparinen and Merilä 2007, Fenberg and Roy 2008, Heino et al. 2015). Accordingly, selection against a large body size is expected to favour slow-growing and early-maturing genotypes, which also tend to have lower fecundity and decreased offspring quality (Walsh et al. 2006, Heino et al. 2013). However, cases remain where exploitation induces no phenotypic change (Hilborn and Minte-Vera 2008, Devine and Heino 2011, Silva et al. 2013, Marty et al. 2014), or a change towards larger body sizes as predicted by density-dependent population models (Hilborn and Minte-Vera 2008). Therefore, whether harvest-induced evolutionary changes occur at all, or are large and rapid enough to influence biomass productivity remains controversial (Andersen and Brander 65 2009, Borrell 2013).

This debate, we feel, is plagued by a pervasive inclination of many researchers to overlook natural selection and to consider selective removal by harvesters as the only dynamic selective force at play. Natural selection, if ever mentioned, is regarded as negligible such that harvest-induced changes are 70 widely considered as slowly reversible (see e.g. the seminal paper by Law 2000). The simplifying assumption that natural selection is negligible further leads to conclude, for instance, that body-size stasis in harvested populations indicates that evolution is absent or has unimportant effects relative to the effects of ecology (e.g., Hilborn and Minte-Vera 2008). As we propose below, body-size stasis may in fact reflect eco-evolutionary dynamics in which natural selection opposes the effects of direct 75 harvest selection. Failure to account for these eco-evolutionary processes might fundamentally hamper our ability to understand and, hence, to manage productivity dynamics in harvested populations.

The objective of this essay is to provide an impetus to the study of natural selection in harvested populations, and may be seen as a complementary follow-up to Kinnison et al. (2015). These authors
80 stressed that our inclination to seek for parsimonious explanations may drive us to overlook the action of complex eco-evolutionary dynamics. Following this idea, we review theoretical, experimental and empirical insights to elaborate plausible scenarios under which complex eco-evolutionary dynamics may affect the yield and resilience of harvested populations in outcomes that may appear very similar to purely ecological or purely evolutionary dynamics. Most of our examples come from fisheries
85 which, because of their ecological, economic and social importance, concentrate the majority of the literature devoted to harvested animal populations.

We first review the mechanisms through which natural selection may favour either large-bodied or small-bodied individuals. Secondly, we build on this knowledge of natural selection to identify the
90 different pathways and directions that size-dependent EEFLs may take in harvested populations when only one single species evolves. In the third section, we extend the approach to EEFLs acting at the two-species and food-web levels. Finally, in the fourth section we conclude with an overview of the methods currently available to advance our empirical knowledge of EEFLs and with a consideration of how EEFLs may change our approach to managing harvested populations.

95

1. Size-dependent natural selection

Ample evidence shows that trait evolution in response to natural selection may be large and fast, hence far from negligible (Hendry and Kinnison 1999, 2001, Grant and Grant 2002, Stockwell et al. 2003,

100 Hairston et al. 2005, Carroll et al. 2007). This section provides details of the mechanisms through which natural selection moulds body sizes.

Table 1. Sources of natural selection predicted and observed to favour either a smaller or larger body size at age or at maturity.

105

Natural selection for a Smaller body size		Natural selection for a Larger body size	
Exploitative competition ¹	(De Roos et al. 2003, Kooijman 2010)	Interference competition	(Le Bourlot et al. 2014)
Long-term food stress ²	(Arendt 1997)	Long-term food stress ²	(Gadgil and Bossert 1970)
Selective predation on large-bodied individuals in prey populations	(Gårdmark and Dieckmann 2006, Heino et al. 2015)	Seasonal food stress	(van de Wolfshaar et al. 2008)
Size-independent predation	(Abrams and Rowe 1996, Gårdmark and Dieckmann 2006, Heino et al. 2015)	Selective predation on small-bodied individuals in prey populations ³	(Day et al. 2002)
Selective predation on juvenile (immature) individuals in prey populations	(Abrams and Rowe 1996)	Cannibalism	(Claessen et al. 2004)
		Selective predation on mature individuals in prey populations	(Ernande et al. 2004, Heino et al. 2015)

110 1: Note, however, that selection may be for a larger (not smaller) body size if attack rate increases faster with body size than maintenance metabolic rate. See text for more details. 2: long-term food stress has opposite effects on size-at-age and on size-at-maturity. 3: direction of selection may be different on size-at-age and on size-at-maturity, see text.

1.1 Natural selection for a smaller body size

Competition may be exploitative, i.e., resource-mediated (or indirect) or interference-mediated, i.e., direct. Both types of competition are expected to generate selection on body size, but only exploitative competition is expected to favour smaller body sizes. Exploitative competition may be usefully construed using the R^* rule, which states that competition selects individuals surviving on the lowest

equilibrium resource level (Tilman 1982). A lower individual R^* (i.e., a higher competitive ability) is achieved by increasing resource intake and/or by decreasing basal metabolic requirements. Note, however, that both resource intake and basal metabolic rate generally increase with body size (Peters 1983, Persson et al. 1998, De Roos et al. 2003, Kooijman 2010). Hence, whether individual R^* increases or decreases with body size depends on the relative strengths of allometric constraints acting on resource intake and metabolic rate. If resource intake increases faster with body size than metabolic rate, R^* decreases with increasing body size and exploitative competition should select for larger body sizes. In contrast, if resource intake increases slower than metabolic rate, R^* increases with body size and exploitative competition should select for smaller body sizes. In fish, available evidence suggests that R^* increases with body size (Persson and De Roos 2006), so that exploitative competition should favour smaller sizes. The argument extends to many other taxa if one assumes that ingestion increases with body surface ($\propto \text{size}^2$) while maintenance increases with body volume ($\propto \text{size}^3$) (Kooijman 2010). Population dynamics consistent with this prediction have been reported in the vendace *Coregonus albula* (Hamrin and Persson 1986), roach *Rutilus rutilus* (Persson et al. 1998) and Japanese medaka *Oryzias latipes* (Edeline et al. 2016).

Competition, if not leading to competitive exclusion, may also select on body sizes indirectly through decreasing the individual resource share on the long term. Available evidence suggests that such food stress has opposite effects on somatic growth rate and age at maturation. Across a wide variety of taxa, food stress favours slower growth rates and smaller size at age, presumably by imposing energy reallocation to the most vital functions (Arendt 1997). In contrast, fitness-maximising models predict that food stress should select for delayed maturation and, hence, for larger size at maturity if somatic growth rate is constant (Gadgil and Bossert 1970), a prediction supported by available empirical

140 evidence (Holliday 1989, Sgrò and Partridge 2000). Therefore, if somatic growth and maturation trade off, the growth-mediated and maturation-mediated effects of food stress on body size oppose each other, and are thus likely to remain inconsistent or cryptic.

Competition is not the only ecological interaction that may select for smaller body sizes. Predators that
145 target large-bodied prey directly select for smaller prey body sizes just like harvesters do (see above). This is for instance the case for fish predation on zooplankton (Brooks and Dodson 1965). If predators are non size-selective, predators still favour earlier maturation in prey and, hence, a smaller size at maturity if somatic growth is constant (Abrams and Rowe 1996). This is because early-maturing individuals have an increased fitness advantage when life expectancy is reduced (Gårdmark and
150 Dieckmann 2006, Heino et al. 2015). Finally, if predation mortality is stage-dependent, higher juvenile (immature) mortality favours earlier maturity which, given a fixed somatic growth rate, also means maturity at a smaller body size (Abrams and Rowe 1996, Heino et al. 2015).

1.2. Natural selection for a larger body size

155 In the wild, survival often increases with larger body sizes (Roff 1992), as demonstrated for instance in juvenile fish (Perez and Munch 2010, Stige et al. 2019), juvenile Soay sheep (*Ovis aries*, Hunter et al. 2018), or adult fish (e.g., Carlson et al. 2007, Olsen and Moland 2011). The mechanism behind this positive survival-size relationship could involve a higher resistance to starvation in larger-bodied individuals (van de Wolfshaar et al. 2008), but also results from strong interference in competitive
160 interactions. While size-selective effects of exploitative competition are dependent upon the allometric scaling exponents of intake and maintenance rates (see above), interference competition almost

universally brings an advantage to large-sized individuals in contests for food (Persson 1985, Post et al. 1999). In fish, dominance hierarchies are highly size-dependent both among and within species (Griffiths et al. 2020, Fausch et al. 2021). In experimental populations of the springtail *Folsomia*
165 *candida*, interference favours large-sized individuals that can monopolize resources (Le Bourlot et al. 2014). Similarly, in wild populations of the brown anole lizard *Anolis sagrei* natural selection for larger body sizes increases in parallel with population density and associated interference competition (Calsbeek and Smith 2007). In these lizards, the strength of competition-induced selection on body size overwhelmed the strength of predation-induced selection (Calsbeek and Cox 2010).

