

STOCHASTIC ECO-EVOLUTIONARY DYNAMICS OF MULTIVARIATE TRAITS

A Framework for Modeling Population Processes Illustrated by the Study of Drifting G-Matrices

Bob Week^{1,2}

¹Zoological Institute, Kiel University, Kiel, Germany and ²Max-Planck Institute for Evolutionary Biology, Plön, Germany

Abstract

I derive a novel stochastic equation for the evolution of the additive genetic variance-covariance matrix \mathbf{G} in response to mutation, selection, drift, and fluctuating population size. Common wisdom holds that the effect of drift on \mathbf{G} is simply to reduce each of its entries by a common proportional amount while preserving its orientation. In contrast, I find that drift causes significant and directional shifts in the orientation of \mathbf{G} by driving genetic correlations to their extremes. Biologically, this is a consequence of linkage build-up introduced by drift. I compare these theoretical results to empirical observations based on experiments conducted by Phillips et al., (2001). Additionally, to derive the model of \mathbf{G} -matrix evolution, I developed a novel synthetic framework for modelling ecological and evolutionary dynamics of populations carrying multivariate traits. This framework is optimized for deriving new models across a wide range of topics in population biology. Foundations of the framework are formalized by the theory of measure-valued processes, but application of the framework only requires multivariate calculus, and heuristics are presented in the main text for making additional calculations involving stochastic processes. Collectively, this work establishes a powerful framework enabling efficient formal analysis of integrated population processes across evolution and ecology, and its potential for making new discoveries is illustrated by novel findings on fundamental aspects of \mathbf{G} -matrix evolution.

Key words: Evolutionary Ecology; Quantitative Genetics; Drift; \mathbf{G} -Matrices; Martingale Problems

1. Introduction

A fundamental principle of evolutionary biology is that random genetic drift erodes heritable variation at a rate inversely proportional to effective population size. At the level of allele frequencies, models for the distribution of genetic variation responding to drift are well-known (Kimura, 1964, 1968; Ewens, 2004; Hill and Robertson, 1966). In contrast, at the level of quantitative characters, models of additive genetic variation responding to drift focus on the average outcome using deterministic models (Latter, 1970; Bulmer, 1972; Lande, 1976, 1980; Chakraborty and Nei, 1982; Turelli and Barton, 1994; Lynch and Hill, 1986; Bürger, 2000; Barton and Turelli, 2004; Débarre and Otto, 2016; Walsh and Lynch, 2018). Empirical work has supported theoretical predictions for the average response of additive genetic variation to drift (Phillips et al., 2001; McGuigan et al., 2005; Whitlock et al., 2002), but has also emphasized the need to predict the distribution of outcomes (Phillips et al., 2001; Whitlock, 1995) especially for multivariate traits and genetic covariances (Phillips and McGuigan, 2006; Mallard et al., 2023a).

In the setting of multivariate traits, a common summary statistic for genetic architecture is the \mathbf{G} -matrix. This matrix has the additive genetic variance of each trait on the associated diagonal entry. Off-diagonal entries quantify genetic covariances between traits which may be maintained by pleiotropic loci and linkage between loci affecting different traits. The standard view is that drift produces a proportional decrease in \mathbf{G} (Phillips and McGuigan, 2006; Cano et al., 2004; McGuigan, 2006; Chapuis et al., 2008; Dugand et al., 2021; Mallard et al., 2023a), and thus independent populations that have diverged due to drift should have proportional \mathbf{G} -matrices (Roff, 2000; Steppan et al., 2002; Aguirre et al., 2013). However, this insight rests on results from a deterministic model for the response of \mathbf{G} -matrices to drift (Lande, 1979) and a model that assumes recombination happens sufficiently fast to break-up linkage produced by selection (Lande, 1980). Hence, there is a need to develop theoretical predictions for the stochastic evolution of \mathbf{G} -matrices driven solely by drift, and especially for the effect of drift on genetic covariances (Mallard et al., 2023a,b).

The reason for this gap in evolutionary theory stems, in part, from the lack of formal approaches to place stochastic

models of \mathbf{G} -matrix evolution on a concrete mathematical foundation. In this paper, I aim to fill this gap by leveraging the powerful theory of measure-valued processes, while also keeping the presentation as accessible as possible. Furthermore, taking this approach to formally derive \mathbf{G} -matrix dynamics revealed a much broader framework for modelling a wide-range of population processes. In particular, given the growing appreciation for the interplay between ecological and evolutionary processes (Reznick, 2015; Hendry, 2017; Kuosmanen et al., 2022), this framework makes an important contribution by enabling the formal interfacing of \mathbf{G} -matrix evolution with models of eco-evolutionary feedbacks (e.g., Patel et al., 2018). I therefore focus this paper on presenting the framework in its full generality, and return to the study of \mathbf{G} -matrix evolution as an example to illustrate its utility.

In its full generality, this framework offers tools to model the integrated ecological and evolutionary dynamics of populations with multivariate traits that respond to mutation, selection (including frequency and abundance dependence), demographic stochasticity, and consequential random genetic drift. In particular, this framework can be used to obtain generalizations of many classical models in evolution and ecology, such as Lotka-Volterra dynamics (Cattiaux and Méléard, 2009; Akjouj et al., 2024), coevolution (Gilman et al., 2012; Débarre et al., 2014), and evolutionary rescue (Klausmeier et al., 2020; Xu et al., 2023). Further details on how to apply this framework to generalize known models are given in the discussion section.

