Interim Report IR-13-032

The evolution of age-dependent plasticity

Barbara Fischer
G. Sander van Doorn
Ulf Dieckmann (dieckmann@iiasa.ac.at)

Approved by
Pavel Kabat
Director General and Chief Executive Officer

June 2015
The evolution of age-dependent plasticity

Barbara Fischer\textsuperscript{1,2,3,4,*}, G. Sander van Doorn\textsuperscript{1,5,*}, Ulf Dieckmann\textsuperscript{2} and Barbara Taborsky\textsuperscript{1,2}

1. Division of Behavioural Ecology, Institute of Ecology and Evolution, University of Bern, Switzerland
2. Evolution and Ecology Program, International Institute for Applied Systems Analysis, Laxenburg, Austria
3. Centre for Ecological and Evolutionary Synthesis, Department of Biology, University of Oslo, Norway
4. Department of Theoretical Biology, University of Vienna, Austria
5. Centre for Ecological and Evolutionary Studies, University of Groningen, the Netherlands

* The first two authors contributed equally to this article

Author E-mail addresses

Barbara Fischer barbara.fischer@ibv.uio.no
Sander van Doorn (corresponding author) g.s.van.doorn@rug.nl
Ulf Dieckmann dieckmann@iiasa.ac.at
Barbara Taborsky barbara.taborsky@iee.unibe.ch

Keywords

Developmental plasticity
Plasticity windows
Reaction norms
Eco-evo-devo
Information sampling
Dynamic optimization

Elements of the manuscript that will appear in the expanded online edition

Online Appendix A with supporting material, supplementary table A1 and supplementary figures A1 and A2

Submitted as an Article
Abstract

When organisms encounter environments that are heterogeneous in time, phenotypic plasticity is often favored by selection. The degree of such plasticity can vary during an organism’s lifetime, but the factors promoting differential plastic responses at different ages or life stages remain poorly understood. Here we develop and analyze an evolutionary model to investigate how environmental information is optimally collected and translated into phenotypic adjustments at different ages. We demonstrate that plasticity must often be expected to vary with age in a non-monotonic fashion. Early in life it is generally optimal to delay phenotypic adjustments until sufficient information has been collected about the state of the environment to warrant a costly phenotypic adjustment. Towards the end of life, phenotypic adjustments are disfavored as well, because their beneficial effects can no longer fully be reaped before death. Our analysis clarifies how patterns of age-dependent plasticity are shaped by the interplay of environmental uncertainty, the accuracy of perceived information and the costs of phenotypic adjustments with life-history determinants such as the relative strengths of fecundity and viability selection experienced by the organism over its lifetime. We conclude by comparing our results with expectations for alternative mechanisms, including developmental constraints, that promote age-dependent plasticity.
Introduction

Phenotypic plasticity is a universal property of living organisms (Tollrian and Harvell 1999, West-Eberhard 2003). Plasticity reveals itself as the capacity of a single genotype to produce different phenotypes in response to environmental influences during development. The adaptive use of information about environmental conditions distinguishes phenotypic plasticity from stochastic switching or bet-hedging (Slatkin, 1974), which is a risk-spreading strategy, frequently employed by microbes (Veening et al. 2008), that helps to ensure long-term survival in an unpredictably varying environment.

A plastic genotype has a selective advantage over a non-plastic one if the former has a higher net fitness than the latter averaged over the environments the organism can encounter (Bradshaw 1965, Levins 1968). Theoretical studies suggest that plastic genotypes are superior in variable environments when sufficiently reliable environmental cues are available and costs of plasticity are low (Via and Lande 1985, Van Tienderen 1991, Gomulkiewicz and Kirkpatrick 1992, Schlichting and Pigliucci 1995, Houston and McNamara 1992, Ernande and Dieckmann 2004).

The plastic adjustment of phenotypes can involve morphological modifications, adaptations of physiological and neural regulation, or behavioural changes. A well-known example of morphological reconstruction is found in Daphnia sp., with individuals adapting to environmental conditions by growing a protective helmet-like structure in response to the presence of predators (Tollrian 1990). Physiological plasticity is observed, for instance, in several closely related species of larks (family Alaudidae), which can adjust their basal metabolic rate to the ambient temperature (Tieleman et al. 2003). In the rat (and several other mammals) the level of maternal care (pup licking / grooming) received early in life has long-lasting effects on the responsiveness to stress, mediated by brain-specific DNA methylation and differential expression of stress hormone receptors in the central nervous system (Szyf et al. 2007). Finally, an example of behavioural plasticity is found in the spider Parawixia bistriata, which adjusts the size and structure of its web to the type of prey it expects to catch (Sandoval 1994).
If organisms were able to acquire full information about which phenotype is optimal in a given situation, and if adjustments would be cost-free and could be realized without time lags, we would expect to see organisms with unlimited plasticity. Such hypothetical organisms (sometimes called ‘Darwinian demons’ after Law 1979) would express highly specialized phenotypes, and constantly switch between them as their environments change, so as to express optimal trait values for every possible environmental situation. This clearly is not what we see in nature. One reason for this is that plasticity generally comes at a cost. Morphological adjustments are likely to be associated with high construction costs and may be difficult to reverse (Brönmark and Miner 1992, Van Buskirk 2000, Callahan et al. 2008), whereas physiological and behavioural plasticity is usually mediated by a redirection of neuroendocrine and hormonal regulatory pathways. The latter is often considered to be less costly than morphological reconstruction, but can be associated nevertheless with a number of costly (e.g., mobilization of energy and tissue nitrogen) and potentially risky (e.g., down-regulation of the immune system) physiological processes (reviewed in Sapolsky et al. 2000, Sapolsky 2002, Badre and Wagner 2006).

Limits to plasticity are also illustrated by the observation that many organisms are more responsive to environmental perturbations during some ages or life stages than during others (e.g., Dufty et al. 2002, Hoverman and Relyea 2007). These patterns are observed to vary across species (Hoverman and Relyea 2007) and traits (e.g., Taborsky 2006, Kotrschal and Taborsky 2010, Arnold and Taborsky 2010, Segers and Taborsky 2012). For instance, bryozoans can grow defensive structures in response to chemical predator cues only early in their life (Harvell 1991), and in rats persistent stress resistance can be induced by maternal care only if experienced in the first week after birth (Szyf et al. 2007). In freshwater snails (Helisoma trivolvis), the ability to build defensive structures against predatory water bugs extends well beyond sexual maturity, whereas a reversal of this trait is only possible during early ontogeny (Hoverman and Relyea 2007). Finally, as an example of a species exhibiting a prolonged high degree of plasticity in a morphological trait, we mention barnacles (Balanus glandula), which maintain a lifelong ability to grow and shrink legs used for suspension feeding in response to flow conditions (Marchinko 2003).
It is not yet understood which factors determine the diverse patterns of age-dependent plasticity across species and traits that are observed in nature. In general, changes in plasticity with age are expected if an organism does not have perfect information at birth, but can improve its estimate of the environmental state by integrating information accumulated over a longer period of time (Dufty et al. 2002). Some theoretical work exists on the evolution of reversible plastic responses (Gabriel et al. 1999, 2005), but, to our knowledge, the evolution of age-dependent phenotypic plasticity has not so far been systematically explored.

Here we study how plasticity is expected to change with age in an environment that varies stochastically over time. To this end, we calculate optimal patterns of age-dependent plasticity and examine how these depend on the rates of environmental fluctuations, the organism’s life history, and the relative strengths of selection on different components of fitness. We model the process of information acquisition, which is crucial for decision-making in uncertain environments (Real 1992, Dall et al. 2005), and consider different degrees of perception accuracy and plasticity costs.

**Model**

The definition of our model will be structured as follows: first, we focus on the environment, which we assume to be both stochastically fluctuating and partially predictable. We then describe how organisms can predict future conditions based on current and past observations of the state of the environment. Next, we explain how organisms adjust their phenotypes depending on the gathered information, given a reaction norm for age-dependent plasticity. As a final step, we specify how the fitness of a reaction norm is calculated and outline the optimization procedure for finding a reaction norm that maximizes fitness. Figure 1 provides a preview of how these steps coincide with life-cycle events in our model and also serves as a reference for some of the notation that will be developed.

