

1	
2	
3	
4	
5	
6	A heuristic model on the role of plasticity in adaptive evolution:
7	plasticity increases adaptation, population viability, and genetic variation
8	
9	
10	
11	Ivan Gomez-Mestre [*] and Roger Jovani
12	
13	Estación Biológica de Doñana, Consejo Superior de Investigaciones Científicas,
14	Avda. Americo Vespucio s/n, Isla de la Cartuja,
15	Seville E-41092, Spain
16	
17	
18	*Corresponding author
19	igmestre@ebd.csic.es
20	
21	
22	
23	
24	
25	
26	
27	
28	SUMMARY

29 An ongoing new synthesis in evolutionary theory is expanding our view of the sources 30 of heritable variation beyond point mutations of fixed phenotypic effects to include 31 environmentally-sensitive changes in gene regulation. This expansion of the paradigm is 32 necessary given ample evidence for a heritable ability to alter gene expression in 33 response to environmental cues. In consequence, single genotypes are often capable of 34 adaptively expressing different phenotypes in different environments, i.e. are adaptively 35 plastic. We present an individual-based heuristic model to compare the adaptive 36 dynamics of populations composed of plastic or non-plastic genotypes under a wide 37 range of scenarios where we modify environmental variation, mutation rate, and costs of 38 plasticity. The model shows that adaptive plasticity contributes to the maintenance of 39 genetic variation within populations, reduces bottlenecks when facing rapid 40 environmental changes, and confers an overall faster rate of adaptation. In fluctuating 41 environments, plasticity is favoured by selection and maintained in the population. 42 However, if the environment stabilises and costs of plasticity are high, plasticity is 43 reduced by selection, leading to genetic assimilation, which could result in species 44 diversification. More broadly, our model shows that adaptive plasticity is a common 45 consequence of selection under environmental heterogeneity, and hence a potentially 46 common phenomenon in nature. Thus, taking adaptive plasticity into account 47 substantially extends our view of adaptive evolution.

48

49 Keywords: adaptation, developmental plasticity, genetic accommodation,

50 heterogeneous environment, selection, bottleneck, genetic variation.

- 51
- 52

53 1. INTRODUCTION

54 Understanding the mechanisms of adaptation is key to understand how life on 55 earth has persisted over widely varying environmental conditions resulting in the 56 observed biodiversity, and to understand how organisms would adapt to current global 57 change. Adaptive evolution requires heritable phenotypic variation for selection to act 58 upon, and the standing paradigm that emerged from the Modern Synthesis argued that 59 random genetic mutations of fixed phenotypic effects are the only source of heritable 60 phenotypic variation fuelling adaptive evolution [1-3]. Under this scenario, mutations 61 accumulate in populations through various combinations of recurrent mutation, drift, 62 recombination, immigration, and selection in heterogeneous environments [4-6]. 63 Selection then acts on this standing genetic variation producing adaptations, and hence 64 the environment acts merely as a sieve for phenotypes.

65 Nevertheless, there is now ample evidence showing that the environment can 66 also act as a phenotypic inducer so that a single genotype is often capable of expressing 67 alternative appropriate phenotypes in response to different environments [7-9]. This 68 phenotypic plasticity is the consequence of environmentally-induced changes in gene 69 expression [10]. Plasticity is often heritable, and it evolves under selection if 70 environmental cues are reliable and gene flow is high among subpopulations [11, 12]. 71 Conversely, local adaptation and reduced plasticity occur when dispersal is low [11] or 72 environmental variation is unpredictable or negligible [13, 14].

Extending the paradigm to include adaptive plasticity is a necessary step in evolutionary biology to extend our understanding of the mechanisms of adaptive evolution [15], and there has been a surge of interest in characterising the evolutionary consequences of environmentally induced variation [16-18]. Previous theoretical studies have greatly contributed to our understanding of different aspects of the evolution of plasticity under particular scenarios, and often using complex quantitative genetic

models [19-22]. These models have shown that plasticity is advantageous in rapidly
changing environments and that it may help colonising new environments [22],
although genetic correlations and costs of plasticity could limit these benefits of
plasticity [23, 24].

83 Adaptive plasticity can also result in evolutionary innovations [18]. If sister 84 lineages evolve independently in different stable environments and ancestral plasticity is 85 costly, divergent reaction norms are expected to evolve through selection on genetic 86 modifiers available in the population [2, 7, 25]. This would lead to genetic 87 accommodation of environmentally induced phenotypes, i.e. adaptive genetic changes 88 in response to selection on the regulation and form of the phenotype [7]. Fixed-effect 89 genes (i.e. not sensitive to environmental input) giving rise to phenotypes with 90 increased fitness in the new environment will be positively selected, and the trait will 91 become genetically assimilated, a particular case of genetic accommodation [7, 26]. 92 Thus, whether resulting in novel or canalised phenotypes, or simply in divergent 93 reaction norms, developmental plasticity can foster speciation and diversification [17, 94 27]. Genetic accommodation and assimilation of plasticity have been experimentally 95 demonstrated [28-30] and also inferred from comparative analyses [31, 32]. Plasticity is 96 thus a common feature of organisms that is favoured by selection precisely under the 97 same circumstances that maintain standing genetic variation, namely environmental 98 heterogeneity and gene flow among subpopulations [11]. However, historically there 99 has been some reluctance to recognise the importance of phenotypic plasticity in 100 evolution [3, 9, 21, 33, 34]. Perhaps simple heuristic models may help illustrating the 101 potential of plasticity in evolution while avoiding the so-often black-box feeling of 102 complex models.