170

Often, predators are size-limited and thus preferentially feed on small-sized prey individuals. This is true for both aquatic and terrestrial systems (Sinclair et al. 2003). In such cases, predators favour prey individuals that grow fast through a “predation window” to rapidly reach a size refuge, i.e., they select for a large body size at a given age (Day et al. 2002). This predation window plays a key role in
175 mediating the population dynamic effect of intraspecific predation (i.e., cannibalism), an interaction that is present in multiple aquatic or terrestrial taxa (Fox 1975, Claessen et al. 2002, 2004). Cannibalism is presumably the mechanism that controlled the positive effect of population density on somatic growth rate in the pike (*Esox lucius*) population of Windermere where, as the density of cannibals increased, survival was biased towards faster-growing individuals (Edeline et al. 2007, 2009).

180

The effect of size-limited predation on age at maturation is less straightforward than on somatic growth. If mortality increases among small-sized individuals, predictions depend on the details of the model. Optimality models predict evolution of delayed maturation at a larger body size (Taylor and Gabriel 1992). In contrast, adaptive dynamics models accounting for a trade off between somatic

185 growth and reproduction and for a positive effect of body size on fecundity lead to more complex
 outcomes: increased mortality among small-sized individuals can increase or decrease maturation size,
 or even lead to the coexistence of both early- and late-maturing individuals when benefits from early
 maturation collide with benefits from growing fast to a size refuge (Gårdmark and Dieckmann 2006).
 The outcome of evolution also depends on whether food availability is sufficient to support fast
 190 somatic growth (Chase 1999). To our knowledge, the available empirical and experimental evidence
 more often supports delayed maturation at a larger body size when predation targets small-sized
 individuals (Edley and Law 1988, Wellborn 1994, Beckerman et al. 2010, Le Rouzic et al. 2020).

Finally, a larger body size further provides females with a higher fecundity in egg-spawning species
 195 (Barneche et al. 2018), and males with a strong advantage in contest sexual selection (e.g., Fleming and
 Gross 1994). Combined together, these multiple positive effects of natural selection on body sizes are
 likely to outweigh the negative effects, as suggested by an overall tendency for natural and sexual
 selection to favour larger body sizes across multiple taxa (Kingsolver and Pfennig 2004). We now
 move to examining how such size-dependent natural selection may interact dynamically with ecology
 200 in EEFLs.

2. Theory and scenarios of harvest-induced EEFLs with one evolving species

Box1. Defining the selection- and evolvability-mediated pathways to eco-evolutionary feedback loops (EEFLs).

205 To study existing feedbacks between ecological and evolutionary dynamics, two main frameworks are
 currently used: quantitative genetics (QG) and adaptive dynamics (AD). Though the two methods

differ, they are both based on the idea that the description of trait dynamics in response to selection requires two fundamental ingredients: trait(s) evolutionary potential (hereafter “evolvability”) and a measure of selection acting on the trait(s) (Abrams 2001).

210

Consider for instance the classical recursive equation of quantitative genetics (QG):

$$\bar{x}_{t+1} = \bar{x}_t + \Delta \bar{x} = \bar{x}_t + \underbrace{Va_x}_{\text{Evolvability}} \underbrace{\frac{\text{COV}(w, x)}{\text{var}(x)}}_{\text{Selection}} \quad \text{Eq. 1,}$$

215

where \bar{x} is the mean population value of a univariate trait x , t is generation index, Va_x is

additive genetic variance, w is relative individual fitness, and $\frac{\text{COV}(w, x)}{\text{var}(x)}$ is the directional

selection gradient, i.e., the slope of the linear regression between relative fitness and trait x (Lande

and Arnold 1983). Provided that the definition of w includes at least density dependence and/or

frequency dependence, Eq. 1 incorporates selection-mediated EEFLs as the ecological context (density

220

or frequency) then impacts the selection term (Abrams 2001). Eco-evolutionary feedback loops may

also occur through the evolvability-mediated pathway in Eq. 1, for instance if Va_x is directly linked

to the demographic context (e.g., an existing correlation between population density and genetic

variability) or if Va_x is an explicit function of the strength of selection since strong directional

selection is expected to decrease additive genetic variances (Crow 2008).

225

Adaptive dynamics (AD) (Dieckmann and Law 1996) readily account for both selection- and evolvability-mediated EEFLs. This essential feature of AD is captured by the canonical equation:

$$\frac{d\bar{x}}{dt} = \underbrace{\frac{1}{2}\mu\sigma^2 N^*(x)}_{\text{Evolvability}} \underbrace{\frac{\partial W(x',x)}{\partial x'}}_{\text{Selection}} \Big|_{x'=x} \quad \text{Eq. 2,}$$

230

where x is a resident trait, x' is a mutant trait, $d\bar{x}/dt$ is a continuous-time analogue of $\Delta\bar{x}$ in Eq. 1, μ is per capita mutation rate, and σ^2 is phenotypic variance from a mutation. $N^*(x)$ is equilibrium population size for the resident trait, and Eq. 2 hence incorporates the evolvability-mediated pathway to EEFLs since evolvability, here determined by the mutation process, is explicitly dependent on equilibrium population size $N^*(x)$, which is set by the value of the resident trait x .

235

$W(x',x)$ is invasion fitness for a mutant trait x' in an environment determined by the resident trait x . Because this fitness definition is based on ecological dynamics, one sees that selection-mediated EEFLs are readily considered in adaptive dynamics models. Finally $(\partial W(x',x))/(\partial x')$ is the directional selection gradient acting on the mutant trait x' , i.e., is the invasion criterion (slope of the fitness landscape for x' evaluated in x).

240

Theory presented in Box 1 predicts that EEFLs may proceed through two different pathways: a selection-mediated and an evolvability-mediated pathways, which we illustrate in Fig. 1. The selection-mediated pathway is captured by Arrow 1 (Fig. 1): the environment of an individual generates natural selection on body size (see Section 1 above). In addition to influencing individual fitness and, from there, population densities, body size has widespread and consistent ecological effects (Peters 1983,

245

Brown et al. 2004, Woodward et al. 2005). Hence, selection-induced change in body size, in turn, may impact the environment through the size-dependency of reproductive success and ecological interactions (Arrow 2). The evolvability-mediated pathway to size-dependent EEFLs is captured by
 250 Arrow 3, and involves mutation-limitation effects linked to population sizes and/or any other existing correlations between genetic diversity and population size (Box 1, Frankham 1996). Such evolvability-mediated pathways to EEFLs are often neglected, but may actually be important from a management or a conservation point of view (Carlson et al. 2014, Marty et al. 2015, Kuparinen and Hutchings 2017). Harvesting may trigger or disrupt size-dependent EEFLs through both direct harvest selection on body
 255 size and through the removal of conspecifics and possibly also heterospecifics (Fig. 1).

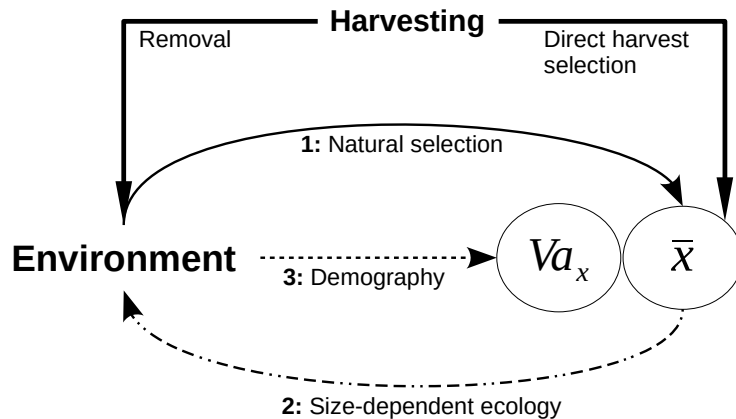


Figure 1: Selection-mediated and evolvability-mediated pathways to size-dependent eco-evolutionary feedback loops (EEFLs). Va_x and \bar{x} stand for additive genetic variance and mean body size, respectively, in the harvested population (Box 1). Arrow 1: natural selection and the selection-mediated pathway to EEFLs, Arrow 2: body size-dependent ecological processes, Arrow 3: Effects of demography on genetic variability and the evolvability-mediated pathway to EEFLs (Box 1). NB 1: phenotypic plasticity, as also captured by Arrow 1, will not be discussed. NB 2: For

the sake of simplicity, we did not consider the potential direct effects of selection on additive genetic variances (e.g. Crow 2008).