*Fluctuations in the state of the environment (Figure 1, step 1)*

We consider a population of organisms living in a variable environment that changes stochastically from one reproductive season to the next. At each such time step, the environment can be in one of two discrete states, denoted A and B, representing different ecological
conditions, such as high-flow and low-flow conditions in an aquatic environment. It should be understood that these two conditions in general do not need to represent a ‘good’ and a ‘poor’ environmental state, even though this latter distinction is common and important. In fact, we are primarily interested in situations in which the two different ecological conditions call for different phenotypic specializations, such that the fitness rank of phenotypes may change when the environment switches from one state to the other.

The lifetime reproductive success of an individual depends on the sequence of environmental states it experiences during its life. We denote this sequence by $E = (E_1, E_2, \ldots, E_T)$, where $E_t = A$ or $B$ represents the state of the environment at time $t$, and $T$ is the maximum lifetime of individuals. For each individual, time is measured relative to the moment of its birth and expressed in discrete time units corresponding to one reproductive season. We assume that the state of the environment at time $t$ is dependent on its state at time $t - 1$, such that the $E_t$ are correlated random variables. Accordingly, we model the environmental fluctuations as a first-order autoregressive stochastic process with two parameters $\alpha$ and $\beta$ that define the rates of switching between environmental states. Specifically, $\alpha$ is the probability at each time step that the environment switches from state $B$ to $A$, which can be expressed as the conditional probability $P[E_t = A \mid E_{t-1} = B]$. Likewise, $\beta$ is the reverse transition probability, i.e., $\beta = P[E_t = B \mid E_{t-1} = A]$. Throughout, we focus on environments for which $0 < \alpha + \beta < 1$. Under this condition, $E_t$ and $E_{t+1}$ are positively correlated, such that, given knowledge of the current state of the environment ($E_t$), the organism can predict the future state $E_{t+1}$ and adjust its phenotype accordingly. The accuracy of this prediction is inherently limited, however, by the fact that $E_t$ and $E_{t+1}$ cannot be perfectly correlated in a changing (i.e., $\alpha + \beta > 0$) environment.

Environmental sampling and the integration of information (Figure 1, steps 2 and 3)

A second factor that limits an organism’s ability to predict future conditions is that the state of the environment may not be directly observable, forcing individuals to infer information from a finite sample of imperfect cues. In our model, we therefore introduce the random variable $O_t$ ($O_t = A$ or $B$) to represent the observation of the state of the environment made by an individual at time $t$. The observed and actual environmental state may be strongly or weakly correlated to each other, depending on the reliability of the information that is accessible to the organism.
Specifically, we assume that, irrespective of the state of the environment, observations are correct with probability $a$, such that $P[O_t = o_t \mid E_t = e_t]$ is equal to either $a$ or $1 - a$, depending on whether the current state of the environment is perceived correctly ($o_t = e_t$) or not ($o_t \neq e_t$). Here and henceforth, $o_t$ ($o_t = A$ or $B$) is used to denote the actual observation at time $t$ made by a particular individual under consideration (i.e., $o_t$ is a realization of $O_t$). A similar consistent use of upper- and lowercase symbols distinguishes between the state of the environment as a random variable ($E_t$) and its realization ($e_t$; see also Online Appendix A, notational conventions). Throughout, we will refer to the parameter $a$ as the sampling accuracy.

Even though a single observation has limited accuracy, older individuals who have repeatedly sampled the environment, may still be able to estimate the state of the environment reliably by integrating information over the sequence of observations $o_t = (o_1, o_2, \ldots, o_t)$ they have made up to their present age. However, earlier observations are inherently less informative than more recent ones, because the environment may have changed in the time since an observation was made. As a result, the organism needs to find a balance between rapidly discounting previous information, so as to minimize the risk of making decisions based on out-of-date observations (adaptive forgetting; Kraemer and Golding 1997), and integrating over a large number of observations so as to avoid being misled by observation errors. An optimal solution for this problem is to use Bayesian updating after each observation, in order to estimate how likely it is that the environment is currently in one state or the other.

Let us therefore assume that the organism is capable of keeping track of a state variable $p_t$ that reflects its best possible estimate for the current state of the environment given the limited information it has access to. As this information is fully contained in the sequence of observations, we define an individual’s estimate $p_t$ as a likelihood

$$p_t = P[E_t = A | O_t = o_t \cap O_{t-1} = o_{t-1} \cap \cdots \cap O_1 = o_1]$$

(1)

that is conditioned on the complete history of observations made by the individual up to its present age.

In Online Appendix A, we derive how each individual can calculate its estimate $p_t$ based on its prior knowledge of the state of the environment (represented by the previous estimate $p_{t-1}$), and its current observation, $o_t$. This dependence can be expressed in the form of a Bayesian update
rule $U$, which maps the previous estimate $p_{t-1}$ to a new, updated estimate $p_t$ after making observation $O_t = o_t$:

$$
p_t = U(p_{t-1}, o_t) = \begin{cases} 
\frac{a[(1 - \beta)p_{t-1} + \alpha(1 - p_{t-1})]}{a + (1 - 2a)[\beta p_{t-1} + (1 - \alpha)(1 - p_{t-1})]} & \text{if } o_t = A, \\
1 - \frac{a[\beta p_{t-1} + (1 - \alpha)(1 - p_{t-1})]}{a + (1 - 2a)[1 - \beta]p_{t-1} + \alpha(1 - p_{t-1})} & \text{if } o_t = B.
\end{cases} \tag{2}
$$

The derivation of this result, which follows from Bayes’ theorem and the laws of probability for conditionally independent events (Online Appendix A), rests on the assumption that the environmental switching rates and the sampling accuracy are ‘known’ in the sense that the considered species has previously adapted to the considered fluctuating environment. As an implication, $p_0$, the initial estimate of a naïve individual who has not yet made any observations, is taken to be equal to the long-term average frequency of environmental state $A$, $P[E_t = A] = \alpha / (\alpha + \beta)$.

Equation (2) conforms to the biological intuition in two ways: first, it confirms that prior information is less valuable in a more variable and less predictable environment. Specifically, in the absence of environmental autocorrelation ($\alpha = 1 - \beta$), knowledge of the previous state of the environment becomes useless for predicting the current state. Therefore, the right-hand side of equation (2) becomes independent of $p_{t-1}$ if $\alpha = 1 - \beta$. Second, it indicates that the value of current information decreases with the frequency of observation errors in an individual’s assessment of the environmental state. In the event that observations are as likely to be correct as not ($a = \frac{1}{2}$), the right-hand side of equation (2) becomes independent of $o_t$. In that case, no information can be accumulated and $p_t$ remains at $p_0 = P[E_t = A] = \alpha / (\alpha + \beta)$.

The typical situation considered in our analysis is when the result of the Bayesian update rule depends both on the current, potentially erroneous observation and on information collected earlier. As an example of such a case, consider an organism in a fluctuating environment with $\alpha = 0.15$ and $\beta = 0.1$. With these switching rates, the long term average frequency of environmental state $A$ is $\alpha / (\alpha + \beta) = 0.6$, such that a naïve organism does best by starting with an initial estimate $p_0 = 0.6$. Suppose that, at age 1, the organism observes that the environment is in state B. Based on equation (2), it will then decrease its estimate $p_1$ to a value less than $p_0$,.
but larger than zero, because generally the organism cannot be certain that the environment truly is in state B based on this single observation. For instance, if the sampling accuracy is $a = 0.7$, we find $p_1 = 0.39$ (after observing B in this particular environment). Subsequent observations of environmental state B at age 2 and 3 would further increase the organism’s confidence that the environment is in state B (application of the Bayesian update rule gives $p_2 = 0.25$ and $p_3 = 0.18$). However, if the organism observes environmental state A at age 4, 5 and 6, the estimates go up again (in this case, equation (2) gives: $p_4 = 0.48$, $p_5 = 0.71$ and $p_6 = 0.83$).