As in classical quantitative genetics, the framework here assumes a linear genotype-phenotype map with fixed mutational covariance. Under this modelling choice, development enters implicitly through the genotype-phenotype map rather than via explicit mechanistic developmental feedbacks (see Milocco and Salazar-Ciudad, 2022; González-Forero, 2022, 2024a,b). This paper therefore focuses on mutation-selection-drift dynamics of multivariate traits, not on the evolution of developmental architecture.

To apply this framework, the most important biological details relevant to a modeler are the mechanisms mediating fitness. In contrast, details involving the genetic architecture of traits are abstracted in a way that captures basic biological principles while optimizing analytical tractability. For instance, asexual reproduction is assumed and mutation is modeled following approaches similar to Kimura (1965) and Débarre and Otto (2016) by assuming offspring traits are distributed around their parental traits. This approach to modelling inheritance is similar to the infinitesimal model in which traits are determined by many loci of small effect, which implies offspring are normally distributed around their parental values (Barton et al., 2017). However, an important distinction is that the infinitesimal model restricts the response of allele frequencies to selection such that changes in genetic variance are only temporary and due to build-up of genetic linkage (Bulmer, 1971). Instead, I begin by assuming offspring traits are normally distributed around their parental traits (referred to as the Gaussian descendants model by Turelli, 2017), in which case the response of population variance to selection can be permanent. Abstract approaches similar to this have been successful for obtaining valuable analytical insights into genetic variation maintained by mutation-selection balance (Kimura, 1965; Lande, 1975; Turelli, 1984, 1986) and by mutation-drift balance (Lande, 1976, 1979; Barton, 1989; Débarre and Otto, 2016). Hence, by optimizing a trade-off between genetic detail and analytical tractability, this framework provides an

accessible approach for obtaining and communicating a wide array of novel theoretical insights.

To establish this framework, I build on the work of Week et al. (2021), which presented a stochastic differential equation framework focused on modelling the simultaneous dynamics of abundances, 1-dimensional mean traits, and 1-dimensional trait variances responding to mutation, selection, demographic stochasticity, and random genetic drift. This 1-dimensional framework was based on the development of heuristics (i.e., methods for performing exact calculations without formal justification) for working with stochastic partial differential equations (spde). However, for multivariate traits, the spde approach breaks down (Dawson, 1993; Etheridge, 2000; Perkins, 2002). To overcome this challenge and establish a rigorous analytical framework for deriving population processes, I take an approach based on so-called *martingale problems* (Ethier and Kurtz, 1986; Dawson, 1993; Stroock and Varadhan, 1997; Rogers and Williams, 2000). Mathematical aspects of this approach are provided in Supplementary Material, Section 4. In the main text, I focus on the resulting dynamical equations and heuristics for performing calculations with minimal technical background.

1.1. Overview

I begin by outlining the derivation of the deterministic version of the framework without making any assumptions on the shape of trait distributions, which is summarized by a system of ordinary differential equations. This leads to expressions of selection in terms of covariances with fitness, which I refer to collectively as the Deterministic Covariance version (or *DC* for short). By assuming traits follow multivariate normal distributions, covariances with fitness are replaced by multivariate gradients of fitness with respect to mean traits and trait variances, and I refer to the resulting system of differential equations as the Deterministic Gradient version (or *DG* for short). To simplify presentation of the deterministic version of the framework, I assume traits are perfectly heritable. However, because this work is motivated by understanding the consequences of drift for \mathbf{G} -matrix evolution, I briefly describe an approach to model imperfect heritability after introducing *DC* and *DG*. This model of imperfect heritability is adopted while introducing the stochastic extensions of the framework.

The stochastic extensions of the framework include the effects of demographic stochasticity (i.e., random reproductive output) and random genetic drift (which occurs here as a consequence of demographic stochasticity). I introduce two stochastic extensions. Both build on *DG* by assuming multivariate normal trait distributions and by expressing selection in terms of fitness gradients. The first form expresses dynamics in terms of Brownian motions as drivers of stochasticity (referred to as the Brownian Motion Gradient version, or *BG* for short), which is particularly useful for numerical analysis. The second form expresses dynamics in terms of a more general underlying martingale process (referred to as the Martingale Gradient version, or *MG* for short), and I use this form to introduce heuristics for deriving analytical models. To demonstrate the heuristics (summarized in Table 1), I derive a stochastic equation for the evolution of additive genetic correlations between trait values. To bring this paper full-circle, I then discuss how this exercise provides novel insights into the evolutionary response of \mathbf{G} -matrices to random genetic drift and compare these theoretical results with

167 observations obtained from experiments (Phillips et al., 2001;
 168 McGuigan et al., 2005; Whitlock et al., 2002).

169 The Supplementary Material provides details to support
 170 conclusions and arrive at expressions presented throughout
 171 the main text. Supplementary Material, Section 1 summarizes
 172 numerical implementations of the framework. Detailed
 173 derivation of the deterministic versions of the framework
 174 are given in Supplementary Material, Section 2. Section
 175 3 of the Supplement formally interfaces the continuous-
 176 time framework presented here with discrete time models of
 177 classical quantitative genetics. The mathematical foundation
 178 of this framework is communicated in Supplementary Material,
 179 Section 4. Using this foundation, Section 5 of the Supplement
 180 presents detailed calculations for the derivation of the
 181 stochastic versions of the framework presented in the main text.
 182 In Supplementary Material, Section 6, I derive the dynamics
 183 of genetic correlations responding to drift. An outline for one
 184 approach to derive this framework from a diffusion-limit is
 185 given in Section 7 of the Supplement. Finally, Supplementary
 186 Material, Section 8 formalizes a connection with Gillespie's
 187 work on the evolutionary consequences of offspring number
 188 variation (Gillespie, 1974, 1975, 1977)