103 Here we built and analysed a simple heuristic individual-based model comparing 104 adaptive evolution in populations composed of either plastic or non-plastic genotypes. 105 We examine how adaptive plasticity evolves under common scenarios assumed to 106 maintain non-environmentally dependent standing genetic variation, and then examine 107 how plasticity affects adaptive evolution because of the role of the environment as a 108 phenotypic inducer. We simulated population dynamics under contrasting combinations 109 of environmental stochasticity, occurrence of genetic changes, levels of plasticity, and 110 costs of plasticity. We specifically explored the conditions under which genetic 111 assimilation occurs, and the relationship between plasticity and standing genetic 112 variation. There is also evidence that in some organisms epigenetic marks allow induced 113 phenotypes themselves (and not just the ability to produce them) to be inherited across 114 multiple generations [35, 36], but that is not the scope of the present study. Here we 115 focus only on plastic genotypes that inherit the ability to produce different adaptive 116 phenotypes according to perceived environmental cues. 117 We used the model to test the following predictions: i) during rapid 118 environmental change or when facing a novel environment, plasticity improves the 119 persistence of populations and reduces the severity of bottlenecks; ii) plasticity 120 contributes to the maintenance of standing genetic variation within populations; iii) by 121 increasing population persistence and maintaining genetic variation, plasticity "buys 122 time" for appropriate genetic variants of fixed phenotypic effect to appear by mutation; 123 iv) costs of plasticity result in genetic assimilation (i.e. loss of plasticity) if 124 heterogeneous environments stabilise. 125

126 **2. THE MODEL**

127 This model description follows the ODD (Overview, Design concepts and Details)

128 protocol for describing individual- and agent-based models [37-39]. The model is

129 implemented in NetLogo 5.0.3 [40], (NetLogo is freely downloadable from

130 <u>http://ccl.northwestern.edu/netlogo/download.shtml</u>) and available in the electronic

131 supplementary material (Model.nlogo).

132 *— Purpose.* The main purpose of the model is to explore the consequences of 133 phenotypic plasticity in adaptive evolution. This is done by simulating population 134 persistence and genetic evolution under environmental change. Simulations are run separately for non-plastics and plastics. Non-plastics evolve by selection on random 135 136 genotypic mutations with fixed phenotypic effects. *Plastics* evolve exactly in the same 137 way, but also through selection on mutations conferring phenotypic plasticity (figure 1) 138 - Entities, state variables and scales. Environmental conditions are simulated by the 139 variable *environment*. The entities of the model are asexual individuals of two kinds: 140 either *non-plastics* or *plastics*. Each individual has a given *genotype* and a *phenotype*. 141 Plastics also have a plasticity-range that allows them to improve their match with the 142 environment. The match is an individual variable calculated as 1 - | phenotype -143 environment, which shapes individual survival and reproduction (see below). The 144 amount of *plasticity-range* used by the individual to improve its phenotypic match with 145 the environment is the *used-plasticity*. For instance, a *genotype* of 0.7 in an *environment* 146 of 0.8 with a *plasticity-range* of 0.2 would only need to use 0.1 of its *plasticity-range* to 147 produce a perfectly matching *phenotype* (i.e. *used-plasticity* = 0.1). Thus, while 148 plasticity-range is an inherited trait of the individual, plasticity-used is a value recorded 149 by the model when the individual develops. One time step of the model corresponds to 150 one generation, and generations are non-overlapping. See table 1 for variable definitions 151 and range of parameterised values.

152 — Process overview and scheduling. See a schematic diagram in figure 1. At birth, 153 individuals inherit from their parent a genotype and (if plastics) a plasticity-range. Both 154 genetic features mutate in the same way (see 'mutation' below). Non-plastics develop a 155 phenotype equal to their genotype. Plastics, however, use their plasticity-range to fit 156 their phenotype as much as possible to the environment (see 'development' below). Non-157 *plastics* and *plastics* have a mortality probability according to their realized *match* to the 158 environment (see 'die-by-mismatch?' below). Subsequently, they can die by negative 159 density-dependence (see 'die-by-negative-density-dependence?' below). Moreover, 160 plastics could die by costs of maintaining a given plasticity-range and the costs of the 161 *plasticity-used* (see '*die-by-plasticity-costs*?' below). These two costs of plasticity are 162 commonly identified in the literature on developmental plasticity as 'maintenance costs' 163 and 'production costs' and correspond to the presumed costs of maintaining a sensory 164 machinery and actually producing alterations on the phenotype, respectively [23, 24]; 165 see electronic supplementary material).

166 Surviving individuals reproduce (see 'reproduction' below) and die immediately 167 after. The *environment* is updated before the new generation is born, starting the cycle 168 again. The *environment* is thus updated between the death of generation t and the birth 169 of generation t+1 (see '*environmental-change*' below). In this way, newborns can adjust 170 (if *plastics*) their *phenotype* according to the *environment* where they will live until 171 death; and this is the *environment* that will affect their survival and reproduction. -Design concepts. Evolution (changes in population mean/variance values of 172 173 genotypes, either plastic or non-plastic, and plasticity-range) and other population 174 dynamics (e.g. stability, bottlenecks, extinction) emerge from the combined effects of 175 heredity, phenotypic plasticity (for *plastics* only), natural selection (differential survival 176 and reproduction of individuals), and demographic (density-dependence) processes.

177 Also, population genetic variability (either *genotype* or *plasticity-range*) is not imposed 178 at initialization, but emerge during the first 100 generations when the population 179 evolves under a mildly fluctuating environment (see 'environmental-change' below). 180 Note that the *genotype* and the *phenotype* could potentially take any real value, but in 181 simulations tended to remain between 0 and 1 because of the selection imposed by the 182 *environment* and the initialization conditions (i.e. genotype = phenotype = 0.5; see 183 figure 2 insets and figure 3c). Stochasticity affects environmental change, mutation, 184 survival probability and reproduction.

185 We recorded the number of individuals at the end of 300 generations (100 of

186 them being the initialization generations). For illustrative purposes, we also recorded for

187 some model runs longitudinal (e.g. environmental fluctuations, population size

188 dynamics, mean population *genotype*, *phenotype*, and *plasticity-range*) and transversal

189 data (e.g. *genotype* of each individual) across and within generations, respectively.

190 — *Initialization*. Simulations were initialized with *environment* = 0.5 and 100

191 individuals (either *mutants* or *plastics*). All individuals started with *genotype* =

192 phenotype = 0.5. Plastics started with plasticity-range = 0.

Input. The model does not have any external input; the *environment* is updated
according to internal model rules.