The range of values that the estimate $p_t$ can take is constrained by the inequalities

$$\alpha (1 - a) / (2a - 1) < p_t < 1 - \beta (1 - a) / (2a - 1)$$

(this lower and upper bound is found by solving $p_t = U(p_t, B)$ and $p_t = U(p_t, A)$ for small $\alpha$ and $\beta$). Certainty about the state of the environment is therefore inherently limited by both the environmental switching rates and the accuracy of individual observations. As a result, there is also a limit to an organism’s knowledge gain through sampling.

Development of the phenotype (Figure 1, step 4)

After the individual has sampled the environment, and has integrated the newly obtained information with previous observations, it may adjust its phenotype. We allow the level of adjustment to depend on the organism’s state, which encompasses its age, its phenotype at the previous time step, and its estimate of the state of the environment. For simplicity, we take the phenotype to be a one-dimensional trait that can take any value between 0 and 1, and describe its development by a recursion

$$x_t = x_{t-1} + h_t(x_{t-1}, p_t).$$

(3)

Here $x_t$ denotes the phenotype at age $t$, and $h_t$ is the reaction norm that captures how the organism adjusts its phenotype depending on its state after sampling the current environment. As for the estimate $p_0$, we assume that the initial phenotype $x_0$ has been set by adaptive evolution. Our further analysis therefore treats $x_0$ as an evolutionary trait that is optimized together with the reaction norm.

Given an initial phenotype $x_0$ and a reaction norm $h$, the recurrence relationship (3) and update rule (2) allow us to calculate an individual’s developmental trajectory $x_0 \rightarrow x_1 \rightarrow \ldots \rightarrow x_T$ from
the sequence of observations the individual makes throughout its life (figure 1). In the next section, we explain how the developmental trajectory determines an individual’s lifetime reproductive success. As a final step, we outline the procedure for maximizing the expectation of this fitness measure over environmental states in order to find the optimal reaction norm.

**Fitness consequences of plasticity (Figure 1, step 5)**

The fitness of a reaction norm $h$ depends on its average performance across all possible realizations of the sequence of environmental states. Moreover, in any given environment, not all individuals will make the same sequence of observations due to errors in the assessment of environmental cues. As these errors can induce a change in the phenotypic trajectory, they represent an additional source of variation for the fitness of the reaction norm. Accordingly, the fitness function $W$, which has to be maximized to identify the optimal reaction norm, is defined by a double average

$$W = \prod_e \left( \sum_o P(O = o | E = e) R_1(o, e) \right)^{P[E = e]}.$$  

(4)

Here, $R_1(o, e)$ denotes the lifetime reproductive success (from age 1 onwards) of an individual with observation sequence $o = (o_1, o_2, \ldots, o_T)$ in environment $e = (e_1, e_2, \ldots, e_T)$. The summation averages individual lifetime reproductive success over the distribution of observation sequences in environment $e$, yielding the population-average fitness of the reaction norm in that environment. The product averages the population’s fitness over all possible realizations of the environment $e$, using the standard geometric mean fitness criterion for evaluating the long-term evolutionary success of a strategy in a stochastic environment (Lewontin and Cohen 1969).

All that remains to complete the definition of the model, is to specify a procedure for determining $R_1(o, e)$. One straightforward but indirect method is to calculate the expected reproductive success of an individual at age $t$ and onwards from the recursion $R_t(o, e) = S_t \left( F_t + R_{t+1}(o, e) \right)$. Here, $S_t$ denotes the survival probability of the individual at age $t$ and $F_t$ denotes its fecundity at that age. Iterating the recursion backwards in time from $t = T$ to $t = 1$ (with the terminal reward $R_{T+1}(o, e)$ defined to be zero), gives an expression for the lifetime reproductive success $R_1(o, e)$.
From here on, fecundity and survival probability will be written as functions \( F_t(x, \Delta x_t) \) and \( S_t(x, \Delta x_t) \), respectively, to emphasize that these fitness components depend on the current environment \( e_t \) (\( e_t = A \) or \( B \)), the current phenotype \( x_t \), and the phenotypic adjustment \( \Delta x_t = |x_t - x_{t-1}| \) made by the individual at age \( t \). The dependence on \( e_t \) and \( x_t \) is critical for modeling the benefits of plasticity (i.e., expressing a phenotype that matches with the environment), while the dependence on \( \Delta x_t \) is included to capture potential costs associated with the process of phenotypic adjustment. Our analysis excludes cases where an organism’s current phenotype determines survival or fecundity later in life, as, for example, when the organism stores energy reserves for later use in reproduction. Such more complex scenarios can be analyzed by introducing additional state variables, which we choose to avoid here.

**Linearization of the fitness function and evolutionary optimization of the reaction norm**

For any given fecundity and survival function, equation (4) can be maximized using evolutionary optimization methods (e.g., individual-based simulation). However, this approach provides limited biological insight. We therefore make a number of simplifying assumptions, which enable us to obtain approximate expressions for the fitness function that clarify how the cost and benefit of plasticity interact with the life-history of the organism. Here, we only give a brief outline of this derivation; technical details are provided in Online Appendix A. The main simplification is that we assume selection to be weak. This allows us to ignore, up to first approximation, interaction effects between components of selection associated with different environmental states or acting on different life-history stages. In addition, we take the costs of phenotypic adjustment to be independent of the state of the environment, and first assume that \( F_t \) and \( S_t \) are linear in their arguments \( x_t \) and \( \Delta x_t \), before generalizing our results to arbitrary nonlinear functions in Appendix A (see also online figure A1).

The first step in simplifying the fitness function is to consider an individual with a fixed phenotype \( x_t = z \) and to use the average life history of this individual as a benchmark against which all fitness effects of plasticity are measured. If selection is weak, all fitness deviations from the reference life history are small, which implies that the environmental fluctuations have modest effects on survival and fecundity. With this in mind, we introduce two sets of (small) selection coefficients. First, the coefficients \( f_t^s = \frac{\partial}{\partial x} F_t(x, 0) / \overline{F}_t \) and \( s_t^s = \frac{\partial}{\partial x} S_t(x, 0) / \overline{S}_t \)
quantify the relative difference in, respectively, fecundity and survival between two individuals whose phenotypes differ by one phenotypic unit. Positive values of these coefficients indicate that selection favors higher values of $x_i$ in environment $e_i$. Second, the coefficients $f'_t = -\frac{\partial}{\partial y} \bar{F}_t(z, y) / \bar{F}_t$ and $s'_t = -\frac{\partial}{\partial y} \bar{S}_t(z, y) / \bar{S}_t$ measure the relative marginal fecundity and survival costs of phenotypic adjustment at age $t$ per unit of phenotype change. Larger positive values of $f'_t$ and $s'_t$ reflect stronger fecundity and viability selection against phenotypic adjustment. Throughout, the use of an overbar, as in $\bar{F}_t(z, y)$ and $\bar{S}_t(z, y)$, will signify an average across environmental states (e.g., $\bar{F}_t(z, y) = \sum_{e=A,B} P[E_t = e] F'_t(z, y)$). The selection coefficients $f'_t$ and $s'_t$ depend only on these averages as a result of our assumption that the marginal costs of phenotypic adjustment do not differ between environmental state A and B.

If selection is weak, the difference in reproductive success between the life history of an individual with reaction norm $h$ and the reference life history can be approximated by a linear function in the selection coefficients. In order to minimize the approximation errors in this step of the analysis, we choose the reference phenotype $z$ equal to the value that maximizes lifetime reproductive success for an individual with a fixed phenotype. Using once more the recursive definition of expected future reproductive success ($R_t = S_t (F_t + R_{t+1})$), the relative fitness advantage of a phenotypically plastic individual can now be expressed in terms of its additional reproductive success from age $t$ onwards, $\partial R_t$, relative to an individual with the fixed phenotype $z$.