In order to fully grasp the basic ideas that underpin size-dependent EEFLs in the system depicted by Fig. 1, we provide a graphical representation of a moving adaptive landscape in Fig. 2. For simplicity,
 260 the fitness landscapes represented on fig 2 ignore frequency dependent selection, so that each phenotype has a given fitness irrespective of its frequency. This fitness would be representative of the

absolute fitness of the corresponding monomorphic population. Relative fitness of this phenotype confronted to another can then simply be read from the relative position on the fitness landscapes, phenotypes with higher fitness being selected. Therefore this representation, though it simplifies the
265 underlying ecological aspects, allows to assess absolute and relative fitness easily. We make this choice because population persistence depends on absolute, not relative fitnesses, and because absolute fitness is therefore more intuitively linked with management aspects. For more discussion on the link between absolute and relative fitness, see Orr (2007).

270 Second, we represent what we believe are the most generic functional forms for absolute fitness functions: harvest selection is often directional against a large body size, and body size is often constrained by trade-offs resulting in stabilizing natural selection (Kingsolver et al. 2001, Kingsolver and Pfennig 2004, Carlson et al. 2007). Note that this assumption of stabilizing natural selection also follows from the general observation that evolution towards smaller body sizes is associated with
275 severe fitness costs (e.g., Walsh et al. 2006), while largest-bodied and oldest individuals may be subject to senescence. Although these settings are very simple and maybe rare in nature, their evolutionary outcome is more easily visualized than when multiple environmental feedbacks operate simultaneously and prevent evolutionary optimization (Metz et al. 2008).

280 In the absence of any direct harvest selection, the population mean body size resides at the naturally-selected body size optimum (dotted blue curve, t_0 in Fig. 2A). The product of natural selection with direct harvest selection (i.e., survival to harvesting, dashed red curve) instantaneously warps the naturally-selected fitness landscape to generate a new, composite fitness landscape (solid grey curve) on which the population mean trait value is associated with a fitness at which the population crosses the

285 extinction threshold (t_1 , maladaptation). Rapid adaptive evolution through a few generations towards the newly-selected adaptive optimum restores a fitness at which the population may persist (t_2 , re-adaptation). If adaptive change occurs fast enough, it may potentially restore a positive population growth and prevent extinction, a process termed “evolutionary rescue” (Glossary, Gomulkiewicz and Holt 1995).

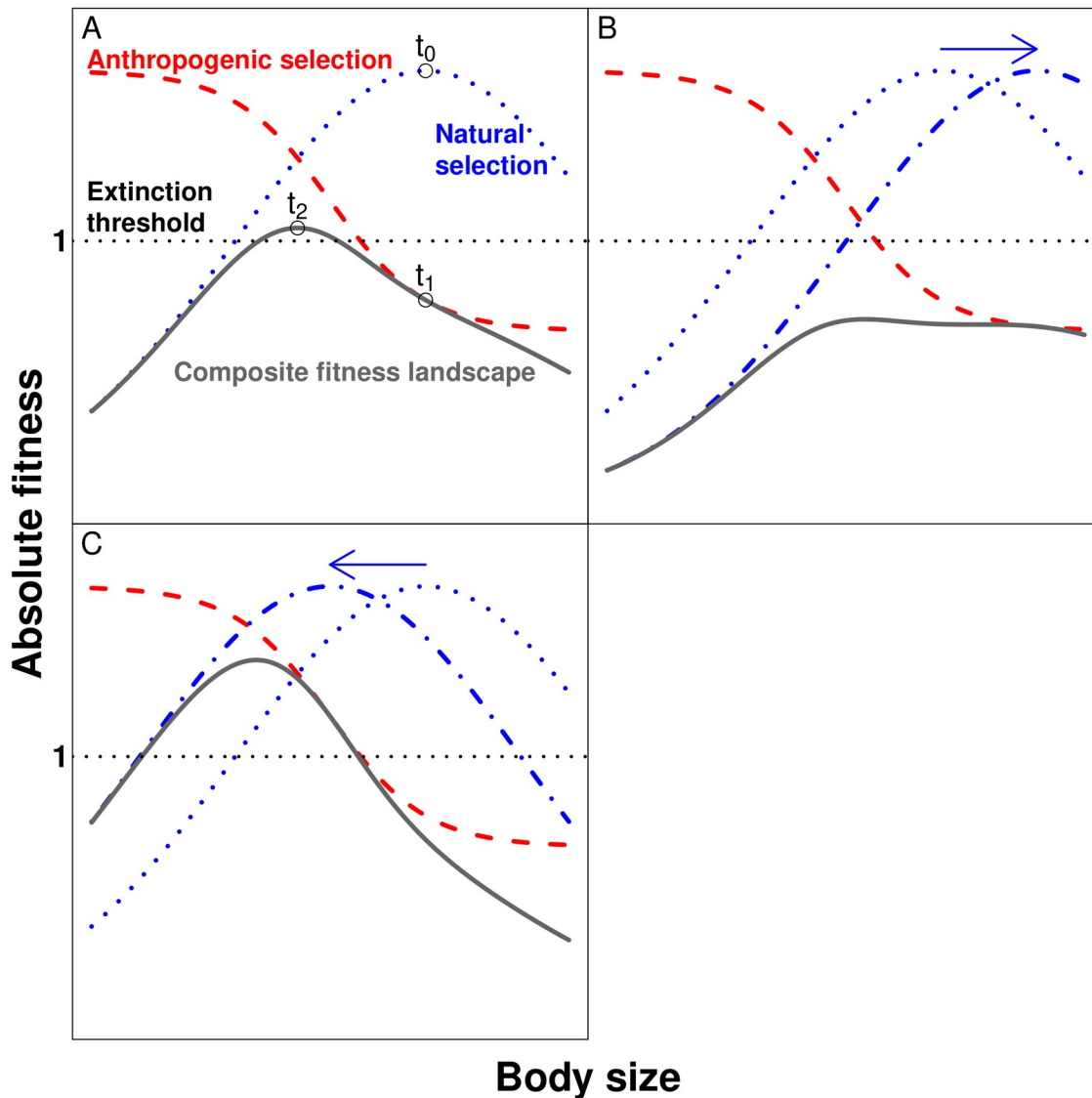


Figure 2: Eco-evolutionary feedbacks in harvested populations. Curves show the relationship between absolute fitness and body size, and the horizontal dotted line shows unity absolute fitness (extinction threshold). **A:** Evolutionary “rescue” (see Glossary) with no eco-evolutionary feedback loop (EEFL). Open circles show the

mean phenotype in the population. **B:** A single density parameter feedbacks on natural selection, generating an antagonistic EEFL. The arrow shows the change in directional natural selection due to the environmental feedback. **C:** The one-dimensional density-dependent feedback generates a synergistic EEFL in which natural selection changes to reinforce the effect of direct harvest selection.

290

The model presented in Fig. 2A makes the simplifying assumption that natural selection does not respond to harvesting. However, in addition to imposing direct harvest selection on body size, harvesting also alters the environment (Fig. 1, Arrow 1) and may thus indirectly change natural selection acting on body size (Bouffet-Halle et al. 2021). We will now examine two scenarios in which
 295 such harvest-induced changes in natural selection either oppose or reinforce the action of direct harvest selection on body size.

2.1. Antagonistic EEFLs

We first consider a feedback in which harvesting changes natural selection towards favouring *larger-*
 300 *than-initial* body sizes (sketched in Fig. 2B). As this selection acts in opposite ways to the direct selective effects of harvesting, we refer to this situation as an **antagonistic EEFL**. Compared to an EEFL-absent case, antagonistic EEFLs magnify warping of the adaptive landscape and thus impair population persistence. At an extreme, the fitness peak may dwindle below the extinction threshold (Fig. 2B). Antagonistic EEFLs are expected whenever density-dependent natural selection favours
 305 small body sizes and harvesting, through reducing densities, relaxes natural selection for a small body size (Fig. 2B). For instance, reduced population densities may relax exploitative competition for resources, and weaken associated selection for smaller body sizes (Table 1). Antagonistic EEFLs may also emerge from changes in predation regimes, as demonstrated by Gårdmark et al. (2003) using a theoretical model in which an age-structured population evolves in response to both harvesting and
 310 predation mortality. Harvesting reduces prey availability so that predator density decreases, thus

inducing relaxed predation and the associated natural selection for smaller body sizes. This result is likely to apply whenever predators of the harvested population directly select for smaller body sizes, i.e., when predators preferentially prey on large-bodied individuals, on juveniles, or when they are non size-selective (Table 1).