189 2. The Framework

190 The framework tracks the dynamics of the population density
 191 across trait space for asexually reproducing populations. To
 192 model d -dimensional traits, I assume trait space is the entire
 193 Euclidean space \mathbb{R}^d . Given the d -dimensional trait $\mathbf{z} = (z_1, \dots, z_d)^\top$ (with \top denoting matrix transposition so \mathbf{z} is
 194 a column vector), I write $\nu(\mathbf{z})$ for the population density
 195 at \mathbf{z} . Then the total abundance of the population is given
 196 by $n = \int_{\mathbb{R}^d} \nu(\mathbf{z}) d\mathbf{z}$. The distribution of the trait \mathbf{z} is then
 197 $p(\mathbf{z}) = \nu(\mathbf{z})/n$, and this is also referred to as the relative
 198 abundance of \mathbf{z} . Using $p(\mathbf{z})$, the mean trait vector is given
 199 by $\bar{\mathbf{z}} = \int_{\mathbb{R}^d} \mathbf{z} p(\mathbf{z}) d\mathbf{z}$, and the trait covariance matrix is $\mathbf{P} =$
 200 $\int_{\mathbb{R}^d} (\mathbf{z} - \bar{\mathbf{z}})(\mathbf{z} - \bar{\mathbf{z}})^\top p(\mathbf{z}) d\mathbf{z}$, with P_{ij} being the covariance between
 201 z_i and z_j . The chosen notation facilitates comparison with
 202 classical multivariate quantitative genetic models (Lande, 1980;
 203 Lande and Arnold, 1983; Jones et al., 2003; Arnold et al., 2008).

204 In the following section (2.1) I describe how to obtain
 205 the deterministic version of the framework using multivariate
 206 calculus. I do this in two parts. The first part (*DC*)
 207 makes no assumption on the shape of $p(\mathbf{z})$. The second part
 208 (*DG*) assumes $p(\mathbf{z})$ is the density of a multivariate normal
 209 distribution. I continue to make this assumption in section 2.2
 210 where I introduce the stochastic extension of this framework.

212 2.1. Deterministic Dynamics

213 2.1.1. The Deterministic Covariance Version (*DC*)

214 To establish a flexible, but tractable framework to model
 215 the dynamics of n , $\bar{\mathbf{z}}$, and \mathbf{P} , I generalize the deterministic
 216 model used by Week et al., (2021), referred to there as the
 217 Deterministic Asexual Gaussian Allelic model (DAGA), to the
 218 multivariate setting. Specifically, DAGA focuses on dynamics
 219 due to just mutation and selection for asexually reproducing
 220 populations. Mutation is modeled as the variance M of a
 221 normal distribution determining offspring traits centered on
 222 their parental traits, and selection results from the covariance
 223 of fitness and phenotype.

224 In the multivariate trait setting, mutation is modeled as
 225 a $d \times d$ covariance matrix \mathbf{M} , with the off-diagonal entries
 226 contributing to trait covariances. Here, I assume \mathbf{M} does not

depend on \mathbf{z} . This setup fits within the conceptual framework
 227 that has been popular for theoretical and empirical studies
 228 of \mathbf{G} -matrix evolution (Jones et al., 2007; Arnold et al.,
 229 2008; Mallard et al., 2023a). Fitness is quantified by a rate
 230 $m(\nu, \mathbf{z})$, which is the growth rate for the sub-population of
 231 individuals with trait value \mathbf{z} in a population summarized by
 232 ν . The dependency of m on ν and \mathbf{z} allows for the modelling
 233 of interwoven ecological and evolutionary dynamics. I will
 234 also refer to $m(\nu, \mathbf{z})$ as a *fitness function*, and this function
 235 may also depend on environmental parameters such as the
 236 trait values of individuals in interacting species. However, I
 237 omit notation accounting for such possibilities to simplify the
 238 frameworks presentation. Putting these components together,
 239 the multivariate generalization of DAGA is given by the partial
 240 differential equation
 241

$$\dot{\nu}(\mathbf{z}) = m(\nu, \mathbf{z}) \nu(\mathbf{z}) + \frac{1}{2} \nabla^\top \mathbf{M} \nabla \nu(\mathbf{z}), \quad (1)$$

242 where $\dot{\nu}(\mathbf{z})$ is the instantaneous rate of change of $\nu(\mathbf{z})$ over time,
 243 the symbol $\nabla := (\partial/\partial z_1, \dots, \partial/\partial z_d)^\top$ is the gradient operator
 244 with respect to the d -dimensional trait \mathbf{z} , and
 245

$$\frac{1}{2} \nabla^\top \mathbf{M} \nabla := \frac{1}{2} \sum_{i,j=1}^d M_{ij} \frac{\partial}{\partial z_i} \frac{\partial}{\partial z_j} \quad (2)$$

246 is the mutation operator. Equation (1) can be used to
 247 obtain a multivariate generalization of Kimura's continuum-
 248 of-alleles model (Kimura, 1965) that assumes diffusion instead
 249 of convolution with an arbitrary mutation kernel. Justification
 250 for multivariate DAGA (equation (1)) as a deterministic
 251 approximation of the full stochastic framework is given in
 252 Supplementary Material, Section 7.