The fitness measure $\partial R_t$ is a function of the state of the individual at age $t$, after it has observed the state of the environment and updated its estimate to $p_t$, but before it has adjusted its phenotype (indicated by the block arrow in figure 1). Based on the derivation in Online Appendix A, $\partial R_t$ is defined by a sum of three terms that correspond to three subsequent steps in the cycle of events that occur in each breeding season:

$$\partial R_t(x_{t-1}, p_t) = -h_t(x_{t-1}, p_t) \left( s'_t + \frac{\bar{F}_t}{\bar{R}_t} f'_t \right)$$

$$+ \left( x_{t-1} + h_t(x_{t-1}, p_t) - z \right) \sum_{e \in \{A,B\}} P[E_t = e | O_t = o_t] \left( s'_t + \frac{\bar{F}_t}{\bar{R}_t} f'_t \right)$$

$$+ \left( 1 - \frac{\bar{F}_t}{\bar{R}_t} \right) \sum_{o \in \{A,B\}} P[O_{t+1} = o | O_t = o_t] \partial R_{t+1}(x_{t-1} + h_t(x_{t-1}, p_t), U(p_t, o)).$$
First, the organism changes its phenotype from the old value $x_{t-1}$ to the new value $x_t = x_{t-1} + h_t(x_{t-1}, p_t)$, at which point it has to pay the cost of plasticity. The resulting fitness reduction, captured by the first term on the right-hand side above, is proportional to the amount of phenotypic adjustment and increases with the marginal fecundity and survival costs of plasticity at age $t$, $f'_t$ and $s'_t$. These two costs are weighted according to their relative impact on the remaining lifetime reproductive success: reduced fecundity only affects the expected reproductive output in the current season (its relative contribution to the remaining reproductive success is given by $\frac{F_t S_t}{R_t}$), whereas reduced survival impacts all further reproductive success from age $t$ onwards (a similar differential weighting applies to the coefficients $f'_t$ and $s'_t$ discussed in the following paragraph).

After phenotypic adjustment, the organism is first subject to viability selection and then to fecundity selection. Accordingly, the second line on the right-hand side of equation (5) measures the fitness effect of expressing the new phenotype $x_t$ (written as $x_{t-1} + h_t(x_{t-1}, p_t)$), relative to the fitness of the reference individual with phenotype $z$. The magnitude of this contribution to $\partial R_t$ depends on the difference between $x_t$ and $z$, as well as on the fitness gradient (given by the term $s'_t + f'_t \frac{F_t S_t}{R_t}$) averaged over the distribution of environmental states across the individuals with observation history $O_t = o_t$ (here and elsewhere, $O_t = o_t$ stands for the composite event $O_t = o_t \cap O_{t-1} = o_{t-1} \cap \ldots \cap O_1 = o_1$). By definition (1), the distribution of environmental states for such individuals is given by $P[E_t = A \mid O_t = o_t] = p_t$ and $P[E_t = B \mid O_t = o_t] = 1 - p_t$, which captures the critical connection between an individual’s estimate of the state of the environment and the selective conditions that it is likely to experience.

The final step in each cycle of events is the transition from the current breeding season to the next, which is associated with a potential change in the state of the environment, a new observation $O_{t+1} = o$, and an update of the estimate $p_t$ to $p_{t+1} = U(p_t, o)$. The last line on the right-hand side of equation (5) takes into account that individuals can be in two different states after these events, depending on their observation at age $t+1$. The contribution of each of the corresponding future life-history trajectories to the remaining lifetime reproductive success is weighted by its probability of occurring, and the entire sum is multiplied by the relative
contribution of future fitness to the current remaining reproductive success, \( 1 - \frac{F_t S_t}{R_t} \). According to equation (A18) in Appendix A, the probabilities \( P[O_{t+1} = A | O_t = o_t] \) and \( P[O_{t+1} = B | O_t = o_t] \) can again be expressed in terms of \( p_t \).

The final step in the linearization procedure is to approximate equation (4) for the long-term average fitness of the reaction norm, using the fact that all \( \partial R_i \) are small. Under this approximation, the optimization task reduces to maximizing the relative difference \( \partial W \) in expected lifetime reproductive success between a plastic individual and an individual with the optimal fixed phenotype \( z \), where \( \partial W \) is given by

\[
\partial W = \frac{W - R_i}{R_i} \approx \sum_{o \in \{A, B\}} P[O_t = o] \partial R_i (x_o, U(p_o, o)).
\]  

As indicated by equation (5), the maximization of \( \partial W \) requires the optimization of a sequence of interdependent functions \( \partial R_i \). Since the dependency between these functions is unidirectional according to equation (5), the optimal reaction norm \( h \) can be found by backward state-dependent optimization. That is, we first maximize \( \partial R_{T_r} \), then \( \partial R_{T_{r-1}} \), and so on, until \( \partial R_i \) has been maximized. The final step of the optimization is to find the optimal initial phenotype \( x_0 \). An annotated version of the C++ code used for the optimization has been deposited on http://www.datadryad.org (doi:10.5061/dryad.kh008).

**Results**

**Optimal reaction norms for a semelparous life history**

In order to calculate the optimal reaction norm \( h \) it is necessary to specify the life history of the organism, as determined by the fecundity and survival probability functions \( F_i^\alpha(x_i, \Delta x_i) \) and \( S_i^\alpha(x_i, \Delta x_i) \). A simple case, which we will consider first, is when the species is semelparous, meaning that individuals reproduce once after reaching maturation at age \( T \) and die afterwards. Lifetime reproductive success then depends on the cumulative survival up to the reproductive event and the organism’s fecundity. We assume that only survival is affected by the phenotype in each respective environment, and take

\[
S_i^\alpha(x_i, \Delta x_i) = \begin{cases} 
1 - s \left(1 - x_i\right) - c \Delta x_i & \text{if } e_i = A, \\
1 - s x_i - c \Delta x_i & \text{if } e_i = B.
\end{cases}
\]  

(7)
Accordingly, at all ages, the optimal phenotype in environment A is $x_t = 1$, whereas $x_t = 0$ is optimal in environment B. The parameter $s$ ($0 < s << 1$), determines the survival disadvantage of maladapted phenotypes and, therefore, measures the strength of selection. In addition, survival at each time step decreases with the current amount of phenotypic adjustment. Parameter $c$ ($0 < c << 1$) measures the cost of plasticity, which we assume to be independent of the state of the environment. For the fecundity function, we take $F_t^{x_t}(x_t, \Delta x_t; t) = 0$ for all $0 < t < T$. The fecundity at age $T$, $F_T^{x_T}(x_T, \Delta x_T ;T) = \varphi$, is independent of $e_T, x_T$ and $\Delta x_T$, with $\varphi > 1$ set by density-dependence, such that the population remains stationary.

With these definitions, the recursion for the expected net fitness effect of plasticity, $\partial R_t$ (equation (5)), simplifies to

$$
\partial R_t(x_{t-1}, p_t) = -c \left[ h_t(x_{t-1}, p_t) \right] + s(2p_t - 1)(x_{t-1} + h_t(x_{t-1}, p_t) - z) + \sum_{o \in \{A,B\}} P[O_{t+1} = o | O_t = o_t] \partial R_{t+1}(x_{t-1} + h_t(x_{t-1}, p_t), U(p_t, o)).
$$

(8)

This expression is accurate up to first order in $s$ and $c$ (Online Appendix A). The first line on the right-hand side quantifies the current cost and benefit of phenotypic adjustment whereas the terms on the second line take into account its future fitness consequences. If the organism has no or little information about the state of the environment ($p_t \approx 0.5$), current survival is maximized if no phenotypic adjustment occurs (i.e., the cost term is minimized by $h_t = 0$). However, when the absolute value of $s(2p_t - 1)$ exceeds $c$, it becomes beneficial to adjust the phenotype to either $x_t = 1$ or $x_t = 0$, depending on what the current state of the environment is estimated to be.