315

A hallmark of eco-evolutionary dynamics is their tendency to remain cryptic if they are not anticipated and, hence, not specifically investigated (Kinnison et al. 2015). Size-dependent, antagonistic EEFLs are no exception, because the changes in natural selection oppose the effects of direct harvest selection and favour body-size stasis, an outcome that may erroneously be interpreted as direct harvest selection being too weak to drive any evolutionary response (e.g., Hilborn and Minto-Vera 2008). In fact, however, body-size stasis of antagonistic EEFLs is associated with a fitness drop that may ultimately prevent evolutionary rescue (Fig. 2B). The fitness drop and resultant decreased population size may further jeopardize body-size evolvability (Box 1, Arrow 2 → 3 sequence in Fig. 1) which, together with a vanishing strength of selection due to a flat composite fitness landscape (Fig. 2B), decreases the probability for recovery. Overall, any situation in which harvesting is associated with body-size stasis but severe population decline may be suspected to reflect an antagonistic EEFL.

320
325

2.2. Synergistic EEFLs

Synergistic EEFLs occur when the environmental feedback changes natural selection towards favouring smaller-than-initial body sizes in synergy with direct harvest selection (Fig. 2C). Synergistic EEFLs may result, for instance, when harvesting, through reducing the density of large-sized individuals in the population, relaxes interference competition and cannibalism and associated natural selection for a large body size (Table 1). Recent experimental evidence in replicated fish populations

330

suggests that this harvest-induced relaxation of interference competition and cannibalism can drive a rapid evolutionary divergence between harvested and non-harvested populations (Bouffet-Halle et al. 2021). Synergistic EEFLs are also expected when predation favours larger body sizes and predators disappear due to harvest-induced prey shortage (Table 1, Jusufovski and Kuparinen 2020).

Qualitatively, the phenotypic outcome from synergistic EEFLs looks similar to the phenotypic outcome from EEFL-absent dynamics (Fig. 2A), though directional selection is stronger and expected trait variation faster. Synergistic EEFLs are thus likely to remain cryptic and to be interpreted as a large and rapid response to direct harvest-selection acting alone (e.g., Darimont et al. 2009). Compared to an EEFL-absent situation (Fig. 2A), however, synergistic EEFLs result in a magnified fitness peak on the composite fitness landscape (Fig. 2B) and, hence, favour larger population sizes at the body-size optimum and higher body-size evolvability (Arrow 2 \rightarrow 3 sequence in Fig. 1). Hence, synergistic EEFLs may favour evolutionary rescue and allow fast evolutionary rebound after relaxation of fishing. This is presumably the configuration that explains why pike, a highly cannibalistic species, showed a fast and large evolutionary response to varying harvesting intensity in Windermere (Edeline et al. 2007, Coltman 2008). Finally, synergistic EEFLs increase slope steepness around the fitness peak on the composite fitness landscape (Fig. 2C), resulting in stronger selection around and faster evolution towards the body-size optimum. Therefore, synergistic EEFLs are consistent with the observation that fishing-induced trait changes are often much faster than predicted by theoretical models that only assume direct harvest selection (Audzijonyte et al. 2013a).

These simple scenarios of antagonistic and synergistic EEFLs focus on the evolution of just one harvested species alone. Fisheries, however, most often target not just one but several species within

the ecological network, so that an ecosystem perspective on fishery management is required (White et al. 2012, Perälä and Kuparinen 2020). Therefore, we now move to examining EEFLs when more than one species evolves.

360

3. Scenarios of harvest-induced EEFLs with multiple evolving species

There is currently an emerging recognition that evolution in a given harvested species can induce co-evolution in other species through changes in ecological interactions (Wood et al. 2018). However, understanding the evolutionary response to harvesting in a multispecific context is highly challenging (Audzijonyte et al. 2013b). Investigation on multispecies EEFLs requires to account simultaneously for the coevolution of the various body sizes, of the network structure, and to consider how one feeds back on the other (Loeuille and Loreau 2005). Direct data investigating the occurrence and magnitude of multispecies EEFLs are scarce. However, different empirical facts suggest that multispecies EEFLs may naturally emerge in exploited ecological networks.

370

First, empirical data suggest that predators are often larger than their prey in both terrestrial and aquatic systems (Cohen et al. 2003, Sinclair et al. 2003, Brose et al. 2006) and that predator-prey body-size ratios determine the strength of predation (Emmerson and Raffaelli 2004, Renneville et al. 2016). Therefore, we expect that selection on body size will change the distribution of interaction strengths, which largely constrains ecosystem functioning and stability (e.g., McCann et al. 1998, Rooney et al. 2006). Ultimately, rewiring and redistribution of interaction strengths may lead to extinctions in the network. For instance, evolution of larger body sizes can decrease the density of the evolving population thereby increasing its vulnerability to demographic stochasticity and potentially facilitating

its extinction (evolutionary deterioration, see Glossary). At the same time, body-size evolution may
 380 undermine predator persistence through weakening trophic links, or compromise prey persistence
 through strengthening trophic links. Similarly, variations in interaction strengths will affect apparent
 competition (Holt et al. 1994), thereby changing coexistence conditions within the food web and
 possibly leading to competitive exclusions. Because the network structure, in turn, constrains the
 fitness of species within the community, multispecies EEFLs naturally emerge (Fig. 3).

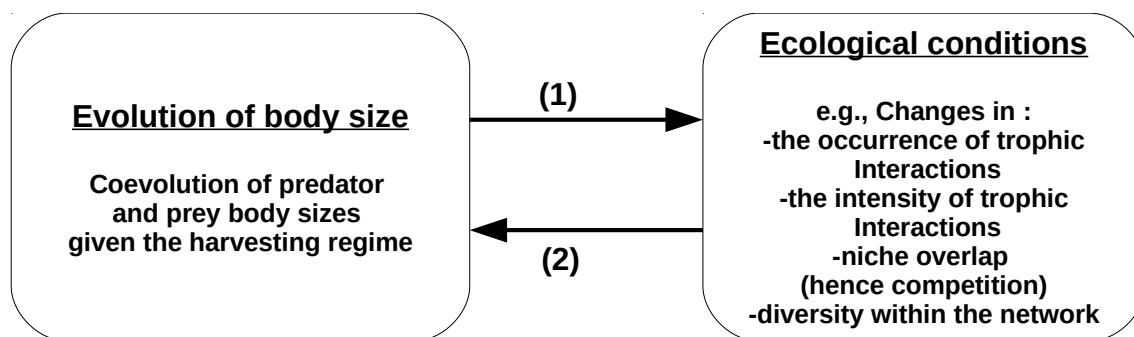


Figure 3: Components of multispecies EEFLs in food webs. (1) Coevolution of body sizes within the network under the new selective regime (harvesting) affects interaction patterns within the network, as well as coexistence conditions. (2) This new ecological context in turn changes the fitness of species (eg, due to changes in predation rates), thereby acting on body size coevolution.

Co-evolution strongly complicates EEFLs. In particular, the graphical framework from Fig. 2 no longer
 applies, because the environment now becomes multidimensional and evolution no longer optimizes
 fitness or population size of any given species (Meszéna et al. 2001, Metz et al. 2008). To keep our
 arguments as simple as possible we focus on a single co-evolving predator-prey pair, in which we
 390 examine two non-exclusive mechanisms for the emergence of EEFLs. We first examine the
 consequences of a “trophic relaxation”, which occurs when decreased densities weaken the strength of
 the predator-prey link. Second, we examine the consequences of predators and prey having

“asymmetric evolvability” for body size and, hence, evolving at a different pace in response to harvesting. For both mechanisms we consider that, before harvesting starts, the predator-prey pair resides at an evolutionary equilibrium.

3.1. Trophic relaxation

Prey may be either smaller or larger than the preferred prey size of the predator. Prey sizes matching the preferred size are rarely expected, as prey may evolve away from such situations, but also because the distribution of body sizes does not usually maximize trophic interactions due to metabolic constraints, competition and the multiplicity of prey and predator species that also act as selective pressures (e.g., Loeuille and Loreau 2005, 2006).