253 If $M_{ij} = 0$ when $i \neq j$ and $M_{ii} = M_{jj}$ is constant across
 254 all ij , then the above mutation operator becomes proportional
 255 to the Laplacian operator on \mathbb{R}^d , which causes symmetric
 256 diffusion of the population density. Hence, unequal entries
 257 imply that mutation results in asymmetric diffusion across trait
 258 space so that mutation affects some traits more than others.
 259 Additionally, if $M_{ij} \neq 0$ when $i \neq j$, mutation contributes to
 260 covariance between traits z_i and z_j . This model of mutation can
 261 be obtained from a diffusion-limit of an individual-based model
 262 that assumes independence of reproduction and mutation in
 263 which the phenotypic effect size of mutation is small (see
 264 Supplementary Material, Section 7). A model where mutation
 265 and reproduction interact has been studied by Wickman et al.,
 266 (2023).

267 Mathematically, assuming n is finite allows us to write $\dot{n} =$
 $\int_{\mathbb{R}^d} \dot{\nu}(\mathbf{z}) d\mathbf{z}$. We can then apply integration-by-parts to obtain
 268

$$\dot{n} = \bar{m} n, \quad (3)$$

269 where $\bar{m}(\nu) = \int_{\mathbb{R}^d} m(\nu, \mathbf{z}) p(\mathbf{z}) d\mathbf{z}$ is mean fitness. Biologically,
 270 n should be very large since drift is being ignored. However,
 271 this approach can still be useful for gaining insights when n
 272 is small, which occurs for example with Lotka-Volterra models
 273 (Akjouj et al., 2024).

274 Dynamics of the mean trait vector are obtained by applying
 275 the quotient rule and integration-by-parts to $\dot{\bar{\mathbf{z}}} = \int_{\mathbb{R}^d} \mathbf{z} \dot{p}(\mathbf{z}) d\mathbf{z}$,
 276 which provides

$$\dot{\bar{\mathbf{z}}} = \text{Cov}(m, \mathbf{z}), \quad (4)$$

277 where $\text{Cov}(m, \mathbf{z})$ is a d -dimensional vector with i -th entry given
 278 by
 279

$$\text{Cov}(m, z_i) = \int_{\mathbb{R}^d} (m(\nu, \mathbf{z}) - \bar{m}(\nu)) (z_i - \bar{z}_i) p(\mathbf{z}) d\mathbf{z}, \quad (5)$$

280 where \bar{z}_i is the i -th entry of $\bar{\mathbf{z}}$.
 281

284 The same techniques can also be applied to obtain $\dot{\mathbf{P}}$ as

285
$$\dot{\mathbf{P}} = \mathbf{M} + \text{Cov}(m, (\mathbf{z} - \bar{\mathbf{z}})(\mathbf{z} - \bar{\mathbf{z}})^\top), \quad (6)$$

286 where $\text{Cov}(m, (\mathbf{z} - \bar{\mathbf{z}})(\mathbf{z} - \bar{\mathbf{z}})^\top)$ is a $d \times d$ matrix with ij -th entry
287 given by $\text{Cov}(m, (z_i - \bar{z}_i)(z_j - \bar{z}_j))$. The response of the trait
288 covariance matrix to selection as a covariance of fitness with
289 $(\mathbf{z} - \bar{\mathbf{z}})(\mathbf{z} - \bar{\mathbf{z}})^\top$ was previously found by Lande and Arnold
290 (1983). The steps required to arrive at equations (3)-(6) are
291 detailed in the Supplementary Material, Section 2.

292 Equations (3), (4), and (6) are especially flexible because
293 they make no assumptions on the trait distribution. In this
294 form, the framework can then be used to study the dynamics
295 of populations with non-trivial higher moments, such as skewed
296 distributions. However, this flexibility of accommodating
297 general trait distributions comes at the cost of a limited range of
298 fitness functions that are amenable to study. This consequence
299 is due to i) the challenge of calculating covariances between
300 arbitrary fitness functions and phenotypic moments, and ii)
301 moment-closing issues that often arise during these calculations
302 (Barton and Turelli, 1987; Gilpin and Feldman, 2019; Guerand
303 et al., 2023). In spite of this limitation, a family of biologically
304 important fitness functions are tractable to study using *DC*.
305 These function take the form

306
$$m(\nu, \mathbf{z}) = r + \mathbf{b}^\top \mathbf{z} - \frac{1}{2}(\boldsymbol{\theta} - \mathbf{z})^\top \Psi(\boldsymbol{\theta} - \mathbf{z}) - c n, \quad (7)$$

307 where $\mathbf{b} = (b_1, \dots, b_d)^\top$ (which does not depend on \mathbf{z})
308 confers directional selection (positive or negative for trait z_i
309 depending on the sign of b_i), and Ψ is a symmetric matrix
310 that mediates stabilizing or disruptive selection around the
311 vector $\boldsymbol{\theta}$. Classically, Ψ is denoted ω^{-1} in quantitative genetic
312 theory (e.g., Lande, 1979; Jones et al., 2012). The absence of
313 stabilizing selection under the classical notation occurs when all
314 eigenvalues of ω diverge towards $+\infty$. To avoid this technicality,
315 I use Ψ so that the absence of stabilizing selection is associated
316 with the zero matrix $\Psi = \mathbf{0}$. Additionally, the parameter $c \geq 0$
317 is the strength of competition, which here acts globally between
318 individuals regardless of trait, and r is the growth rate in the
319 absence of selection and competition (i.e., when \mathbf{b}, Ψ , and c
320 are all zero). It is possible to generalize this family so that
321 competition is non-global, but this requires additional technical
322 details as the fitness function becomes operator-valued (which
323 is treated, for example, by Volpert, 2014; and by Etheridge
324 et al., 2024). Supplementary Material, Section 3 connects these
325 fitness functions to those known from classical discrete time
326 quantitative genetic models.