195 — Submodels

196 — *'environmental-change'*: During the first 100 generations of a simulation the

environment tightly fluctuates around 0.5. This is achieved by changing the

198 *environment* towards 0.5 by increasing (or decreasing) the *environment* by a

199 pseudorandom number extracted from a normal distribution with mean = 0.5 and

200 variance arbitrarily fixed at 0.01 to ensure small fluctuations of the *environment*

around 0.5. For the next 200 generations, the *environment* fluctuates every

202 generation according to the value of a pseudorandom number extracted from a 203 normal distribution with zero mean and *Std-Dev-environment-change* variance. To 204 test the adaptive response to rapid directional changes and the role of costs of 205 plasticity in causing genetic assimilation, we also modelled a scenario in which the 206 *environment* fluctuates during the first 100 generations as in the other simulations, 207 but then rapidly drift upwards in steps of 0.015 from 0.5 to 1, then remaining at 1 for 208 the rest of the simulation.

209 — *'reproduction'*: Each individual produce $match \times 2$ individuals, rounded to the

210 nearest integer; i.e. they produce either 0, 1 or 2 individuals according to their *match*.

211 — 'mutation': The genotype and the plasticity-range (if plastics) inherited from the

212 parent mutate by extracting a pseudorandom number from an exponential decay

213 distribution with mean *mean-mutational-change* (see electronic supplementary

214 material). This number is either added or extracted to the inherited trait with equal

215 probability. In this way, we are jointly modelling the probability of mutation and the

216 magnitude of its effect on the phenotype. Given the many sources and kinds of

217 mutations, we preferred this approach over simply modelling a per base per

218 generation substitution rate (see electronic supplementary material).

219 — 'development': Non-plastics develop a phenotype = genotype. Plastics, however,

220 use their *plasticity-range* to produce a *phenotype* as close as possible (given their

221 *plasticity-range*) to the *environment*. The amount of *plasticity-range* eventually used

is called *used-plasticity* (i.e. $0 \le used-plasticity \le plasticity-range$).

223 — '*die-by-mismatch?*': Individuals can die because of a low *match* with the

- 224 *environment*. They do so with probability 1 *match*, i.e. extracting a pseudorandom
- number from a uniform distribution from 0 to 1, dying if this number is > *match*.

226 - 'die-by-negative-density-dependence?': Plastics and non-plastics die because of 227 negative density-dependence when (before reproduction) population size is above 228 100 individuals. The dying individuals are those with lower *match* with the 229 *environment* (note that in any given model run all individuals are either *plastics* or 230 *non-plastics*, so there is no competition between these types). 231 — 'die-by-plasticity-costs?': With the same approach, *plastics* can also die first 232 with probability = *plasticity-range* * *plasticity-costs*, and then also with probability = 233 used-plasticity * plasticity-costs. That way, increased plasticity costs penalise 234 separately plasticity maintenance and plasticity use. Maintenance is associated with 235 the ability of being plastic, i.e. *plasticity-range*; the broader the range of possible 236 phenotypes, the highest the cost. Production costs, however, are the costs incurred 237 when actually altering the phenotype (i.e. *used-plasticity*; see electronic 238 supplementary material).

239

240 Simulations

241 Simulations for *non-plastics* and *plastics* are run independently but using the same 242 pseudorandom generator seed to make results fully comparable. For each group we ran 243 a total of 200 simulations for each of the 4,056 combinations of 26 (equally spaced) 244 values for Std-Dev-environment-change, 26 different values for mean-mutational-245 change and six values of plasticity-cost i.e. a total of 811,200 model runs (see Table 1 246 for parameter details). For each of the 4,056 parameter combinations we calculated 247 (separately for *non-plastics* and *plastics*) population size at the end of the simulations 248 and the cumulated population size along the 200 generations after initialization. Note 249 that we run 200 simulations for each of the 4,056 parameter combinations for *plastics* 250 and *non-plastics* although parametrisations only differing in the *plasticity-cost* value do

not affect *non-plastics*. This way results from *plastics* were directly comparable with
simulations (with same pseudorandom generation seeds) for *non-plastics*. To test
hypothesis (iv) regarding genetic assimilation in a novel environment we also modelled
a scenario with an abrupt directional environmental change, which then stabilised (see
above). This could represent either the colonisation of a novel habitat, or a rapid
environmental transformation such as those occurring as a consequence of global
change across the world.

258 **3. RESULTS**

259 During the first 100 generations of the model runs the environment was forced to 260 remain close to 0.5 and the initial generation had genotype = 0.5 and *plasticity-range* (if 261 *plastics*) = 0. In all simulation runs, *plastic* and *non-plastic* populations survived these 262 initial generations, generating standing genetic variation and (in *plastics*) variation in 263 *plasticity-range* (figure 2 insets). As plasticity costs increased, population size during 264 the first 100 generations of initialisation was lower for *plastics* than for *non-plastics* 265 (see examples for intermediate plasticity costs in figure 3b), indicating that under low 266 environmental fluctuations plasticity costs may outweigh the benefits of plasticity.

267 Adapting to a fluctuating environment

268 Afterwards, when the *environment* was allowed to vary stochastically along 200

269 generations, the *plastic* and *non-plastic* populations began evolving to adapt to the

270 changing *environment*. Both *plastic* and *non-plastic* populations were capable of

271 persisting over simulated environmental fluctuations provided that the *mean*-

272 mutational-change was high, but population viability was severely compromised as

environmental fluctuations increased (figures 2 and 3). At low environmental

274 fluctuations, *plastics* always performed slightly worse than *non-plastics* during the next

200 generations (figure 3*a*, and first panel of figure 3*b*). This also supports the idea that
plasticity even at low plasticity costs has demographic consequences when occurring at
low environmental fluctuations.

278 Selection favoured increased plasticity during bouts of rapid, recurrent, or wide 279 environmental shifts (figure 2 main panels), often being the most plastic genotypes the 280 ones that persisted (see examples in figures 3c and 4a). Costs of plasticity reduced the 281 effectiveness of the plastic response and when taken to the extreme ultimately made 282 *plastic* genotypes evolve analogously to *non-plastic* ones (figure 3a). Except in such 283 scenarios of extreme costs of plasticity, *plastic genotypes* always showed a better 284 phenotypic match to the environment than non-plastic ones, even at high mean-285 *mutational-change* (figure 2 main panels).