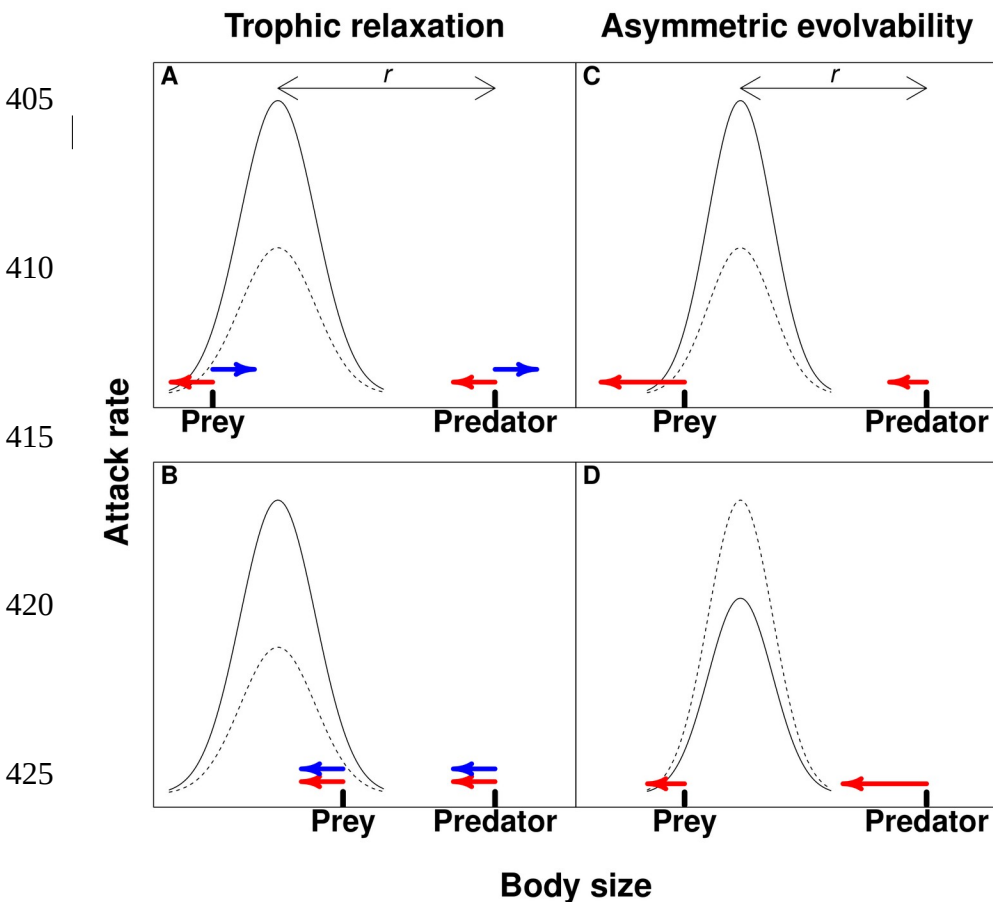


Figure 4. Alternative settings in coevolving predator-prey pairs. The Gaussian curves show the predation intensity before (solid lines) and after (dashed lines) harvesting starts. Optimal prey size is set by the fixed ratio r . Red horizontal arrows show potential for body size evolution (i.e., body-size evolvability) in response to direct harvest selection. Blue arrows show natural-selection response to harvesting, i.e., the EEFL. **A:** Trophic relaxation leading to increased natural selection for a larger body size. **B:** Trophic relaxation leading to natural selection for a smaller body size. **C:** Body-size evolvability is larger in the prey than in the predator. **D:** Body-size evolvability is larger in the predator than in the prey.

We now start harvesting both the prey and predator which, hence, are both under direct harvest selection for a smaller body size (red arrows in Fig. 4). In Figs. 4A and 4B, red arrows have similar
430 lengths indicating that both species evolve smaller body sizes at a similar pace, such that no change is to be expected in their realized body-size ratio. However, because harvesting reduces population density in both the prey and predator, we expect a relaxation in the strength of the predator-prey link. This “trophic relaxation” is outlined in Figs. 4A and 4B by a decreased predation intensity (dotted Gaussian curves). Such a trophic relaxation may lead to opposite eco-evolutionary outcomes depending
435 on whether initial prey size is smaller or larger than the predator’s optimal prey size.

In Fig. 4A, prey size is initially smaller than optimal for the predator, and the trophic relaxation thus results in relaxed natural selection for smaller body sizes in both the prey and predator (blue arrows). In other words, change in natural selection acts in opposition with direct harvest selection in an
440 antagonistic EEFL. In case 4B, in contrast, prey size is initially larger than optimal predator size, and the trophic relaxation thus results in relaxed natural selection for larger body sizes, thus creating a synergistic EEFL. Of course, these relatively simple outcomes are complicated by feedbacks from intraspecific interactions (competition, cannibalism) that may either reinforce or oppose the effects of the predator-prey feedback (see Section 2).

445

3.2. *Asymmetric evolvability*

Assuming symmetric evolvability in the prey and predator (Figs. 4A and 4B) is likely unrealistic for most situations. Rather, body-size may be more evolvable in prey than predators (hence the longer red arrow in Fig. 4C), either because the trait is determined by different gene networks for the two species,
450 or because the two species have very different population sizes, hence differing in accumulation of

mutations or standing genetic variability. For instance, smaller (prey) body sizes are often associated with larger population numbers (Woodward et al. 2005) and with a higher genetic variability (Romiguier et al. 2014, De Kort et al. 2021). Under these settings, prey evolve smaller body sizes faster than their predator, move further away from predator's preferred prey size, and ultimately benefit from an evolution-induced trophic relaxation (Fig. 4C). The predator on the other hand, may become resource limited, so that further declines in predator population are expected. This is different from trophic relaxation in cases 4A and 4B which was the driver of evolution.

In Fig 4D, we sketch an opposite, perhaps less common situation in which predators have a higher body-size evolvability than their prey. This configuration may potentially result from prey being close to a lower evolutionary limit for body size (Le Rouzic et al. 2020, Renneville et al. 2020). Under these settings, predators evolve smaller body sizes faster than prey, such that preferred prey size moves closer to prey size and a trophic magnification results. Such a coevolution therefore favours the maintenance of the trophic interaction. Note that these outcomes depend on prey being smaller than predator's preferred prey size in Figs. 4C and 4D, and are reversed when prey are larger than the preferred prey size of the predator (i.e., trophic magnification in Fig. 4C and trophic relaxation Fig. 4D).

3.3. *More complex interaction networks*

In more complex networks, the multiplicity of trophic and non-trophic interactions may generate a variety of counteracting selection gradients, so that evolution might be more constrained than in a single predator-prey link. If this hypothesis is true, EEFLs might well be more important in explaining evolutionary and ecological stasis rather than change (Ellner et al. 2011, Strauss 2014, Kinnison et al.

2015). Beyond very specific scenarios, network and eco-evolutionary complexities under harvesting
475 scenarios are virtually impossible to grasp intuitively, and are even hard to handle through a
mathematical analysis. However, numerical simulations are certainly possible. In this regard, the
development of evolutionary models of food webs based on body size offer promising venues, as they
already consider simultaneously evolution of body size and changes in the network structure (Loeuille
and Loreau 2005, 2009, Brännström et al. 2011, Allhoff et al. 2015). Harvesting scenarios may be
480 implemented in such models (Perälä and Kuparinen 2020), as has been done in other contexts (eg,
climate warming, Weinbach et al. 2017, Yacine et al. 2020).

4. Management and perspectives

So far, the vast majority of models used to project the eco-evolutionary consequences of fishing ignore
485 natural selection on body size (but see Jusufovski and Kuparinen 2020). Hence, although quite
elaborated, these models are likely to underestimate either the demographic consequences of harvesting
when antagonistic EEFLs are involved, or the rates of evolutionary change and recovery when
synergistic EEFLs are involved. We recognize, however, that more empirical and experimental studies
are needed to document the pathways, directions and strength of density-dependent selection acting on
490 body size in harvested systems. In particular, it is important to document whether and when harvest-
induced EEFLs can be simplified into a one-dimensional, density-dependent process that can be
handled by optimality approaches such as that outlined in Fig. 2. In Box 2, we provide an overview of
the empirical methods currently available to progress in that direction.

495 **Box 2. Empirical exploration of size-dependent EEFLs: where to go next?**

Demonstrating a full selection-mediated EEFL requires showing both that natural selection drives evolutionary trait change and, in turn, that the resultant trait evolution alters the environment in such a way that natural selection acting back on the trait is modified (Figs. 1, 2). Considering also the evolvability-mediated pathways to EEFLs requires to further measure the effects of environmental changes on trait evolvability. Tackling such a complexity is challenging but, as we show below, not beyond of reach.

Measuring natural selection and trait response to selection

The form and strength of selection are most accurately measured by estimating fitness-traits relationships at the individual level (Arnold 2003) using, e.g., capture-recapture techniques. Alternatively, the directional component of selection may also be estimated from population and trait time series using the “Geber method”, the age-structured price equation or integral projection models (Hairston et al. 2005, Ellner et al. 2011, van Benthem et al. 2017, Govaert 2018). A drawback of all these methods is that they measure selection acting on phenotypes, while evolution is concerned only by selection acting on the heritable component of phenotypes (Morrissey et al. 2010). To circumvent this problem, statistical approaches making use of the “animal model” (AM) of quantitative genetics were developed to specifically measure selection acting on the additive genetic component of traits and, hence, to accurately predict evolution (Hadfield 2008, Morrissey et al. 2010, Stinchcombe et al. 2014). AM-based approaches require pedigree data and are thus more readily implementable in small, closed systems than in large-scale fisheries (but see Koch et al. 2008).