327 Combining equations (3), (4), and (6) with equation (7)
328 provides (derivation given in Supplementary Material, Section
329 2.3)

330
$$\dot{n} = \left(r + \mathbf{b}^\top \bar{\mathbf{z}} - \frac{1}{2}(\boldsymbol{\theta} - \bar{\mathbf{z}})^\top \Psi(\boldsymbol{\theta} - \bar{\mathbf{z}}) - \frac{1}{2}\text{tr}(\Psi \mathbf{P}) - c n \right) n, \quad (8a)$$

331
$$\dot{\bar{\mathbf{z}}} = \mathbf{P} \mathbf{b} + \mathbf{P} \Psi(\boldsymbol{\theta} - \bar{\mathbf{z}}) - \frac{1}{2} \mathbf{S} : \Psi, \quad (8b)$$

332
$$\dot{\mathbf{P}} = \mathbf{M} + \frac{1}{2}\text{tr}(\Psi \mathbf{P}) \mathbf{P} + \mathbf{S} \cdot (\Psi(\boldsymbol{\theta} - \bar{\mathbf{z}})) + \mathbf{S} \cdot \mathbf{b} - \frac{1}{2} \mathbf{K} : \Psi, \quad (8c)$$

333 where \mathbf{S} is the third-order skew tensor of \mathbf{z} defined by $S_{ijk} =$
334 $\int_{\mathbb{R}^d} (z_i - \bar{z}_i)(z_j - \bar{z}_j)(z_k - \bar{z}_k)p(\mathbf{z})d\mathbf{z}$. Products with \mathbf{S} are
335 given by $(\mathbf{S} : \Psi)_i = \sum_{jk} S_{jki} \Psi_{jk}$, $(\mathbf{S} \cdot \Psi)_{ij} = \sum_k S_{ijk} \Psi_{kj}$,
336 and $(\mathbf{S} \cdot \mathbf{b})_{ij} = \sum_k S_{ijk} b_k$. In addition, \mathbf{K} is the fourth-
337 order kurtosis tensor of \mathbf{z} defined by $K_{ijkl} = \int_{\mathbb{R}^d} (z_i -$
338 $\bar{z}_i)(z_j - \bar{z}_j)(z_k - \bar{z}_k)(z_l - \bar{z}_l)p(\mathbf{z})d\mathbf{z}$. The product with \mathbf{K} is

339 given by $(\mathbf{K} : \Psi)_{ij} = \sum_{kl} K_{ijkl} \Psi_{kl}$. Equation (8) collects
340 the abundance, mean, and covariance dynamics (including
341 skewness and kurtosis contributions) into a single system
342 for clarity. While related formulations exist, this particular
343 consolidation does not appear to be standard in the literature.
344

345 Equations (8) demonstrate that, for fitness functions taking
346 the form of (7), the dynamics of abundance, mean trait,
347 and trait covariance matrix depend on higher phenotypic
348 moments for non-normal trait distributions. In fact, a complete
349 description requires an infinite number of equations. However,
350 by combining equation (7) with multivariate DAGA (i.e.,
351 with equation (1)), I demonstrate in Supplementary Material,
352 Section 2.5 that, when Ψ is positive definite, $\nu(\mathbf{z})$ has a globally
353 asymptotically stable equilibrium proportional to the density of
354 a multivariate normal distribution with covariance matrix
355

356
$$\hat{\mathbf{P}} = \Psi^{-1/2} (\Psi^{1/2} \mathbf{M} \Psi^{1/2})^{1/2} \Psi^{-1/2} \quad (9)$$

357 (with matrix square roots defined using eigenvalue decomposition:
358 $\mathbf{Q} = \mathbf{U} \Lambda \mathbf{U}^{-1}$ implies $\mathbf{Q}^{1/2} = \mathbf{U} \Lambda^{1/2} \mathbf{U}^{-1}$) and equilibrium
359 mean vector

360
$$\hat{\mathbf{z}} = \Psi^{-1} \mathbf{b} + \boldsymbol{\theta}, \quad (10)$$

361 and with a total abundance

362
$$\hat{n} = \frac{1}{c} \left(r + \mathbf{b}^\top \boldsymbol{\theta} + \frac{1}{2} \mathbf{b}^\top \Psi^{-1} \mathbf{b} - \frac{1}{2} \text{tr}(\sqrt{\mathbf{M} \Psi}) \right). \quad (11)$$

363 Derivations for these equilibrium expressions are given in
364 Supplementary Material, Section 2.4.

365 The mutation-selection balance of phenotypic variance $\hat{\mathbf{P}}$ is
366 a multivariate generalization of the univariate classical result
367 obtained from multi-locus models (e.g., Bulmer, 1972) and
368 continuum-of-alleles models (e.g., Bürger, 1986). Lande (1980)
369 found a similar result using a multilocus approach that included
370 recombination. In particular, $\hat{\mathbf{P}}$ has the same matrix-square-
371 root structure as the equilibrium found by Lande (1980),
372 and the two coincide when linkage is negligible and when
373 \mathbf{M} corresponds to the sum of Lande's locus-specific mutation
374 matrices. Further work is needed to extend these results for
375 more general fitness functions and higher phenotypic moments,
376 which are already known to have important evolutionary
377 consequences (Débarre et al., 2015). A new approach to study
378 the dynamics of higher moments for univariate traits was
379 recently introduced by Gilpin and Feldman (2019).