286 At higher *Std-Dev-environment-change* selective sweeps of poorly matched 287 genotypes were more frequent and resulted in population bottlenecks (figure 3b), 288 reducing the likelihood of persistence for both *plastic* and *non-plastic* genotypes (figure 289 3a). Population viability of *non-plastics* was restricted to low environmental 290 fluctuations and high *mean-mutational-change* (figure 3a). *Plastic genotypes*, however, 291 experienced attenuated population bottlenecks because a greater fraction of *genotypes* 292 within the population were capable of expressing appropriate *phenotypes*, confirming 293 our first prediction (figures 2 and 3). Plasticity allowed the persistence of populations 294 even at low rates of *mean-mutational-change* and high environmental fluctuations, 295 unless *plasticity-costs* were high (0.7 and above; figure 3*a*). 296 The maintenance of an average greater population size and alleviation of

297 bottlenecks also contributed to increased genetic variation in the *plastic* populations

298 (figure 2 insets). Moreover, because large *plasticity-ranges* allowed *genotypes* that

299 would otherwise have had a poorly fitted *phenotype* to improve their *match*, the effect 300 of selection was buffered and higher genotypic diversity within populations was 301 retained in plastic populations at all times, confirming our second prediction. The strong 302 genetic response to selection of *non-plastics*, however, resulted in a better *match* 303 between average *genotype* and the *environment* for *non-plastic* than for *plastic* 304 genotypes (figure 2). Consequently, in fluctuating environments plasticity allowed the 305 phenotype to closely match the environment while slowing down the genotypic response 306 to selection (figure 2). At low *plasticity-costs*, the average genotypic value was 307 maintained around the average value of the environmental conditions experienced 308 throughout the simulations while at the same time retaining large genotypic variance 309 (figure 2b,c). In consequence, low plasticity-costs allowed increased plasticity to evolve 310 (figure 2b,c), leading to a higher genotype variance (figure 2 b,c insets) and thus 311 increasing the chances that appropriate genetic variants of fixed phenotypes arose by

312 mutation.

313 Environmental stabilisation and genetic assimilation

314 To test the prediction that costs of plasticity result in loss of plasticity upon environment 315 stabilisation, we simulated a fast environmental transition from *environment* = 0.5 to 1, 316 followed by *environment* stabilisation at 1, such as it would occur for instance due to 317 human activity or if a population was to enter a distinct ecological region (figure 4). As 318 in previous analyses (figures 2 and 3) our model exploration showed that adaptation to 319 the novel environment in the non-plastic population depended on mean-mutational-320 change relative to environmental change (results not shown). Also, if the environment 321 changed too abruptly given their mean-mutational-change, the *non-plastic* population 322 failed to adapt and went extinct. Plastic genotypes, however, managed to persist even 323 with a low *mean-mutational-change* and despite rapid transitions to the novel

324 environment. It was possible because their plasticity-range allowed them to manifest 325 phenotypes that better matched the environment at any given time. As shown in figure 4a, plasticity-range was strongly positively selected during the abrupt environmental 326 327 change and only the most plastic genotypes survived the sharp environmental transition, 328 because only very plastic genotypes were capable of producing extreme phenotypes. 329 Nevertheless, plastic *genotypes* lagged substantially behind their *phenotype* (figure 4b). 330 In other words, plasticity bought time for adaptive fixed (i.e. non *environment* sensitive) 331 genetic changes to occur because individuals expressed the appropriate *phenotype* soon but it often still took the genotype many generations to match the environment (figure 332 333 4b). When costs of plasticity were high and the new *environment* remained stable, 334 plasticity quickly decreased to background levels maintained by mutation, resulting in 335 genetic assimilation of the environmentally-induced phenotypes (figure 4a,b).

336

337 4. DISCUSSION

338 With this simple heuristic model we integrated adaptive plasticity into an explicit 339 population genetic framework, and examined some fundamental consequences of 340 plasticity in adaptive evolution. We found that fluctuating or rapid directional 341 environmental change strongly selected for plastic genotypes. This result is in 342 accordance with previous modelling approaches [22, 41, 42], especially when 343 environmental fluctuations are modelled to act after development but before selection 344 [43]. In our model, increased plasticity allowed genotypes to produce phenotypes better 345 matching the changing environmental conditions at each generation, hence showing a 346 high potential for rapid adaptation to new environments. This relationship between 347 plasticity and adaptive potential to novel environments has been suggested in some

cases, as in invasive plant species having greater plasticity than non-invasive ones [44];
plasticity mediating rapid adaptation to introduced predators in zooplanktonic species
[45]; or adaptations to climate change in birds [46].

Plasticity led to faster phenotypic modifications of whole populations because adaptive phenotypes were induced concurrently by environmental cues available to all individuals, instead of requiring the time for beneficial mutations to spread throughout the population by differential survival and reproduction [7]. This allowed populations composed of plastic genotypes to suffer fewer and lesser demographic bottlenecks despite steep fluctuations in the environment (figures 2 and 3).

357 An important result emerging from this model is that adaptive plasticity 358 contributes to the maintenance of genetic variation within population (figure 2 insets) in 359 two ways. First, plastic populations had higher genetic variation because plasticity 360 shielded a broader range of genotypes from purifying selection by allowing them to 361 express well-matched phenotypes. Second, plasticity reduced the effect of genetic drift 362 as a consequence of maintaining greater population sizes (i.e. by reducing population 363 bottlenecks). This result is supported by a very different modelling approach that has 364 also recently proposed that plasticity tends to lead to populations with greater 365 mutational and standing genetic variance [47].

It has often been debated whether plasticity fosters evolution by facilitating adaptation to novel environments or rather impede divergence by shielding genetic variation from divergent selection [17, 48, 49]. We show that plasticity allows phenotypically cryptic (or unexpressed) genetic variation to build up within populations by conferring similar fitness to distinct genotypic variants (see also [18, 50]). Adaptive plasticity also allows otherwise imperilled populations to persist until appropriate genetic variants appear (figures 2 and 4). Moreover, the accumulated genetic variation

can be rapidly released and manifested in the face of further environmental or
mutational changes, enabling rapid adaptive divergences [6, 17, 51, 52]. Our study
suggests that plasticity facilitates adaptation to novel environments by allowing a
synchronic phenotypic shift in response to the environment, while at the same time
maintaining genetic variation that would otherwise be selected out (figure 2 insets),
even though phenotypic plasticity slows down the response to selection (figures 2 and
4b).