Measuring the dependency of natural selection on the environment (Fig. 1, Arrow 1)

A pivotal condition for the emergence of selection-mediated EEFLs is that natural selection dynamically changes due to changes in the environment (Figs. 1, 2, Govaert et al. 2019). This may be checked *a posteriori* through measuring genotype-by-food interactions on body sizes. For instance, Bouffet-Halle et al. (2020) used this approach to show that harvest-induced evolution towards smaller body sizes in experimental populations of medaka fish (*Oryzias latipes*) had evolved in a low-food but not in a high-food environment. This result suggested that medaka had evolved in response to density-dependent natural selection at high population density (low food), but not in response to direct harvest selection at low population density (high food). This approach, however, remains fragile because our understanding of genotype-by-food interactions remains limited, and other complementary results may be necessary to back-up conclusions from genotype-by-food analyses (Bouffet-Halle et al. 2021). When possible, selection-environment relationships should be measured directly using individual capture-recapture techniques (e.g., Haugen et al. 2007, Calsbeek and Smith 2007, Calsbeek and Cox 2010), keeping in mind the problems highlighted above of measuring selection at the phenotype level. Here also, these problems may be solved if the data permits applying the AM, which may be extended to estimate environment-selection relationships acting at the additive genetic level (Hunter et al. 2018).

535 *Measuring the trait dependency of ecological dynamics (Fig. 1, Arrow 2)*

Time series data may be used to quantify the feedback from phenotypic trait change to environmental variables. Since the inception of the Geber Method by Hairston et al. (2005) and Ellner et al. (2011), a multiplicity of more sophisticated methods have flourished. These methods are based either on inferring parameters for dynamic models from data (e.g., Rudy et al. 2017 and references therein), on non-parametric approaches such as Recurrent Neural Networks, or on hybrid approaches combining

differential equations with neural networks (e.g., Bonnaffé et al. 2020 and references therein).

Reviewing these methods is beyond the scope of this paper.

545 *Measuring the effects of the environment on trait evolvability (Fig. 1, Arrow 3)*

Evolvability may be measured using multiple metrics (e.g., Hansen et al. 2011, 2019), which condition approaches to exploring evolvability-mediated EEFLs. Here, we focused on additive genetic variance V_A which is a commonly-used measure of evolvability (Box 1), and which we assumed to be positively linked to population size (see Reed and Frankham 2001 for a contrasted view). Estimation of V_A relies on the AM, using either pedigrees or genetic markers of coancestry to construct relatedness matrices, with some caveats stressed by Lynch and Walsh (2018). The AM may further be extended to incorporate effects of environmental covariates on V_A in so-called random regression approaches (Schaeffer 2004).

550

555 Importantly, our review suggests that the ecological and evolutionary consequences of harvesting will largely depend on the ecological factors that regulate the population and, hence, will likely be constrained by the details of the local network context. However, based on our above analysis we may still propose some general management rules accounting for size-dependent EEFLs. As highlighted by Engen et al. (2014), a very general consequence of density-dependent selection is that the more ecologically-sustainable strategies will also produce the less evolutionary changes. Therefore, preventing population declines and alleviating evolutionary change are not independent lines of management but are instead highly intertwined management targets. If possible, management rules should further account for the probability of EEFLs to be either antagonistic or synergistic, because the

560

former are far more detrimental than the later to population persistence and recovery and, hence, would impose lower exploitation rates. Ideally, an *a priori* knowledge of the direction of density-dependent natural selection acting on body sizes could be gained using *had hoc* approaches (Box 2). Alternatively, a basic knowledge of the dominant ecological interactions could be used (Table 1).

In co-evolving predator-prey pairs, managers may also account for body-size ratios and potential asymmetries in body-size evolvability, so as to classify their harvested system into one of the four categories depicted in Fig. 4. Body-size ratios are well documented in the literature, and identification of a context prone to trophic relaxation or magnification should be relatively simple and lead to prudent exploitation. Prudent exploitation is also recommended if asymmetric body-size evolvability is suspected, especially when prey can escape predation (Fig. 4D), a situation in which exploitation rates should be stronger on the faster-evolving species so as to resorb asymmetry in evolvability. This recommendation somehow converges towards “balanced harvesting”, a management approach based on spreading fishing mortality across the widest possible range of species and sizes in proportion to their natural productivity. Interestingly, such balanced strategies have already been advocated to conciliate yield and sustainability even in models that ignore evolution (Tromeur and Loeuille 2017). Although more research is clearly needed to test whether and under which conditions these general recommendations hold true, we believe that far enough evidence is already available showing that a consideration of natural selection is highly needed if we are to improve our ability to accurately predict and manage the dynamics of harvested populations.

585 **Acknowledgements:** EE acknowledges financial support from the Norwegian Research Council
(projects EvoSize RCN 251307/F20 and REEF RCN 255601/E40) and from Rennes Métropole (project
AIS 18C0356).

Conflict of interest disclosure: The authors of this article declare that they have no financial conflict
590 of interest with the content of this article. NL is one of the PCIEcology recommenders.

REFERENCES

- Abrams 2001. Modelling the adaptive dynamics of traits involved in inter- and intraspecific interactions: an assessment of three methods. - *Ecol. Lett.* 4: 166–175.
- Abrams, P. A. and Rowe, L. 1996. The effects of predation on the age and size of maturity of prey. - *Evolution* 50: 1052–1061.
- Allhoff, K. T. et al. 2015. Evolutionary food web model based on body masses gives realistic networks with permanent species turnover. - *Sci. Rep.* 5: 10955.
- Andersen, K. H. and Brander, K. 2009. Expected rate of fisheries-induced evolution is slow. - *Proc. Natl. Acad. Sci. U. S. A.* 106: 11657–11660.
- Arendt, J. D. 1997. Adaptive intrinsic growth rates: an integration across taxa. - *Q. Rev. Biol.* 72: 149–177.
- Arnold, S. J. 2003. Performance surfaces and adaptive landscapes. - *Integr. Comp. Biol.* 43: 367–375.
- Audzijonyte, A. et al. 2013a. How fast is fisheries-induced evolution? Quantitative analysis of modelling and empirical studies. - *Evol. Appl.* 6: 585–595.
- Audzijonyte, A. et al. 2013b. Ecological consequences of body size decline in harvested fish species: positive feedback loops in trophic interactions amplify human impact. - *Biol. Lett.* 9: 20121103.
- Barneche, D. R. et al. 2018. Fish reproductive-energy output increases disproportionately with body size. - *Science* 360: 642.
- Beckerman, A. P. et al. 2010. The reaction norm of size and age at maturity under multiple predator risk. - *J. Anim. Ecol.* 79: 1069–1076.

- Biro, P. A. and Post, J. R. 2008. Rapid depletion of genotypes with fast growth and bold personality traits from harvested fish populations. - Proc. Natl. Acad. Sci. U. S. A. 105: 2919–2922.
- Bonnaffé, W. et al. 2020. Neural ordinary differential equations for ecological and evolutionary time series analysis. - Authorea in press.
- Borrell, B. 2013. Ocean conservation: a big fight over little fish. - Nature 493: 597–598.
- Bouffet-Halle, A. et al. 2021. Density-dependent selection mediates harvest-induced trait changes. - Ecol. Lett. in press.
- Brännström, Å. et al. 2011. Emergence and maintenance of biodiversity in an evolutionary food-web model. - Theor. Ecol. 4: 467–478.
- Brooks, J. L. and Dodson, S. I. 1965. Predation, body size, and composition of plankton. - Science 150: 28.
- Brose, U. et al. 2006. Consumer-resource body-size relationships in natural food webs. - Ecology 87: 2411–2417.
- Brown, J. H. et al. 2004. Toward a metabolic theory of ecology. - Ecology 85: 1771–1789.
- Calsbeek, R. and Smith, T. B. 2007. Probing the adaptive landscape using experimental islands: density-dependent natural selection on lizard body size. - Evolution 61: 1052–1061.
- Calsbeek, R. and Cox, R. M. 2010. Experimentally assessing the relative importance of predation and competition as agents of selection. - Nature 465: 613.
- Carlson, S. M. et al. 2007. Four decades of opposing natural and human-induced artificial selection acting on Windermere pike (*Esox lucius*). - Ecol. Lett. 10: 512–521.
- Carlson, S. M. et al. 2014. Evolutionary rescue in a changing world. - Trends Ecol. Evol. 29: 521–530.
- Carroll, S. P. et al. 2007. Evolution on ecological time-scales. - Funct. Ecol. 21: 387–393.
- Chase, J. M. 1999. To grow or to reproduce? The role of life-history plasticity in food web dynamics. - Am. Nat. 154: 571–586.
- Claessen, D. et al. 2002. The impact of size-dependent predation on population dynamics and individual life history. - Ecology 83: 1660–1675.
- Claessen, D. et al. 2004. Population dynamic theory of size-dependent cannibalism. - Proc. R. Soc. Lond. B Biol. Sci. 271: 333.
- Cohen, J. E. et al. 2003. Ecological community description using the food web, species abundance, and body size. - Proc. Natl. Acad. Sci. U. S. A. 100: 1781–1786.