2.1.2. The Deterministic Gradient Version (DG)

380 As noted above, deriving population dynamics from arbitrary
381 fitness functions can be challenging in the more general
382 framework based on covariances between fitness and phenotype.
383 To overcome this we can make the useful simplifying
384 assumption that traits follow a multivariate normal distribution.
385 Under this assumption, all phenotypic moments (such as
386 skew and kurtosis) can be expressed in terms of the mean
387 and variance which greatly simplifies expressions such as
388 equations (8). While deviations from normality can have
389 consequences for ecological and evolutionary processes (Turelli,
390 1988; Débarre et al., 2015), normality has been an important
391 initial assumption for studying a wide range of topics such as
392 reaction-norm evolution (Lande, 2014), coevolving mutualistic
393 networks (Nuismer et al., 2018), and niche construction
394 (Fogarty and Wade, 2022). Furthermore, normality is a well-
395 established approximation that holds under many genetic and
396 selective scenarios (Turelli and Barton, 1994). Then, as a first
397 pass, I assume traits are multivariate normally distributed for
398 the rest of this paper.

In the context of this framework, the assumption of multivariate normality is particularly useful because it allows us to rewrite covariances between fitness and phenotype as gradients of fitness functions with respect to moments of the trait distribution. Such gradients can be analytically calculated for a broad range of fitness functions. The calculations to obtain these expressions begin with the definitions of covariances between fitness and phenotype, and then apply properties of the multivariate Gaussian function and integration-by-parts (see Supplementary Material, Section 2.3). As a result, the deterministic *DG* version of the framework is given by

$$\dot{n} = \bar{m} n, \quad (12a)$$

$$\dot{\bar{z}} = \mathbf{P} (\nabla_{\bar{z}} \bar{m} - \overline{\nabla_{\bar{z}} m}), \quad (12b)$$

$$\dot{\mathbf{P}} = \mathbf{M} + 2 \mathbf{P} (\nabla_{\mathbf{P}} \bar{m} - \overline{\nabla_{\mathbf{P}} m}) \mathbf{P}, \quad (12c)$$

where $\nabla_{\bar{z}} \bar{m}$ and $\overline{\nabla_{\bar{z}} m}$ are d -dimensional vectors that respectively capture the effects of frequency independent and frequency dependent selection on mean trait evolution. More precisely, writing $\partial_i = \partial/\partial \bar{z}_i$ as the partial derivative operator with respect to the i -th mean trait, the i -th entry of $\nabla_{\bar{z}} \bar{m}$ and $\overline{\nabla_{\bar{z}} m}$ are respectively given by $\partial_i \bar{m}$ and $\overline{\partial_i m} = \int_{\mathbb{R}^d} p(\mathbf{z}) [\partial_i m(\nu, \mathbf{z})] d\mathbf{z}$. Similarly, $\nabla_{\mathbf{P}} \bar{m}$ and $\overline{\nabla_{\mathbf{P}} m}$ are $d \times d$ matrices that respectively quantify the dependence of fitness on trait variances and covariances. Writing $\partial_{ij} = \partial/\partial P_{ij}$ as the partial derivative operator with respect to the covariance between trait components i and j , the ij -th entries of $\nabla_{\mathbf{P}} \bar{m}$ and $\overline{\nabla_{\mathbf{P}} m}$ are respectively given by $\partial_{ij} \bar{m}$ and $\overline{\partial_{ij} m} = \int_{\mathbb{R}^d} p(\mathbf{z}) [\partial_{ij} m(\nu, \mathbf{z})] d\mathbf{z}$.

In Supplementary Material, Section 2.3, I show how to apply integration-by-parts to obtain the alternative expressions $\dot{\bar{z}} = \mathbf{P} \overline{\nabla m}$ and $\dot{\mathbf{P}} = \mathbf{M} + \mathbf{P} \overline{\nabla^2 m} \mathbf{P}$ where $(\nabla)_i = \partial/\partial z_i$ (so $\overline{\nabla m}$ is the trait averaged gradient of fitness with respect to phenotype \mathbf{z}) and $(\nabla^2)_{ij} = \partial^2/\partial z_i \partial z_j$ (so $\overline{\nabla^2 m}$ is the trait averaged Hessian of fitness with respect to phenotype \mathbf{z}). A discrete-time counterpart to $\overline{\nabla^2 m}$ is given by equation (14b) in Lande and Arnold (1983).

The Hessian form for the response of \mathbf{P} to selection is similar to previously known discrete-time expressions (Lande and Arnold, 1983; Débarre et al., 2014; Phillips and Arnold, 1989; Mullon and Lehmann, 2019). An important difference is that the discrete-time expressions include an additional term equal to the outer product of the per-generation response of the mean trait to selection, commonly denoted $(\Delta \bar{z})(\Delta \bar{z})^\top$, which appears with a negative sign in the discrete-time update of \mathbf{P} . However, this extra term disappears under weak selection approximations often made in discrete-time quantitative genetic models (Lande, 1976, 1982; Turelli, 1984; Barton and Turelli, 1989; Turelli and Barton, 1990; Lynch and Walsh, 1998; Bürger, 2000; Walsh and Lynch, 2018). Furthermore, under a continuous-time scaling, the mean changes only $O(\Delta t)$ over a small interval, so its contribution to covariance is $O(\Delta t^2)$. Dividing by Δt to form a rate equation and letting $\Delta t \rightarrow 0$ removes this quadratic term, leaving only the curvature-driven (Hessian) part of selection in the continuous-time covariance dynamics, in agreement with equation (12c).