380 Overall, shielding of genetic variation by plasticity may only be a transient effect 381 of an otherwise rapid process of adaptation to divergent environments by genetic 382 accommodation, as we found that plastic genotypes always showed a greater adaptive 383 potential to a changing environment (figures 2-4). Congruently, there are many cases of 384 rapidly diversifying groups of species where genetic accommodation of plasticity is 385 likely to have been the main driver for divergence [53], as in sticklebacks [54, 55], 386 anole lizards [56], or arctic charrs [57]. Rapid adaptive transitions between 387 environments are more easily achieved by plastic than non-plastic genotypes (figures 3 388 and 4), and we show that genetic assimilation of induced phenotypes and the associated 389 loss of plasticity will occur if costs of plasticity are high and the environment stabilises 390 (figure 4).

391 Plasticity costs have been elusive and difficult to measure empirically [58-60],
392 but there is evidence for plasticity costs from plants to invertebrates and vertebrates [6163]. Moreover, patterns of evolution of plasticity are often congruent with theoretical
and expectations of the consequences of costs of plasticity, namely reduced plasticity under
stable environmental conditions. American spadefoot toads, for instance, have evolved a
canalised accelerated larval development with respect to the slow but plastic
397 development ancestral to the group as a result of their adaptation to ephemeral desert

398 ponds [31]. Accelerated development has become nearly genetically assimilated, and 399 plasticity has been lost to a great extent in desert spadefoot toads so they are no longer 400 capable of long larval periods [31, 64]. Such translation of ancestral environmentally 401 induced changes in development within populations into adaptive constitutive 402 divergences among taxa is a clear path connecting micro- and macroevolution [2, 7, 31]. 403 Because environmental variation is the rule in nature [65] and it often selects for 404 adaptive plasticity [16, 18, 66], the evolutionary paradigm needs to be extended to 405 include environmentally-dependent regulation of gene expression as a heritable source of phenotypic variation, whether genetic or epigenetic [9, 35, 67-69]. Whether the 406 407 incorporation of adaptive plasticity constitutes an extension of the paradigm emerged 408 from the Modern Synthesis or a new paradigm, may ultimately be better evaluated 409 retrospectively. To some extent, adaptive plasticity simply extends and strengthens the 410 current paradigm, as it improves our understanding of the maintenance of genetic 411 variation in populations, facilitates rapid adaptive shifts between adaptive peaks, and 412 helps explaining the adaptive radiations and recurrent parallel speciation. However, at 413 the same time, accounting for adaptive plasticity expands the Modern Synthesis 414 paradigm in several meaningful aspects that may warrant a new paradigm. Our model 415 illustrates these aspects in a fairly simple and intiutive way. First, during organismal 416 development the environment acts as a phenotypic inducer in addition to its traditional 417 role as a mere selective sieve. This is important because environmental induction may 418 act simultaneously on most genotypes in a population inducing synchronous phenotypic 419 shifts in the direction of the new local adaptive optimum. Second, plasticity increases 420 the match of the phenotype to the environment, reducing bottlenecks and hence 421 increasing population viability. Lastly, plasticity contributes to the maintenance of 422 genetic variation within populations both by shielding many genetic variants from

selection and by reducing genetic drift, and can become quickly accomodated betweenlineages evolving in divergent environments.

425 In this line of thought, our model shows the high relevance of plasticity to 426 evolution and population ecology, while at the same time it shows that incoporating 427 plasticity is conceptually as simple as acknowledging the fact that genotypes may have 428 the potential to use environmental information to express better fit phenotypes. Other 429 central tenets of mainstream evolutionary thought (i.e. random mutation and selection of 430 phenotypes according to environmenal conditions) evidently remain unchanged. The 431 simple addition of environmentally-sensitive adaptive gene regulation, however, 432 provides a demonstrated mechanism for swift adaptation to rapidly changing 433 environments that may have often lead to lineage diversification and evolutionary 434 innovations.

435

436 **5. ACKNOWLEDGMENTS**

437 We thank P. Edelaar, S. Sultan, J. Moyá and C. M. Herrera, and two anonymous

438 reviewers for their comments and suggeted improvements to the model. This research

439 was supported by the Ramón y Cajal Program of the Spanish Ministerio de Ciencia e

440 Innovación (MICINN) to IGM (RYC-2008-03519) and to RJ (RYC-2009-03967).

441

442 **6. LITERATURE CITED**

443 1. Futuyma D.J. 2009 *Evolution*. Sunderland, MA, Sinauer Associates.

444 2. Moczek A.P. 2007 Developmental capacitance, genetic accommodation, and

445 adaptive evolution. *Evol. Devel.* **9**, 299-305.

- 446 3. Pigliucci M., Müller G.B. 2010 Elements of an extended evolutionary synthesis.
- 447 In Evolution The extended synthesis (eds. Pigliucci M., Müller G.B.), pp. 3-17.

448 Cambridge, MA, MIT Press.

- 449 4. Gillespie J.H., Turelli M. 1989 Genotype-environment interactions and the
 450 maintenance of polygenic variation. *Genetics* 121, 129-138.
- 451 5. Byers D.L. 2005 Evolution in heterogeneous environments and the potential of
 452 maintenance of genetic variation in traits of adaptive significance. *Genetica* 123, 107-

453 124.

- 454 6. Barrett R.D.H., Schluter D. 2008 Adaptation from standing genetic variation.
- 455 *Trends Ecol. Evol.* **23**, 38-44.
- 456 7. West-Eberhard M.J. 2003 Developmental plasticity and evolution. Oxford,
- 457 Oxford University Press; 794 p.
- 458 8. Pigliucci M. 2001 Phenotypic plasticity: beyond nature and nurture. Baltimore,
- 459 Johns Hopkins University Press; 384 p.
- 460 9. Cabej N.R. 2012 *Epigenetic principles of evolution*. London, Elsevier.
- 461 10. Aubin-Horth N., Renn S.C.P. 2009 Genomic reaction norms: using integrative
- 462 biology to understand molecular mechanisms of phenotypic plasticity. *Mol. Ecol.* 18,

463 3763-3780.