- Conover, D. O. and Munch, S. B. 2002. Sustaining fisheries yields over evolutionary time scales. - *Science* 297: 94–96.
- Crow, J. F. 2008. Maintaining evolvability. - *J. Genet.* 87: 349–353.
- Darimont, C. T. et al. 2009. Human predators outpace other agents of trait change in the wild. - *Proc. Natl. Acad. Sci. U. S. A.* 106: 952–954.
- Day, T. et al. 2002. The role of size-specific predation in the evolution and diversification of prey life histories. - *Evolution* 56: 877–887.
- De Kort, H. et al. 2021. Life history, climate and biogeography interactively affect worldwide genetic diversity of plant and animal populations. - *Nat. Commun.* 12: 516.
- De Roos, A. M. et al. 2003. The influence of size-dependent life-history traits on the structure and dynamics of populations and communities. - *Ecol. Lett.* 6: 473–487.
- Devine, J. A. and Heino, M. 2011. Investigating the drivers of maturation dynamics in Barents Sea haddock (*Melanogrammus aeglefinus*). - *Fish. Res.* 110: 441–449.
- Dieckmann, U. and Law, R. 1996. The dynamical theory of coevolution: a derivation from stochastic ecological processes. - *J. Math. Biol.* 34: 579–612.
- Edeline, E. et al. 2007. Trait changes in a harvested population are driven by a dynamic tug-of-war between natural and harvest selection. - *Proc. Natl. Acad. Sci. U. S. A.* 104: 15799–15804.
- Edeline, E. et al. 2009. Harvest-induced disruptive selection increases variance in fitness-related traits. - *Proc. R. Soc. Lond. B Biol. Sci.* 276: 4163–4171.
- Edeline, E. et al. 2016. Empirical evidence for competition-driven semelparity in wild medaka. - *Popul. Ecol.* 58: 371–383.
- Edley, M. T. and Law, R. 1988. Evolution of life histories and yields in experimental populations of *Daphnia magna*. - *Biol. J. Linn. Soc.* 34: 309–326.
- Ellner, S. P. et al. 2011. Does rapid evolution matter? Measuring the rate of contemporary evolution and its impacts on ecological dynamics. - *Ecol. Lett.* 14: 603–614.
- Emmerson, M. C. and Raffaelli, D. 2004. Predator-prey body size, interaction strength and the stability of a real food web. - *J. Anim. Ecol.* 73: 399–409.
- Engen, S. et al. 2014. Evolutionary consequences of nonselective harvesting in density-dependent populations. - *Am. Nat.* 184: 714–726.
- Ernande, B. et al. 2004. Adaptive changes in harvested populations: plasticity and evolution of age and size at maturation. - *Proc. Biol. Sci.* 271: 415–423.

- Fausch, K. D. et al. 2021. Interspecific social dominance networks reveal mechanisms promoting coexistence in sympatric charr in Hokkaido, Japan. - *J. Anim. Ecol.* 90: 515–527.
- Fenberg, P. B. and Roy, K. 2008. Ecological and evolutionary consequences of size-selective harvesting: how much do we know? - *Mol. Ecol.* 17: 209–220.
- Fleming, I. A. and Gross, M. R. 1994. Breeding competition in a Pacific salmon (coho: *Oncorhynchus kisutch*): measures of natural and sexual selection. - *Evolution* 48: 637–657.
- Fox, L. R. 1975. Cannibalism in natural populations. - *Annu. Rev. Ecol. Syst.* 6: 87–106.
- Frankham, R. 1996. Relationship of genetic variation to population size in wildlife. - *Conserv. Biol.* 10: 1500–1508.
- Gadgil, M. and Bossert, W. H. 1970. Life historical consequences of natural selection. - *Am. Nat.* 104: 1–24.
- Gårdmark, A. and Dieckmann, U. 2006. Disparate maturation adaptations to size-dependent mortality. - *Proc. R. Soc. Lond. B Biol. Sci.* 273: 2185–2192.
- Gårdmark, A. et al. 2003. Life-history evolution in harvested populations: the role of natural predation. - *Evol. Ecol. Res.* 5: 239–257.
- Gomulkiewicz, R. and Holt, R. D. 1995. When does evolution by natural selection prevent extinction? - *Evolution* 49: 201–207.
- Govaert, L. 2018. Eco-evolutionary partitioning metrics: a practical guide for biologists. - *Belg. J. Zool.* 148: 167–202.
- Govaert, L. et al. 2019. Eco-evolutionary feedbacks—Theoretical models and perspectives. - *Funct. Ecol.* 33: 13–30.
- Grant, P. R. and Grant, B. R. 2002. Unpredictable evolution in a 30-year study of Darwin’s finches. - *Science* 296: 707–711.
- Griffiths, J. I. et al. 2020. Individual differences determine the strength of ecological interactions. - *Proc. Natl. Acad. Sci.* 117: 17068.
- Hadfield, J. D. 2008. Estimating evolutionary parameters when viability selection is operating. - *Proc. R. Soc. B Biol. Sci.* 275: 723–734.
- Hairston, N. G. et al. 2005. Rapid evolution and the convergence of ecological and evolutionary time. - *Ecol. Lett.* 8: 1114–1127.
- Hamrin, S. F. and Persson, L. 1986. Asymmetrical competition between age classes as a factor causing population oscillations in an obligate planktivorous fish species. - *Oikos* 47: 223–232.

- Hansen, Thomas F. et al. 2011. Heritability is not evolvability. - *Evol. Biol.* 38: 258–277.
- Hansen, T. F. et al. 2019. Predicting evolutionary potential: a numerical test of evolvability measures. - *Evolution* 73: 689–703.
- Haugen, T. O. et al. 2007. Density dependence and density independence in the demography and dispersal of pike over four decades. - *Ecol. Monogr.* 77: 483–502.
- Heino, M. et al. 2013. Can fisheries-induced evolution shift reference points for fisheries management? - *ICES J. Mar. Sci. J. Cons.* 70: 707–721.
- Heino, M. et al. 2015. Fisheries-induced evolution. - *Annu. Rev. Ecol. Evol. Syst.* 46: 461–480.
- Hendry, A. P. and Kinnison, M. T. 1999. The pace of modern life: measuring rates of contemporary microevolution. - *Evolution* 53: 1637–1653.
- Hendry, A. P. and Kinnison, M. T. 2001. An introduction to microevolution: rate, pattern, process. - *Genetica* 112: 1–8.
- Hilborn, R. and Walters, C. 1992. *Quantitative fisheries stock assessment: choice, dynamics and uncertainty.* - Springer US.
- Hilborn, R. and Minto-Vera, C. V. 2008. Fisheries-induced changes in growth rates in marine fisheries: are they significant? - *Bull. Mar. Sci.* 83: 95–105.
- Holliday, R. 1989. Food, reproduction and longevity: is the extended lifespan of calorie-restricted animals an evolutionary adaptation? - *BioEssays News Rev. Mol. Cell. Dev. Biol.* 10: 125–127.
- Holt, R. D. et al. 1994. Simple rules for interspecific dominance in systems with exploitative and apparent competition. - *Am. Nat.* 144: 741–771.
- Hunter, D. C. et al. 2018. Quantification and decomposition of environment-selection relationships. - *Evolution* 72: 851–866.
- Jusufovski, D. and Kuparinen, A. 2020. Exploring individual and population eco-evolutionary feedbacks under the coupled effects of fishing and predation. - *Fish. Res.* 231: 105713.
- Kingsolver, J. G. and Pfennig, D. W. 2004. Individual-level selection as a cause of Cope's rule of phyletic size increase. - *Evolution* 58: 1608–1612.
- Kinnison, M. T. et al. 2015. Cryptic eco-evolutionary dynamics. - *Ann. N. Y. Acad. Sci.* 1360: 120–144.
- Koch, M. et al. 2008. Pedigree reconstruction in wild cichlid fish populations. - *Mol. Ecol.* 17: 4500–4511.
- Kooijman, S. A. L. M. 2010. *Dynamic energy budget theory for metabolic organisation.* - Cambridge University Press.