As explained in the previous section, the estimate $p_t$ changes in response to the sequence of observations made by the individual, according to the Bayesian update rule (2). Consider, for example, an individual with maturation age $T = 6$, who makes the observations $o = (B, B, B, A, A, A)$ during its life. In an environment with switching rates $\alpha = 0.15$ and $\beta = 0.1$ and sampling accuracy $a = 0.7$ (the parameters used earlier for illustrating equation (2)), the estimate of the focal individual changes from $p_0 = 0.6$ to $p_1 = 0.39$, $p_2 = 0.25$, $p_3 = 0.18$, $p_4 = 0.48$, $p_5 = 0.71$ and $p_6 = 0.83$ (this sequence is indicated by grey lines and circles in the left part of figure 2a).
In accordance with the preceding discussion of equation (8), we find that individuals with the optimal reaction norm (found for $s = 0.05$ and $c = 0.02$ by backward state-dependent optimization) switch between $x_t = 1$ and $x_t = 0$ only when they are sufficiently confident that their current phenotype is suboptimal under the present environmental conditions. For the example individual with observation sequence $o = (B, B, A, A, A)$ this means that the phenotype switches from $x_1 = 1$ to $x_2 = 0$ after the individual observes for the second time that the environment is in state B (when its estimate is $p_2 = 0.25$). At a later stage, the phenotype switches back again from $x_4 = 0$ to $x_5 = 1$ after state A has been observed twice, first at age 4 and then at age 5 (the estimate is then $p_5 = 0.71$). In both cases, the switching points are correctly predicted by the condition $s |2 p_t - 1| > c$ (but see the discussion on the time-dependency of the reaction norm below). The phenotype trajectory $x_0 \rightarrow x_1 \rightarrow \ldots \rightarrow x_6$ for the example individual is highlighted in figure 2b (left part; grey lines and circles).

So far, we have focused on a single observation sequence. With $T = 6$, there are $2^6 = 64$ possible sequences of observations, which collectively give rise to a bifurcating tree of estimate and phenotype trajectories (shown in black in the left column of figure 2). Which path through the tree an individual will take is determined by its sequence of observations: each branch in the tree of estimates (figure 2a) splits into two new branches at the next observation event, from where the individual will follow the right path if it observed A, or the left path if it observed B. Accordingly, the rightmost and leftmost path in the tree correspond to the observation sequences (A, A, A, A, A, A) and (B, B, B, B, B, B), respectively. The phenotype tree (figure 2b) does not necessarily split after each observation because the optimal reaction norm induces a phenotypic switch only when the individual is sufficiently confident that its current phenotype is suboptimal.

In general, not all observation sequences have the same probability of occurrence. First, if the environment is strongly auto-correlated and the sampling accuracy is high, sequences with no or very few switches like (A, A, A, A, A, A), will be much more likely to occur than sequences with many switches like (A, B, A, B, A, B). This effect is visible to some extent in figure 2a, where the likelihood that a particular path occurs is indicated by its line width relative to that at the root of the tree. Paths in the interior of the tree in figure 2a are less likely than paths with
fewer switches that lie on the outside. This pattern becomes more pronounced at higher sampling accuracy and lower rates of switching (not shown). A second asymmetry is caused by unequal switching rates, which bias the weights of paths along the estimate tree towards the environmental state that is more frequent. In figure 2a, this effect reveals itself by the slightly increased thickness of paths in the right part of the tree.

The phenotype tree (figure 2b) is generally highly asymmetric because the optimal initial phenotype for a naïve individual, \(x_0\), is adapted to the most likely environmental state (in this case, state A). This is the typical outcome if the survival and fecundity functions are linear and the two environmental states are not equally frequent. As indicated by the relative thickness of the terminal branches of the phenotype tree, the initial phenotype has a prolonged effect on the phenotype distribution: at the final age \(T\), \(\beta / (\alpha + \beta) = 40\%\) of the individuals are in an environment in state B, but the optimal reaction norm induces less than 30% of the individuals to actually exhibit the phenotype \(x_T = 0\) adapted to this state. The reason is that some individuals in environment B made observation errors, preventing them from adjusting their phenotype from its initial value \(x_0 = 1\).

To quantify the rate of information accumulation and the degree of plasticity at various ages, we calculated the absolute change in forecasting probabilities \(\Delta p_t = |p_t - p_{t-1}|\) and phenotypes \(\Delta x_t = |x_t - x_{t-1}|\) for all sequences of observations, and averaged these values across the tree, weighting by the likelihood of each observation sequence across all possible realizations of the environment. The rate of information accumulation decreases monotonically with age (figure 2a, right part), before it asymptotes towards a stable level. This shows that organisms become better at estimating environmental states the more often they sample, although they are limited in the level of certainty they can achieve. Phenotypic plasticity (measured as \(E[\Delta x_t]\); figure 2b, right part) reaches a maximum in the second season and decrease over the final three seasons.

For the parameters considered in figure 2, no individuals adjust their phenotype in the first or the third season.

To illustrate the structure of the optimal reaction norm, we maximized equation (8) while treating \(p_t\) as a continuous state variable (in reality, \(p_t\) can only take a discrete set of values, one for each possible observation sequence). The resulting representation of the optimal reaction
norm $h_t$ (figure 3 shows results for $h_6$) reveals three regions in state space with qualitatively different optimal responses. First, there is a plateau at intermediate levels of $p_t$, where the optimal adjustment $h_t(x_{t-1}, p_t)$ is zero. This indicates that organisms have to acquire a particular level of certainty about environmental conditions before they adapt their phenotype. When the estimate $p_t$ lies either to the left or to the right of the plateau, it is beneficial to adjust the phenotype. If the fitness function is linear, it is always optimal to change to either $x_t = 0$ (at low values of $p_t$) or $x_t = 1$ (at high values). Indicated by black dots and curves, respectively, are the states and the transitions between states of the example individual from figure 2. Note that multiple, consistent observations are necessary to traverse the plateau and enter the region of phenotypic adjustment, helping to buffer the organism against observation errors.

The width of the plateau at age $T$ is equal to $c / s$ (see Appendix A), and phenotypic adjustment occurs only if $p_t < 1/2 - 1/2 c / s$ or $p_t > 1/2 + 1/2 c / s$. Therefore, as one would expect, phenotypic adjustment becomes less likely if the cost of plasticity, $c$, is high, or if the benefit of expressing an adapted phenotype, $s$, is low. The plateau disappears if $c = 0$. If $c > s$, on the other hand, organisms never adjust their phenotype in their final season, but they may still do so earlier in life. In line with this result, we observe the optimal reaction norm to depend on time. The width of the plateau is maximal at $t = T$ (for comparison, dashed lines in figure 3 outline the contours of $h_1$) such that there are states close to the edges of the plateau, for which organisms adjust their phenotype when they are young, but not when they are older.

The time-dependency of the reaction norm is strongest at the end of life, when it is necessary to compensate for the reduced levels of plasticity in the final life stages (particularly if $c > s$). However, these compensatory effects dampen out generally within a few backwards optimization steps, such that the reaction norms at early ages are indistinguishable in practice. The biological implication is that end-of-life-effects on patterns of plasticity are likely to be confined to the last few stages of an individual’s life history.

Depending on how organisms update their estimate $p_t$ after each observation, and how wide the plateau of the reaction norm is, the optimal reaction norm can be associated with a variety of realized phenotype sequences and resulting patterns of plasticity. Figure 4 illustrates the main effects of the various parameters of the model. In stable environments (figure 4a, left),
individuals adjust their phenotype early in life once they have become sufficiently confident that their initial phenotype is suboptimal. Trait reversal later in life is rare. The frequency of reversal to the initial phenotype goes up as the rate of environmental fluctuations increases, leading to a high average amount of phenotypic adjustment at intermediate values of $\alpha$ and $\beta$ (data not shown). Yet, in highly variable environments (figure 4a, middle), the organism can not always build up a confident estimate before the environment switches again, and any phenotypic adjustments that do occur are likely to be beneficial for only a short time. Hence, the overall level of plasticity decreases once the inherent unpredictability of the environment starts to limit the future benefits of phenotypic adjustment. In the example shown in the middle panel of figure 4a, we still find a plasticity window in the mid-life period, when the expected future benefits of phenotypic adjustment are still considerable and when at least a small subset of the organisms have made a series of consistent observations justifying an adjustment of the phenotype.