- 464 11. Sultan S.E., Spencer H.G. 2002 Metapopulation structure favors plasticity over
- 465 local adaptation. *Amer. Nat.* **160**, 271-283.
- 466 12. Scheiner S.M., Barfield M., Holt R.D. 2012 The genetics of phenotypic
- 467 plasticity. XI. Joint evolution of plasticity and dispersal rate. *Ecol. Evol.* **2**, 2027-2039.
- 468 13. Hallsson L.R., Björklund M. 2012 Selection in a fluctuating environment leads
- to decreased genetic variation and facilitates the evolution of phenotypic plasticity. J.
- 470 Evol. Biol. 25, 1275-1290.

- 471 14. Roff D.A. 1997 *Evolutionary quantitative genetics*. New York, NY, Chapman &
 472 Hall.
- 473 15. Losos J.B., Arnold S.J., Bejerano G., Brodie E.D., III, Hibbett D., Hoekstra
- 474 H.E., Mindell D.P., Monteiro A., Moritz C., Orr H.A., et al. 2013 Evolutionary Biology
- 475 for the 21st Century. *PLoS Biol* **11**, e1001466. (doi:10.1371/journal.pbio.1001466).
- 476 16. West-Eberhard M.J. 2002 Development and selection in adaptive evolution.
- 477 *Trends Ecol. Evol.* **17**, 65-65.
- 478 17. Pfennig D.W., Wund M.A., Snell-Rood E.C., Cruickshank T., Schlichting C.D.,
- 479 Moczek A.P. 2010 Phenotypic plasticity's impacts on diversification and speciation.
- 480 *Trends Ecol. Evol.* **25**, 459-467.
- 481 18. Moczek A.P., Sultan S., Foster S., Ledon-Rettig C., Dworkin I., Nijhout H.F.,
- 482 Abouheif E., Pfennig D.W. 2011 The role of developmental plasticity in evolutionary
- 483 innovation. Proc. Roy. Soc. B 278, 2705-2713.
- 484 19. Ancel L.W. 1999 A Quantitative Model of the Simpson-Baldwin Effect. J.
- 485 *Theor. Biol.* **196**, 197-209.
- 486 20. Berrigan D., Scheiner S.M. 2004 Modeling the evolution of phenotypic
- 487 plasticity. In *Phenotypic plasticity* (eds. DeWitt T.J., Scheiner S.M.), pp. 82-97. Oxford,
- 488 Oxford University Press.
- 489 21. de Jong G. 2005 Evolution of phenotypic plasticity: patterns of plasticity and the
 490 emergence of ecotypes. *New Phytol.* 166, 101-118.
- 491 22. Lande R. 2009 Adaptation to an extraordinary environment by evolution of
- 492 phenotypic plasticity and genetic assimilation. J. Evol. Biol. 22, 1435-1446.
- 493 23. DeWitt T.J., Sih A., Wilson D.S. 1998 Costs and limits of phenotypic plasticity.
- 494 *Trends Ecol. Evol.* **13**, 77-81.

- 495 24. Auld J.R., Agrawal A.A., Relyea R.A. 2009 Re-evaluating the costs and limits
- 496 of adaptive phenotypic plasticity. *Proc. Roy. Soc. B* 277, 503-511.
- 497 25. Pigliucci M., Murren C.J. 2003 Genetic assimilation and a possible evolutionary
- 498 paradox: Can macroevolution sometimes be so fast as to pass us by? *Evolution* **57**,
- 499 1455-1464.
- 500 26. Crispo E. 2007 The Baldwin effect and genetic assimilation: Revisiting two
- 501 mechanisms of evolutionary change mediated by phenotypic plasticity. *Evolution* 61,
 502 2469-2479.
- 503 27. Foster S.A., Wund M.A. 2011 Epigenetic contributions to adaptive radiations:
- 504 insights from threespine sticklebacks. In Epigenetics Linking genotype and phenotype in
- 505 *development and evolution* (eds. Hallgrimsson B., Hall B.K.), pp. 317-336. Berkeley,
- 506 CA, University of California Press.
- 507 28. Waddington C.H. 1952 Selection of the genetic basis for an acquired character.
 508 *Nature* 169, 278.
- 509 29. Waddington C.H. 1959 Canalization of development and the inheritance of
- 510 acquired characters. *Nature* **183**, 1654-1655.
- 511 30. Suzuki Y., Nijhout H.F. 2006 Evolution of a polyphenism by genetic
- 512 accommodation. *Science* **311**, 650-652.
- 513 31. Gomez-Mestre I., Buchholz D.R. 2006 Developmental plasticity mirrors
- 514 differences among taxa in spadefoot toads linking plasticity and diversity. *Proc. Nat.*
- 515 Acad. Sci., USA **103**, 19021-19026.
- 516 32. Ledon-Rettig C.C., Pfennig D.W., Nascone-Yoder N. 2008 Ancestral variation
- 517 and the potential for genetic accommodation in larval amphibians: implications for the
- 518 evolution of novel feeding strategies. *Evol. Devel.* **10**, 316-325.
- 519 33. Simpson G.G. 1953 The Baldwin Effect. *Evolution* 7, 110-117.