- Kristiansen, T. S. and Svåsand, T. 1998. Effect of size-selective mortality on growth of coastal cod illustrated by tagging data and an individual-based growth and mortality model. - *J. Fish Biol.* 52: 688–705.
- Kuparinen, A. and Merilä, J. 2007. Detecting and managing fisheries-induced evolution. - *Trends Ecol. Evol.* 22: 652–659.
- Kuparinen, A. and Hutchings, J. A. 2017. Genetic architecture of age at maturity can generate divergent and disruptive harvest-induced evolution. - *Philos. Trans. R. Soc. B Biol. Sci.* 372: 20160035.
- Lande, R. and Arnold, S. J. 1983. The measurement of selection on correlated characters. - *Evolution* 37: 1210–1226.
- Law, R. 2000. Fishing, selection, and phenotypic evolution. - *ICES J. Mar. Sci. J. Cons.* 57: 659–668.
- Le Bourlot, V. et al. 2014. Interference versus exploitative competition in the regulation of size-structured populations (Mathew A. Leibold and Judith L. Bronstein, Eds.). - *Am. Nat.* 184: 609–623.
- Le Rouzic, A. et al. 2020. Unidirectional response to bidirectional selection on body size. II Quantitative genetics. - *Ecol. Evol.* 10: 11453–11466.
- Loeuille, N. and Loreau, M. 2005. Evolutionary emergence of size-structured food webs. - *Proc. Natl. Acad. Sci. U. S. A.* 102: 5761–5766.
- Loeuille, N. and Loreau, M. 2006. Evolution of body size in food webs: does the energetic equivalence rule hold? - *Ecol. Lett.* 9: 171–178.
- Loeuille, N. and Loreau, M. 2009. Emergence of complex food web structure in community evolution models. - In: Verhoef, H. A. and Morin, P. J. (eds), *Community ecology: processes, models, and applications*. 1st ed.n. Oxford University Press, pp. 163–178.
- Lynch, M. and Walsh, B. 2018. *Evolution and selection of quantitative traits*. - Oxford University Press.
- Marty, L. et al. 2014. Temporal trends in age and size at maturation of four North Sea gadid species: cod, haddock, whiting and Norway pout. - *Mar. Ecol. Prog. Ser.* 497: 179–197.
- Marty, L. et al. 2015. Fisheries-induced neutral and adaptive evolution in exploited fish populations and consequences for their adaptive potential. - *Evol. Appl.* 8: 47–63.
- McCann, K. et al. 1998. Weak trophic interactions and the balance of nature. - *Nature* 395: 794.
- Meszéna, G. et al. 2001. Evolutionary optimisation models and matrix games in the unified perspective of adaptive dynamics. - *Selection* 2: 193–220.
- Metz, J. A. J. et al. 2008. When does evolution optimize? - *Evol. Ecol. Res.* 10: 629–654.

- Morrissey, M. B. et al. 2010. The danger of applying the breeder's equation in observational studies of natural populations. - *J. Evol. Biol.* 23: 2277–2288.
- Olsen, E. M. and Moland, E. 2011. Fitness landscape of Atlantic cod shaped by harvest selection and natural selection. - *Evol. Ecol.* 25: 695–710.
- Orr, H. A. 2007. Absolute fitness, relative fitness, and utility. - *Evolution* 61: 2997–3000.
- Perälä, T. and Kuparinen, A. 2020. Eco-evolutionary dynamics driven by fishing: from single species models to dynamic evolution within complex food webs. - *Evol. Appl.* in press.
- Perez, K. O. and Munch, S. B. 2010. Extreme selection on size in the early lives of fish. - *Evolution* 64: 2450–2457.
- Persson, L. 1985. Asymmetrical competition: are larger animals competitively superior? - *Am. Nat.* 126: 261–266.
- Persson, L. and De Roos, A. M. 2006. Food-dependent individual growth and population dynamics in fishes. - *J. Fish Biol.* 69: 1–20.
- Persson, L. et al. 1998. Ontogenetic scaling of foraging rates and the dynamics of a size-structured consumer-resource model. - *Theor. Popul. Biol.* 54: 270–293.
- Peters, R. H. 1983. *The ecological implications of body size.* - Cambridge University Press.
- Post, J. R. et al. 1999. Density-dependent processes in structured fish populations: interaction strengths in whole-lake experiments. - *Ecol. Monogr.* 69: 155–175.
- Reed, D. H. and Frankham, R. 2001. How closely correlated are molecular and quantitative measures of genetic variation? A meta-analysis. - *Evolution* 55: 1095–1103.
- Renneville, C. et al. 2016. Morphological drivers of trophic cascades. - *Oikos* 125: 1193–1202.
- Renneville, C. et al. 2020. Unidirectional response to bidirectional selection on body size. I. Phenotypic, life-history, and endocrine responses. - *Ecol. Evol.* 10: 10571–10592.
- Roff, D. A. 1992. *The evolution of life histories.* - Chapman & Hall.
- Romiguier, J. et al. 2014. Comparative population genomics in animals uncovers the determinants of genetic diversity. - *Nature* 515: 261–263.
- Rooney, N. et al. 2006. Structural asymmetry and the stability of diverse food webs. - *Nature* 442: 265–269.
- Rudy, S. H. et al. 2017. Data-driven discovery of partial differential equations. - *Sci. Adv.* 3: e1602614.

- Schaefer, M. B. 1954. Some aspects of the dynamics of populations important to the management of commercial marine fisheries. - *Inter-Am. Trop. Tuna Comm. Bull.* 1: 23–56.
- Schaeffer, L. R. 2004. Application of random regression models in animal breeding. - *Livest. Prod. Sci.* 86: 35–45.
- Sgrò, C. M. and Partridge, L. 2000. Evolutionary responses of the life history of wild-caught *Drosophila melanogaster* to two standard methods of laboratory culture. - *Am. Nat.* 156: 341–353.
- Silva, A. et al. 2013. Long-term changes in maturation of sardine, *Sardina pilchardus*, in Portuguese waters. - *Sci. Mar.* 77: 429–438.
- Sinclair, A. R. E. et al. 2003. Patterns of predation in a diverse predator-prey system. - *Nature* 425: 288–290.
- Stige, L. C. et al. 2019. Density- and size-dependent mortality in fish early life stages. - *Fish Fish.* 20: 962–976.
- Stinchcombe, J. R. et al. 2014. Estimating uncertainty in multivariate responses to selection. - *Evolution* 68: 1188–1196.
- Stockwell, C. A. et al. 2003. Contemporary evolution meets conservation biology. - *Trends Ecol. Evol.* 18: 94–101.
- Strauss, S. Y. 2014. Ecological and evolutionary responses in complex communities: implications for invasions and eco-evolutionary feedbacks. - *Oikos* 123: 257–266.
- Swain, D. P. et al. 2007. Evolutionary response to size-selective mortality in an exploited fish population. - *Proc. R. Soc. B Biol. Sci.* 274: 1015.
- Taylor, B. E. and Gabriel, W. 1992. To grow or not to grow: optimal resource allocation for *Daphnia*. - *Am. Nat.* 139: 248–266.
- Tilman, D. 1982. Resource competition and community structure. - Princeton University Press.
- Trippel, E. A. 1995. Age at maturity as a stress indicator in fisheries. - *BioScience* 45: 759–771.
- Tromeur, E. and Loeuille, N. 2017. Balancing yield with resilience and conservation objectives in harvested predator–prey communities. - *Oikos* 126: 1780–1789.
- van Benthem, K. J. et al. 2017. Disentangling evolutionary, plastic and demographic processes underlying trait dynamics: a review of four frameworks. - *Methods Ecol. Evol.* 8: 75–85.
- van de Wolfshaar, K. E. et al. 2008. Population feedback after successful invasion leads to ecological suicide in seasonal environments. - *Ecology* 89: 259–268.

- Verhulst, P. F. 1838. Notice sur la loi que la population suit dans son accroissement. - Corresp. Mathématique Phys. 10: 113–121.
- Walsh, M. R. et al. 2006. Maladaptive changes in multiple traits caused by fishing: impediments to population recovery. - Ecol. Lett. 9: 142–148.
- Weinbach, A. et al. 2017. Selective effects of temperature on body mass depend on trophic interactions and network position. - bioRxiv: 233742.
- Wellborn, G. A. 1994. Size-biased predation and prey life histories: a comparative study of freshwater amphipod populations. - Ecology 75: 2104–2117.
- White, C. et al. 2012. The value of coordinated management of interacting ecosystem services. - Ecol. Lett. 15: 509–519.
- Wood, Z. T. et al. 2018. Eco-evolutionary feedbacks from non-target species influence harvest yield and sustainability. - Sci. Rep. 8: 6389.
- Woodward, G. et al. 2005. Body size in ecological networks. - Trends Ecol. Evol. 20: 402–409.
- Yacine, Y. et al. 2021. Collapse and rescue of evolutionary food webs under global warming. - J. Anim. Ecol. in press.