The amount of sampling that is needed to establish the current state of environment with a sufficient level of confidence is determined by the sampling accuracy. If observation errors are rare (figure 4b, middle) a single observation can be enough to trigger a phenotype change, whereas at lower sampling accuracy, organisms maintain their initial phenotype for a while before they start to specialize (figure 4b, left). Moreover, once specialized, individuals rarely reverse their phenotype. These results are explained by the fact that the sampling accuracy is related to how much the estimate $p_t$ changes after an observation (equation (2)). The estimate changes in small steps if the sampling accuracy is low, such that it may take several consistent observations to traverse the plateau of the reaction norm and enter the region of state space where phenotypic adjustment is beneficial. By contrast, when the sampling accuracy is high, the change in $p_t$ induced by an observation can be sufficient to jump over the plateau in one step, leading to an immediate adjustment of the phenotype after each observation.

Similar effects are observed by varying the cost of phenotypic adjustment (figure 4c). If adjusting the phenotype is costly (figure 4c, middle), the plateau of the reaction norm is wider, such that traversing the plateau requires a larger number of consistent observations (equivalent to decreasing the sampling accuracy). Conversely, if the cost of plasticity is low (figure 4c,
middle), the plateau is easily traversed in a single step, analogous to the situation at high
sampling accuracy.

*Iteroparous life histories with fecundity or viability selection*

Our main result for the fitness consequences of phenotypic adjustment (equation (5)) suggests
that the life history of an organism strongly influences its optimal plasticity schedule. For
example, a combination of life-history parameters appears as a factor $1 - \bar{F}_t \bar{S}_t / \bar{R}_t$ in front of
the expected future fitness effect, on the third line of equation (5). Life-history differences
therefore affect the relative weighting of current and future consequences of plasticity.
Furthermore, this weighting is different depending on whether the costs and benefits of
plasticity act on fecundity or on survival (the fecundity effects $f'_t$ and $f''_t$ are preceded by a
factor $\bar{F}_t \bar{S}_t / \bar{R}_t$, which reflects the relative importance of current reproduction).
To quantify the effects of life history on plasticity, we introduce a heuristic measure $I_t$ that
captures how important the immediate effects of phenotypic adjustment are relative to their
effects on future fitness components in the calculation of lifetime reproductive success
(equation (5)). Our definition is as follows:

$$I_t = \frac{s'_t + \bar{s}_t + \frac{F_t}{R_t} (f'_t + \bar{f}_t)}{s'_t + \bar{s}_t + \frac{F_t}{R_t} (f'_t + \bar{f}_t) + \left(1 - \frac{F_t}{R_t}\right) (s'_t + f'_t + \bar{s}_t + \bar{f}_t)}, \quad (9)$$

where $\bar{f}_t = \left(\alpha |f'_t| + \beta |f''_t|\right)/(\alpha + \beta)$ and $\bar{s}_t = \left(\alpha |s'_t| + \beta |s''_t|\right)/(\alpha + \beta)$ represent the average
strength of fecundity and viability selection at age $t$ across environments. The value of $I_t$ lies
between 0 and 1, with $I_t = 0$ corresponding to a situation in which current phenotypic
adjustments have no consequences for lifetime reproductive success (this may occur when the
cost and benefit of plasticity manifest themselves in the form of fecundity selection, and current
fecundity is negligible relative to the expected reproductive fitness in the future), and $I_t = 1$
indicating that only current reproductive success is relevant to the optimization of the reaction
norm (as, for example, at $t = T$). Accordingly, we refer to $I_t$ as the impact of current phenotypic
adjustment on the remaining lifetime reproductive success.

Low values of $I_t$ are expected to favor delayed phenotypic adjustment, for the reason that
postponing plasticity has limited consequences for current reproductive success, whereas it will
allow for additional observations before the organism commits to a costly phenotypic change. Given that \( I_t \) increases monotonically with \( F_t / \overline{R} \), we expect that, in iteroparous life histories, plasticity will be concentrated at those ages where individuals realize a large fraction of their lifetime reproductive success. Furthermore, this bias is predicted to be more pronounced if the cost and benefit of plasticity are mediated by effects on fecundity (as opposed to survival, as we have thus far assumed).

To illustrate these predictions, we calculated the optimal reaction norm for an example iteroparous life history based on published data from a life-table response experiment using the estuarine polychaete *Streblospio benedicti* (Levin et al. 1996; figure 5). *Streblospio benedicti* occupies soft mucoid sediment tubes from where it feeds either by extending its tentacles up into the water column or by sweeping its feeding palps across the sediment surface. We will therefore consider feeding mode as a potentially plastic phenotype that we will assume to be under divergent selection across environmental states. In our calculations, the observed fecundity and survival parameters from the original life-table response experiment (\( \phi_t \) and \( \sigma_t \); specified in online table A1 and plotted in figure 5) were modified by (hypothetical) costs of feeding-mode adjustments and the fitness advantage of expressing an adapted foraging strategy.

We considered two scenarios for this iteroparous life history, labeled as ‘viability selection’ (figure 6a) and ‘fecundity selection’ (figure 6b). In addition, we calculated the optimal reaction norm for a comparable semelparous life history (figure 6c), using identical values for the parameters \( T, \alpha, \beta, a, s \) and \( c \).

For the ‘viability selection’ scenario we assumed that all fitness effects of plasticity manifested themselves as changes in survival. The fecundity and survival functions were defined by

\[
F_t^A(x_t, \Delta x_t) = F_t^B(x_t, \Delta x_t) = \phi_t \quad \text{and} \quad \sigma_t^A(x_t, \Delta x_t) = \begin{cases} 
\sigma_t \exp\left(-s(1-x_t) - c \Delta x_t\right) & \text{if } e_t = A, \\
\sigma_t \exp\left(-s x_t - c \Delta x_t\right) & \text{if } e_t = B.
\end{cases}
\]

The optimal phenotype tree under these conditions (figure 6a; left) is difficult to distinguish from the result for the semelparous history (figure 6c; left): small differences in the expected amount of phenotype change occur from age 5 onwards (figure 6ac; right). These findings are consistent with the impact profiles \( I_t \) of the two life histories (figure 6ac; middle), which are
overall comparable, except for the final age classes, where $I_t$ for the iteroparous life history increases as a result of the decline of fecundity rates towards the end of life.

The fecundity and survival schedules in the ‘fecundity selection’ scenario were defined as:

$$ F^c_t(x_t, \Delta x_t) = \begin{cases} \varphi_t \exp(-s(1-x_t)) & \text{if } e_t = A, \\ \varphi_t \exp(-s x_t) & \text{if } e_t = B, \end{cases} $$

$$ S^A_t(x_t, \Delta x_t) = S^B_t(x_t, \Delta x_t) = \sigma_t \exp(-c \Delta x_t), $$

such that the costs of plasticity reduced survival, while the expression of an adapted phenotype was favored by fecundity selection. In this case, as reflected by the impact profile, plasticity provides limited benefits before the organism has actually started to reproduce, leading to a delay in the onset of plasticity relative to the semelparous life history (figure 6bc). Also in this case, a comparison of the impact profiles explains the main differences between the plasticity schedules of the iteroparous and semelparous life history. However, without a base for comparison, the impact profile is a poor predictor of the absolute levels of phenotypic adjustment, because the schedule of plasticity is affected primarily by the dynamics of information accumulation. For instance, even in figure 6b, there is a peak of plasticity early in life at the onset of reproduction, when the impact $I_t$ is still relatively low.

**Discussion**

The responsiveness of phenotypically plastic organisms to cues from the environment often varies with age. Various empirically observed patterns of age-dependent plasticity have been suggested to result from changes in the availability, reliability and usefulness of environmental information over the course of an individual’s life (Dufty et al., 2002). To formally evaluate this idea, we have modeled the developmental trajectory of an organism living in a stochastically fluctuating environment, about which the organism obtains information by sampling at regular intervals throughout its life. The evolutionarily optimal response for such an organism is to adjust its phenotype only if it is sufficiently confident of the current state of the environment. Accordingly, for linear and certain nonlinear (online figure A1b) fitness functions, a characteristic feature of the optimal reaction norm is that it has a plateau at intermediate values of the state variable $p_t$, which represents the organism’s current estimate of
the state of the environment (figure 2). The width of the reaction norm’s plateau is dependent on the ratio between the cost of phenotype adjustment and the benefit of expressing the optimal phenotype, $c / s$. In our model, the dynamic of an individual’s state in the state space spanned by the optimal reaction norm $h_t$ is specified by a Bayesian update rule (2), which takes into account the reliability of a single observation, measured by the sampling accuracy $a$, and the inherent uncertainty of the environment, captured by the switching rates $\alpha$ and $\beta$. These parameters determine how many observations are needed for the estimate $p_t$ to traverse the width of the plateau and, correspondingly, how quickly the organism will respond to a change in its environment.