- 520 34. Orr H.A. 1999 Evolutionary biology: An evolutionary dead end? *Science* 285,
 521 343-344.
- 522 35. Jablonka E., Lamb M.J. 2010 Transgenerational epigenetic inheritance In
- 523 Evolution The extended synthesis (eds. Pigliucci M., Müller G.B.), pp. 137-174.
- 524 Cambridge, MA, The MIT Press.
- 525 36. Holeski L.M., Jander G., Agrawal A.A. 2012 Transgenerational defense
- 526 induction and epigenetic inheritance in plants. *Trends Ecol. Evol.* 27, 618-626.
- 527 37. Grimm V., Railsback S.F. 2005 Individual-based modeling and ecology.
- 528 Princeton, USA, Princeton University Press.
- 529 38. Grimm V., Berger U., Bastiansen F., Eliassen S., Ginot V., Giske J., Goss-
- 530 Custard J., Grand T., Heinz S.K., Huse G., et al. 2006 A standard protocol for
- 531 describing individual-based and agent-based models. *Ecol. Model.* **198**, 115-126.
- 532 39. Grimm V., Berger U., DeAngelis D.L., Polhill J.G., Giske J., Railsback S.F.
- 533 2010 The ODD protocol: A review and first update. *Ecol. Model.* 221, 2760-2768.
- 534 40. Wilensky U. 1999 (Northwestern University, Evanston, IL.
- 535 <u>http://ccl.northwestern.edu/netlogo</u>, Center for Connected Learning and Computer-
- 536 Based Modeling.
- 537 41. Scheiner S.M. 1998 The genetics of phenotypic plasticity. VII. Evolution in a
- 538 spatially-structured environment. J. Evol. Biol. 11, 303-320.
- 539 42. Chevin L.M., Lande R. 2011 Adaptation to marginal habitats by evolution of
- 540 increased phenotypic plasticity. J. Evol. Biol. 24, 1462-1476.
- 541 43. Scheiner S.M., Holt R.D. 2012 The genetics of phenotypic plasticity. X.
- 542 Variation versus uncertainty. *Ecol. Evol.* 2, 751-767.

- 543 44. Davidson A.M., Jennions M., Nicotra A.B. 2011 Do invasive species show
- higher phenotypic plasticity than native species and, if so, is it adaptive? A meta-analysis. *Ecol. Lett.* 14, 419-431.
- 546 45. Scoville A.G., Pfrender M.E. 2010 Phenotypic plasticity facilitates recurrent
- 547 rapid adaptation to introduced predators. *Proc. Nat. Acad. Sci.* **107**, 4260-4263.
- 548 46. Charmantier A., McCleery R.H., Cole L.R., Perrins C., Kruuk L.E.B., Sheldon
- 549 B.C. 2008 Adaptive phenotypic plasticity in response to climate change in a wild bird
- 550 population. *Science* **320**, 800-803.
- 551 47. Draghi J.A., Whitlock M.C. 2012 Phenotypic plasticity facilitates mutational
- variance, genetic variance, and evolvability along the major axis of environmental
- 553 variation. *Evolution* **66**, 2891-2902.
- 48. Price T.D., Qvarnstrom A., Irwin D.E. 2003 The role of phenotypic plasticity in
 driving genetic evolution. *Proc. Roy. Soc. B* 270, 1433-1440.
- 556 49. Ghalambor C.K., McKay J.K., Carroll S.P., Reznick D.N. 2007 Adaptive versus
- 557 non-adaptive phenotypic plasticity and the potential for contemporary adaptation in new
- 558 environments. *Funct. Ecol.* **21**, 394-407.
- 559 50. Flatt T. 2005 The evolutionary genetics of canalization. *Quart. Rev. Biol.* 80,
 560 287-316.
- 561 51. Gibson G., Dworkin I. 2004 Uncovering cryptic genetic variation. Nat. Rev.
- 562 *Genet.* **5**, 681-690.
- 563 52. Le Rouzic A., Carlborg ñ. 2008 Evolutionary potential of hidden genetic
- variation. *Trends Ecol. Evol.* **23**, 33-37.
- 565 53. West-Eberhard M.J. 1989 Phenotypic plasticity and the origins of diversity. Ann.
- 566 *Rev. Ecol. Syst.* **20**, 249-278.

- 567 54. Wund M., Baker J.A., Clancy B., Golub J.L., Foster S.A. 2008 A test of the
- 568 'Flexible Stem' model of evolution: ancestral plasticity, genetic accommodation, and
- 569 morphological divergence in the threespine stickleback radiation. *Amer. Nat.* **172**, 449-
- 570 462.
- 571 55. Wund M.A., Valena S., Wood S., Baker J.A. 2012 Ancestral plasticity and
- allometry in threespine stickleback reveal phenotypes associated with derived,
- 573 freshwater ecotypes. *Biol. J. Linn. Soc.* **105**, 573-583.
- 574 56. Losos J.B., Creer D.A., Glossip D., Goellner R., Hampton A., Roberts G.,
- 575 Haskell N., Taylor P., Ettling J. 2000 Evolutionary implications of phenotypic plasticity
- 576 in the hindlimb of the lizard *Anolis sagrei*. *Evolution* **54**, 301-305.
- 577 57. Adams C.E., Huntingford F.A. 2004 Incipient speciation driven by phenotypic
- 578 plasticity? Evidence from sympatric populations of Arctic charr. *Biol. J. Linn. Soc.* 81,
- 579 611-618.
- 580 58. Scheiner S.M., Berrigan D. 1998 The genetics of phenotypic plasticity. VIII. The
 581 cost of plasticity in *Daphnia pulex*. *Evolution* 52, 368-378.
- 582 59. Johansson F. 2002 Reaction norms and production costs of predator-induced
- 583 morphological defences in a larval dragonfly (Leucorrhinia dubia : Odonata). Can. J.
- 584 Zool. **80**, 944-950.
- 585 60. Van Buskirk J., Steiner U.K. 2009 The fitness costs of developmental
- 586 canalization and plasticity. J. Evol. Biol. 22, 852-860.
- 587 61. DeWitt T.J. 1998 Costs and limits of phenotypic plasticity: Tests with predator-
- induced morphology and life history in a freshwater snail. J. Evol. Biol. 11, 465-480.
- 589 62. Agrawal A.A., Conner J.K., Johnson M.T.J., Wallsgrove R. 2002 Ecological
- 590 genetics of an induced plant defense against herbivores: additive genetic variance and
- 591 costs of phenotypic plasticity. *Evolution* **56**, 2206-2213.