Mean trait and trait covariance dynamics can be expressed in index form as

$$\dot{z}_i = \sum_{j=1}^d P_{ij} (\partial_j \bar{m} - \overline{\partial_j m}), \quad (13a)$$

$$\dot{P}_{ij} = M_{ij} + 2 \sum_{k,l=1}^d P_{ik} (\partial_{kl} \bar{m} - \overline{\partial_{kl} m}) P_{lj}. \quad (13b)$$

Inclusion of the terms $\overline{\nabla_{\bar{z}} m}$ and $\overline{\nabla_{\mathbf{P}} m}$ in equations (12b) and (12c) make it clear that this framework accounts for frequency-dependent selection. The decomposition of selection into components arising from frequency-dependent and frequency-independent sources has been made explicit in previous quantitative genetic treatments of mean trait dynamics (see eqn. (9) of Lande, 1976), but was only implicit in models of trait variance evolution due to their expressions in terms of Hessians (Lande and Arnold, 1983; Débarre et al., 2014; Phillips and Arnold, 1989; Mullon and Lehmann, 2019). A decomposition of the Hessian of fitness into frequency-dependent and frequency-independent components does not appear to be standard in existing treatments of trait-covariance evolution. Additionally, the approach outlined here can make a useful complement to adaptive dynamics and evolutionary game-theoretic treatments of frequency-dependent selection (Dieckmann and Law, 1996; Hofbauer and Sigmund, 1998; Traulsen et al., 2005).

An important caveat of both *DC* and *DG* versions of the framework taking the form presented above is that they rely on perfect heritability of trait values. However, traits are in general not perfectly heritable, and this is of fundamental importance in evolutionary biology. I therefore address this additional complexity in the following sub-subsection. I note here that extending *DC* and *DG* to include imperfect heritability results in nearly identical equations, the important difference is that average growth rates must additionally be averaged over a phenotypic residual term. I state this explicitly for the *DG* version below.

2.1.3. Imperfect Inheritance

Following classical quantitative genetics, imperfect inheritance of trait values can be accounted for by assuming traits decompose into an additive genetic component and a residual component: $\mathbf{z} = \mathbf{g} + \mathbf{e}$ (Lynch and Walsh, 1998).

Biologically, this decomposition assumes that non-additive genetic effects and genotype-environment interactions are absent, so that \mathbf{g} reflects only additive genetic effects, while \mathbf{e} reflects the remaining non-heritable factors. This justifies interpreting \mathbf{e} as the residual from fitting a linear statistical model of trait values to genetic predictors (Falconer and Mackay, 1996; Lynch and Walsh, 1998). The absence of non-additive genetic effects is especially important in this asexual framework, where the lack of recombination would otherwise allow multilocus allelic associations to render such effects heritable (Barton and Turelli, 1991). It is further assumed that the only trait-mediating factors that are heritable are additive genetic factors (i.e., cytoplasmic inheritance, persistent epigenetic marks, maternal effects, microbiome transmission, and cultural inheritance are all excluded). Under these assumptions, \mathbf{e} is uncorrelated with \mathbf{g} and is uncorrelated among individuals.

In contrast to \mathbf{e} , the additive genetic component \mathbf{g} follows the same Gaussian mutation model described above. In particular, given that \mathbf{g} is the additive genetic component of the trait of a parent, an offspring will have an additive genetic component that is multivariate normally distributed with mean \mathbf{g} and covariance matrix \mathbf{M} . Assuming the residual \mathbf{e} is independent and identically distributed for all individuals with mean zero and covariance matrix \mathbf{E} , and denoting $\gamma(\mathbf{g})$ the population density at genetic value \mathbf{g} , and $\varepsilon(\mathbf{e})$ the distribution

of the residuals, the population density of trait values is given by $\nu(\mathbf{z}) = \int_{\mathbb{R}^d} \varepsilon(\mathbf{e}) \gamma(\mathbf{z} - \mathbf{e}) d\mathbf{e}$. This preserves total abundance so that $n = \int_{\mathbb{R}^d} \nu(\mathbf{z}) d\mathbf{z} = \int_{\mathbb{R}^d} \gamma(\mathbf{g}) d\mathbf{g}$. These assumptions also imply that the growth rate for the sub-population of individuals carrying genetic value \mathbf{g} is given by $\tilde{m}(\gamma, \mathbf{g}) = \int_{\mathbb{R}^d} \varepsilon(\mathbf{e}) m(\nu, \mathbf{g} + \mathbf{e}) d\mathbf{e}$. Furthermore, the fitness function for trait values $m(\nu, \mathbf{z})$ and the fitness function for additive genetic values $\tilde{m}(\gamma, \mathbf{g})$ have the same mean value across the population

$$\begin{aligned} \bar{\tilde{m}} &= \frac{1}{n} \int_{\mathbb{R}^d} \tilde{m}(\gamma, \mathbf{g}) \gamma(\mathbf{g}) d\mathbf{g} \\ &= \frac{1}{n} \int_{\mathbb{R}^d} \int_{\mathbb{R}^d} m(\nu, \mathbf{z}) \varepsilon(\mathbf{e}) \gamma(\mathbf{z} - \mathbf{e}) d\mathbf{e} d\mathbf{z} \\ &= \frac{1}{n} \int_{\mathbb{R}^d} m(\nu, \mathbf{z}) \nu(\mathbf{z}) d\mathbf{z} = \bar{m}. \end{aligned} \quad (14)$$

This approach to deriving the fitness associated with an additive genetic value given the fitness associated with a trait value and the distribution of residuals was originally introduced by Kimura and Crow (1978). It also appears in equation (4) of Lande (1979). The dynamics of γ are given in analogy to $\dot{\nu}$ as

$$\dot{\gamma}(\mathbf{g}) = \tilde{m}(\gamma, \mathbf{g}) \gamma(\mathbf{g}) + \frac{1}{2} \nabla^\top \mathbf{M} \nabla \gamma(\mathbf{g}). \quad (15)$$