According to our analysis, the interplay of environmental uncertainty and the accuracy of perceived information with life-history determinants and the fitness consequences of phenotypic adjustments must be expected to result in three distinct features of the pattern of age-dependent plasticity. First, the plateau of the reaction norm and the limited accuracy of perceived information typically cause a delay in the response of the organism to its environment, during which it integrates multiple observations into a sufficiently reliable estimate of the state of the environment. Moreover, older individuals take longer to respond to an environmental change during their lifetime than it takes a newborn individual to adjust its phenotype to the environmental condition at the start of its life. This is because newborn individuals have limited prior information about the state of the environment, whereas older individuals are biased by the information they have accumulated earlier. Correspondingly, a naïve newborn individual starts sampling with a state located on the reaction norm’s plateau, and can therefore more easily be induced to adjust its phenotype than an older individual, whose estimate of the state of the environment, $p_t$, must generally first traverse at least the entire width of the plateau before a phenotypic adjustment will occur. Finally, we observed a reduction of plasticity towards the end of life under most parameter conditions and life histories explored by us. This effect is particularly pronounced if the benefit of expressing the optimal phenotype is small relative to the cost of phenotypic adjustment, such that it does not pay to adjust the phenotype unless the individual can profit from this adjustment for several additional time steps before it dies. The combination of these early-, mid- and late-life effects can produce a variety
of optimal age-dependent plasticity patterns, which are found to be non-monotonic in general. A common pattern, which may be more or less pronounced depending on the parameters considered (c.f. figure 2b, 4, 6), features a (delayed) peak of phenotypic adjustments early in life (corresponding to the initial phenotypic adjustment by young individuals after they have accumulated sufficient information). After a period of reduced plasticity, the first peak of plasticity is followed by a second, broader one, which is caused by individuals responding to a change of the environment during their lifetime who still expect to live long enough to benefit from a phenotypic adjustment.

To further explore the effects of life history on optimal patterns of age-dependent plasticity, we extended our analysis to an iteroparous example life history based on demographic data of the estuarine polychaete *S. benedicti* (Levin et al. 1996). Phenotypes were exposed either to simulated viability or fecundity selection. Under viability selection, results for the iteroparous life-history are similar to the predictions for a basic semelparous life-history: the calculated optimal schedule of phenotypic switches is nearly identical (figure 6ac). If the fitness effects of phenotypic changes are mediated by differences in survival, it is risky to postpone phenotypic switches if they are beneficial, because both current and future reproductive success are conditional on current survival. By contrast, under fecundity selection it is optimal to delay phenotypic adjustments until shortly before reproduction takes place, thus allowing for the accumulation of additional information and the subsequent maximization of the benefits of plasticity at the time of reproduction (figure 6b; cf. figure 5). Therefore, under fecundity selection, we would predict a single adjustment to the current environment at the penultimate time step before reproduction in a semelparous life history, and a corresponding delay in plasticity until the onset of reproduction in an iteroparous life history (figure 6b). These predictions need to be adjusted in situations where organisms use more than one season to accumulate the resources necessary for reproduction. As indicated earlier, a formal analysis of such cases needs to take into consideration additional state variables (e.g., the amount of energy reserves stored for reproduction). Although we have not performed this analysis, we expect that storing resources for reproduction would have similar effects for the expression of plasticity as
shifting part of the reproductive output to earlier reproductive seasons, i.e., plasticity would be expressed earlier in life.

An alternative mechanism also mentioned by Dufty et al. (2002) that could possibly be responsible for age-dependent plasticity are developmental constraints arising in the course of ontogeny. Developmental constraints would lead to increasingly canalized phenotypes while organisms pass through certain ontogenetic stages. Our model assumes that the range of attainable phenotypic states does not decrease with age. In this way, we could show that information gain can give raise to age-dependent changes of plasticity as an emergent pattern without a priori introducing hard constraints on the attainable range of phenotypes at different ages. Such constraints could, however, easily be included in our model to produce more detailed predictions. Dufty et al. (2002) proposed that later in life, information is only used for phenotypic fine-tuning, since developmental trajectories have been fixed already early in life. However, life-long plasticity in leg length of barnacles (Marchinko 2003) is a counter example to this suggestion. Further research is needed to clarify whether there are generalities in the way developmental constraints change during ontogeny. Our results show that, even without developmental constraints, plasticity later in life is generally expected to be lower than early in life.

Age-dependence of phenotypic adjustment costs constitutes a third alternative mechanism that might cause age-dependent plasticity. Certain phenotypic responses induced during late ontogeny might cause greater (or smaller) costs than if induced early in life (Hoverman and Relyea 2007, Callahan et al. 2008). Likewise sampling accuracy might change with age, for instance, because accuracy is enhanced over time by learning. Similar to developmental constraints, specific assumptions on age-dependent costs and sampling accuracy could readily be included in the model, but for the sake of simplicity and generality, we did not do so during this study. We also did not account for maintenance costs of plasticity in our model, which are associated with developing and maintaining the sensory and neural machinery necessary for processing environmental information (Scheiner and Berrigan 1998, DeWitt 1998, Van Buskirk and Steiner 2009, Auld et al. 2010). The magnitude and importance of maintenance costs of plasticity is debated. Recent empirical studies suggest that maintenance costs may be modest in
the majority of cases (Van Buskirk and Steiner 2009). We expect that maintenance cost would influence the optimal level of plasticity but not otherwise affect the optimal pattern of age-dependent plasticity.

To our knowledge, no theoretical study has so far investigated possible mechanisms for the evolution of age-dependent plasticity. However, some theory exists on the evolution of reversible plasticity (Gabriel 1999, Gabriel et al. 2005). These studies investigated how lag times in phenotypic responses and the quality of an organism's environmental information affect optimal plasticity. Gabriel et al. (2005) considered the two extreme cases of complete information and no information gain through sampling only. Their analysis shows that non-specialist phenotypes are superior to phenotypes that track environmental change if there is a lag in the phenotypic response, since lag-times cause temporary maladaptation, reducing the fitness of plastic phenotypes. These models also predict that organisms should express less specialized phenotypes when information is incomplete than with perfect information, a result that is also supported by our findings. Our model adds a life-history perspective to the existing theory on the evolution of plastic responses by showing that plasticity can vary not only between different environments (Marchinko 2003, Relyea 2003) but also with age.

We are not aware of empirical studies that have tracked plastic adjustments throughout different individual life histories, so the predictions of our model can only be tested indirectly against empirically observed patterns of plasticity. For example, in barnacles, *Balanus glandula*, wave action is highly correlated with leg length (Arsenault et al. 2001). Barnacle leg length is a plastic trait that responds very rapidly to new flow conditions. Hence, our model predicts that the either the sampling accuracy must be high in this system or the cost-to-benefit ratio of plasticity $c/s$ must be low. Barnacles are iteroparous, hermaphroditic, sessile organisms, which reproduce several times per year. Non-adjusted leg length leads to a suboptimal food intake, which is likely to affect immediate reproductive success through viability or fecundity selection. Transitions between environmental states (high flow vs. low flow) are frequent (Arsenault 2001). In our model, the combination of these factors would tend to favor lifelong plasticity and frequent phenotypic adjustments, corresponding to the pattern observed in barnacles
(Marchinko 2003). However, the degree of plasticity predicted by our model depends strongly on the adjustment cost, which has not yet been estimated empirically.