- 592 63. Relyea R.A. 2002 Costs of phenotypic plasticity. Amer. Nat. 159, 272-282.
- 593 64. Kulkarni S.S., Gomez-Mestre I., Moskalik C.L., Storz B.L., Buchholz D.R. 2011
- 594 Evolutionary reduction of developmental plasticity in desert spadefoot toads. J. Evol.
- *Biol.* **24**, 2445-2455.
- 596 65. Ricklefs R.E., Schluter D. 1993 Species diversity in ecological communities:
- 597 *historical and geographical perspectives*. Chicago, University of Chicago Press.
- 598 66. Scheiner S.M. 1993 Genetics and evolution of phenotypic plasticity. *Ann. Rev.*599 *Ecol. Syst.* 24, 35-68.
- 600 67. Bossdorf O., Richards C.L., Pigliucci M. 2008 Epigenetics for ecologists. *Ecol.*
- 601 *Lett.* **11**, 106-115.
- 602 68. Carroll S.B. 2008 Evo-devo and an expanding evolutionary synthesis: A genetic
- theory of morphological evolution. *Cell* **134**, 25-36.
- 604 69. Richards C.L., Bossdorf O., Pigliucci M. 2010 What role does heritable
- 605 epigenetic variation play in phenotypic evolution? *BioScience* **60**, 232-237.
- 606
- 607
- 608
- 609

Table 1. Variables and parametrisation. All variables and parameters can take continuous values.

	Initialization	Constraints during	Description
		simulations	
- Parameters			
Std-Dev-environment-change	(0.04-1)	Initialization value	Determines the degree of environmental stochasticity
			$environment_{t+1} = environment_t + N~(0,Std-Dev-Environment-Change)$
plasticity-costs	(0-1)	Initialization value	Determines whether plasticity carries a load reducing odds of survival and
			reproducing
mean-mutational-change	(0-0.002)	Initialization value	Determines both the probability of occurrence of genetic changes and
			their effect size on the <i>phenotype</i>
- Emergent values			
environment	0.5	[0,1]	Expresses the environmental conditions on a single dimension, the same
			one used to describe the <i>phenotype</i> , the <i>genotype</i> and the <i>plasticity-range</i>
- non-plastics and plastics			
phenotype	0	-	Phenotypic value expressed in the same dimension as the environment

genotype	0	-	In the absence of plasticity, the <i>phenotype</i> = genotype
match	NA	-	Absolute difference between the phenotypic value and the environmental
			value; the phenotype is optimised if match=1
			1 - environment - phenotype
- plastics only			
plasticity-range	0	-	The maximum phenotypic adjustment that a <i>genotype</i> is capable to
			increase match
used-plasticity	NA	$0 \leq used$ -plasticity	Amount of the <i>plasticity-range</i> that is actually used by an individual
		\leq plasticity range	during development

Figure legends

Figure 1. Schematic representation of the individual-based model comparing adaptive evolution in populations composed of *plastic* or *non-plastic* genotypes. They are all clonal organisms with no recombination so that *non-plastic* genotypes map directly into phenotypes and their odds of surviving and reproducing depend on the *match* with the *environment*. In contrast, *plastic* genotypes can respond to the *environment* modifying their *phenotype* to reduce the mismatch to the extent that their plasticity-range allows. In both cases the *environment* acts as a selective factor, but for *plastic genotypes* it is also a phenotypic inducer.

Figure 2. Examples of adaptive evolution of *plastic* and *non-plastic* populations under medium-low environmental fluctuations (*Std-Dev-environment-change* = 0.1) and different scenarios of *mean-mutational-change* and *plasticity-costs*. (*a*) At high *mean-mutational-change* and high *plasticity-costs*, *plastics* performed similar to *non-plastics*. Here, a high *mean-mutational-change* allowed both populations to closely track the *environment*. *Plasticity-range* was reduced compared to scenarios with lower costs but maintained due to environmental fluctuations. (*b*) Under high *mean-mutational-change* but with low *plasticity-costs*, plasticity allowed a close phenotypic *match* to the *environment* and the persistence of the *plastic* population, but often *non-plastics* went extinct as shown in this example. (*c*) Under low *mean-mutational-change* and low *plasticity-costs*, *plastic genotypes* produced *phenotypes* that closely matched the *environment* while their genotypic values were intermediate across environmental fluctuations, and plasticity increased. *Non-plastic* genotypes could not adapt fast enough and quickly went extinct. At any given time and in all scenarios, genotypic variation

was higher in the *plastic* population than in the *non-plastic* one. This is shown in inset boxplots in each panel, where blue boxes depict genetic variation of the *non-plastic* population and orange boxes that of the *plastic* population, sampled every 25 generations.

Figure 3. (*a*) Results for population size for populations composed of either *plastic* or *non-plastic* genotypes from simulations sweeping over all parameter combinations of environmental stochasticity (*Std-Dev-environment-change*), mutation rate (*mean-mutational-change*), and *plasticity-costs*. Populations composed of *plastic* genotypes persisted over a much broader range of environmental stochasticity than populations of *non-plastic* genotypes, unless *plasticity-costs* were high, in which case they performed worse than *non-plastic* genotypes. (*b*) Examples of population dynamics for *plastic* and *non-plastic* populations at different levels of environmental stochasticity and *mean-mutational-change* = 0.04 and *plasticity-costs* = 0.6; panel numbers relate (*a*) to (*b*). (*c*) Example of clonal lineages trajectories (each line is a lineage) according to *genotype* and (for plastics) *plasticity-range* (lighter green colour depicts higher *plasticity-range*). Note that only very plastic lineages survived the strongest population bottleneck (as shown in corresponding (*b*) panel).

Figure 4. Example of model run for a scenario of directional environmental change, where environment changed abruptly from 0 to 1 and then stabilised at 1 with *mean-mutational-change* = 0.005 and *plasticity-costs* = 0.7. (*a*) Shows for *plastic* individuals their position in the *genetoype* vs. *plasticity-range* space. The arrow indicate the pass of time (in generations), beginning with all individuals with *genotype* = 0.5 and *plasticity-range* = 0 (initialization conditions) and ending at the end of the simulation with individuals with *genotypes* close to 1 and reduced *plasticity-range*. (*b*) Same as in figure

2. It is shown how plasticity increased temporarily under selection and the plasticpopulation expressed well-matched phenotypes, allowing the population to persist overenough generations to allow genotypes to slowly evolve towards the new optimum.Once the environment stabilises, plasticity is rapidly reduced due to costs of plasticity,causing genetic assimilation.