Another example is provided by the snail *Helisoma trivolvis*, which can adjust the size of its shell to the presence of predatory water bugs (*Belostoma flumineum*) in its environment (Hoverman and Relyea 2007). *Helisoma trivolvis* is an iteroparous species living in semi-permanent ponds that can be colonized by water bugs at any time during development or adulthood. Adult water bugs as well as their nymphs are aquatic and prey on snails, so plasticity is likely to confer a viability selection advantage. Snails respond to the presence of predators by producing larger shells. Reversal of this trait is possible only in early development, whereas induction is possible for longer. Partial irreversibility of plasticity is predicted by our model when the environment is relatively stable and the sampling accuracy is low (e.g., figure 4b).

Yet, for *Helisoma* also developmental constraints are likely to play an important role, because the shape of the shell cannot be altered once deposited (Hoverman and Relyea 2007).

Organisms living in a stochastically fluctuating environment with a limited ability to read environmental cues need to integrate current and past information, in order to optimally adjust their phenotype to the state of the environment. We conclude that the accumulation of information during life and the optimal response of the organism in the context of its life history are sufficient to produce striking patterns of age-dependent plasticity. Depending on the rate of environmental fluctuations, the accuracy of sampling, phenotypic adjustment costs and on the fitness component that is most strongly affected by selection (i.e., survival or reproduction), a diversity of age-dependent plasticity patterns can emerge. While these patterns correspond to the wide variety of plasticity schedules observed in nature and expressed across species, it is unlikely that these organisms use the exact complex Bayesian update rule assumed in our analysis. Instead, biological organisms often build on simple 'rules of thumb' when navigating complex environments (Welton et al. 2003, McNamara and Houston 2009). Future research should therefore explore whether there are simple decision rules for age-dependent plasticity that generate similarly efficient responses to stochastic environments as the rules assumed in our analysis.
Acknowledgements

We are grateful to J. Johansson, Peter Taylor and four anonymous reviewers for their valuable comments on various versions of this manuscript. This study was funded by the Austrian Science Fund FWF (grant P18647-B16 to B.T.), the Swiss National Science Foundation SNF (grant 3100A0-111796 to B.T.), the Research Council of Norway (grant 214285 to B.F.), the Netherlands Organization for Scientific Research (NWO VIDI grant 864.11.012 to G.S.vD.) and the European Research Council (ERC Starting Grant 30955 to G.S.vD.). U.D. gratefully acknowledges financial support by the European Commission, the European Science Foundation, the Austrian Science Fund, the Austrian Ministry of Science and Research, and the Vienna Science and Technology Fund.
Literature cited


**Figure legends**

**Figure 1.** Order of events during a single time step.

The sequence of events that occur during a single time step starts with the determination of the environmental state (step 1). Each individual then samples the state of the environment (step 2), and uses the resulting observation to update its personal estimate $p_t$ (step 3). Based on its new state, each individual then decides if and by how much it will adjust its phenotype (step 4). Viability selection acts at the end of each time step, potentially followed by the production of offspring. Both survival and fecundity are allowed to depend on the match between the current phenotype and the environment. In addition, in order to incorporate costs of plasticity, both fitness components may decrease as a function of the absolute size $\Delta x_t = |x_t - x_{t-1}|$ of the latest phenotypic adjustment step.

**Figure 2.** Estimated environmental conditions and resultant plastic phenotypes for a semelparous life history.

In the left column, grey lines and circles indicate (a) the estimated environmental conditions $p_t$ and (b) the resultant phenotypes $x_t$ throughout the lifetime of a single individual that is making the sequence of observations $o = (B, B, B, A, A, A)$. Also in the left column, black lines show (a) the tree of estimates and (b) the tree of phenotypes characterizing the ensemble of many individuals, each experiencing its own personal sequence of observations for a randomly drawn realization of the sequence of environmental states. The likelihood of a particular estimate or phenotype to occur along the tree is proportional to line thickness and depends on the rate of environmental fluctuations and on the sampling certainty. In the right column, black bars indicate the absolute change of (a) $p_t$ and (b) $x_t$, averaged over the distribution of all observation sequences. For this example, the optimal reaction norm leads to plasticity at age 2 and during the second half of life. Parameters: $T = 6$, $\alpha = 0.15$, $\beta = 0.1$, $a = 0.7$, $s = 0.05$, and $c = 0.02$.

**Figure 3.** Optimal reaction norm, describing the phenotypic adjustment $h_t(x_{t-1}, p_t)$ as a function of the current estimate $p_t$ of environmental conditions and the previous phenotype $x_{t-1}$.

The shaded surface depicts the optimal reaction norm $h_6$ at age 6, while the dashed lines outline
the optimal reaction norm $h_1$ at age 1. The plateau at intermediate values of $p_1$ applies to individuals that are not sufficiently certain about the state of the environment to adjust their phenotypes. This plateau is flanked by two ranges of conditions under which individuals change their phenotype to either $x_1 = 0$ (left-hand side, where $p_1$ is low) or $x_1 = 1$ (right-hand side, where $p_1$ is high). The width of the plateau equals $c / s$ at age $T$ (appendix A), being more narrow at younger ages. Filled circles connected by curved black lines indicate the change of state variables for the particular individual shown in figure 2, which is making the sequence of observations $o = (B, B, B, A, A, A)$. Parameters are as in figure 2.

**Figure 4.** Dependence of optimal plasticity patterns on model parameters.

Comparisons between optimal patterns of plasticity ($a$) in environments with rare transitions (approximately once every four lifetimes; black lines in left column) and frequent transitions (approximately twice per lifetime; grey lines in middle column) between the two environmental states, ($b$) between low (black lines in left column) and high (grey lines in middle column) sampling certainty, and ($c$) between low (black lines in left column) and high (grey lines in middle column) cost of plasticity. As in figure 2$b$, the right column shows the absolute phenotypic adjustment, averaged over the distribution of all observation sequences, with black bars corresponding to the left column and grey bars to the middle column. Parameters, where not indicated otherwise: $T = 8$, $\alpha = 0.12$, $\beta = 0.1$, $a = 0.7$, and $s = c = 0.05$.

**Figure 5.** Age-specific fecundity and survivorship schedules for *S. benedicta*.

Data points show the survivorship (filled circles) and weekly fecundity (open circles; normalized so as to yield a lifetime reproductive success of 1) observed in the control treatment of a life-table response experiment with the estuarine polychaete *Streblospio benedicta* (Levin et al. 1996; data were collected for a cohort of 50 individuals). Black lines show the observed survivorship smoothed over a three-week period using least-squares smoothing (figures 2 and 3 of Levin et al. 1996). Before using these empirical observations to parameterize our model, we first partitioned the data into 12 age classes, and calculated the expected survival and fecundity over each of the resulting seven-week periods (values provided in online table A1):
the thus determined survivorship and normalized fecundity are presented as dark-grey and light-grey histograms, respectively, and have retained the main features of the original data.

**Figure 6.** Dependence of optimal plasticity patterns on life-history types and selection regimes. Shown in the left column are the trees of phenotypes resulting from the optimal reaction norm for \((a, b)\) an iteroparous life history and \((c)\) for a semelparous life history. Adapted phenotypes benefit either from reduced mortality \((a\ and\ c;\ viability\ selection)\) or from increased fecundity \((b;\ fecundity\ selection)\). The central and right column, respectively, show the relative importance of immediate and future fitness effects, measured by the impact \(I_t\) of current phenotypic adjustment, and the average absolute phenotypic adjustment for the phenotype trees on the left. Parameters: \(T = 12, \alpha = 0.081, \beta = 0.086\) (i.e., the environment switches between states on average once per 12 time steps), \(a = 0.667, s = 0.05,\) and \(c = 0.02\). Using the definition in the main text, the impact of current phenotypic adjustment is calculated as follows for \(t < T\): \[(a)\ I_t = \frac{1}{2},\ (b)\ I_t = \frac{R_t}{(2\ R_t - \phi_t \sigma_t)},\ and\ (c)\ I_t = \frac{(c\ R_t + s\ \phi_t \sigma_t)}{(2\ c + s\ R_t - c\ \phi_t \sigma_t)}.\] In all three cases, \(I_T = 1\). The life-history parameters \(\phi_t, \sigma_t,\) and \(R_t\) for the iteroparous life history are listed in online table A1.