The distribution of additive genetic values in the population is given by $\tilde{p}(\mathbf{g}) = \gamma(\mathbf{g})/n$. In combination with the model for phenotypic residuals, the mean trait vector is calculated as $\bar{\mathbf{z}} = \bar{\mathbf{g}}$ and the trait covariance matrix is $\mathbf{P} = \mathbf{G} + \mathbf{E}$. To simplify calculating fitness, and to accommodate the assumption of multivariate normal traits, I assume $\varepsilon(\mathbf{e})$ is the density of a multivariate normal distribution (with mean zero and covariance matrix \mathbf{E}).

Under these assumptions the expression for abundance dynamics does not change, but the mean trait dynamics can be calculated as

$$\dot{\bar{\mathbf{z}}} = \mathbf{G} (\nabla_{\bar{\mathbf{z}}} \bar{m} - \overline{\nabla_{\bar{\mathbf{z}}} m}), \quad (16)$$

and \mathbf{G} -matrix dynamics are given by

$$\dot{\mathbf{G}} = \mathbf{M} + 2 \mathbf{G} (\nabla_{\mathbf{G}} \bar{m} - \overline{\nabla_{\mathbf{G}} m}) \mathbf{G}. \quad (17)$$

I apply this model of imperfect inheritance while describing the stochastic extensions of the framework below. Further details about how imperfect inheritance is interfaced with the stochastic extension of this framework, and how to obtain equations (16) and (17) are provided in the Supplementary Material, Section 5.

In summary, this section derives deterministic equations for abundance, mean traits, and trait covariances from the DAGA formulation (equation (1)). It presents both the non-closed moment system for general trait distributions (equations (3)–(8)) and the closed gradient/Hessian form obtained under normality with imperfect inheritance (equations (12a), (16), and (17)), which the stochastic extensions build on below.

2.2. Stochastic Dynamics

In this section, I extend the framework to the case where demographic stochasticity induces random genetic drift. This extension introduces the stochastic version of the framework and builds on the Deterministic Gradient (*DG*) formulation developed above. Two closely related stochastic variants are presented.

The first variant, called the Brownian Motion Gradient version (*BG*), expresses stochastic dynamics with respect to Brownian motion processes, as is standard for stochastic differential equations (Øksendal, 2013; Evans, 2012). *BG* is particularly useful for numerical applications, and can be implemented using the Euler-Maruyama algorithm (Bayram et al., 2018). I illustrate this during a brief study of genetic correlations evolving in response to random genetic drift. Further information is provided in Supplementary Material, Section 1. Implementations of this approach using the DifferentialEquations.jl package in Julia (Rackauckas and Nie, 2017), and a manual implementation are provided at the github repository github.com/bobweek/multi-mtgc.

The second variant, called the Martingale Gradient version (*MG*), is useful for deriving the dynamics of more specific quantities. This use-case is illustrated below to formally obtain a stochastic differential equation tracking the evolution of genetic correlations in response to random genetic drift. To do so, I introduce a powerful set of novel heuristics (i.e., methods for performing exact calculations without formal justification, see Table 1) that can be used to derive an array of models from this variant of the framework. These heuristics are obtained by examining a so-called martingale process underlying this entire framework, which is also the namesake of *MG*. Mathematical details are provided in Section 4 of the Supplement.

Both stochastic extensions of the framework include an additional variable v , which is associated with the variance of lifetime reproductive output of individuals. Supplementary Material, Section 7.1 shows how to derive v from an individual-based model. This parameter can be interpreted as the population average of what Gillespie (1974; 1975; 1977) calls the variance in offspring number. Supplementary Material, Section 8 formally demonstrates this interpretation. Importantly, v can also be interpreted as either the rate of reproduction for semelparous populations (demonstrated in Supplementary Material, Section 7) or the sum of birth and death rates in iteroparous populations (discussed in Week et al., 2021). I refer to v as the *reproductive variance*.

Previous work has shown that a novel form of selection can emerge when v depends on trait value, referred to as *noise-induced selection*, and for which evolutionary responses behave radically different from classical deterministic selection (Gillespie, 1974, 1977; Constable et al., 2016; Parsons et al., 2010; Kuosmanen et al., 2022; Bhat and Guttal, 2025). However, accounting for noise-induced selection leads to significantly more complex expressions for the evolution of mean trait vector and trait covariance matrix (e.g., see Week et al., 2021, for the univariate case). While genotype-dependent differences in the variance of offspring number have been documented in laboratory asexual systems (Turner and Chao, 1999), there is no evidence that such covariance is widespread, and selection acting explicitly on this variance component has not been directly measured in empirical populations. Then, as a first pass, I keep the expressions relatively simple by assuming v is constant across additive genetic values.

2.2.1. The Brownian Motion Gradient Version (*BG*)

Accounting for the effects of demographic stochasticity, the abundance dynamics can now be expressed as the following stochastic differential equation

$$dn = \bar{m} n dt + \sqrt{v n} dB_n, \quad (18)$$

where the noise process B_n is a standard Brownian motion (i.e., $B_n(t)$ has variance equal to t and $B_n(0) = 0$). Assuming multivariate normality, and the above model of imperfect inheritance, the mean trait dynamics (derived in Supplementary Material, Section 5.3) are given by

$$d\bar{\mathbf{z}} = \mathbf{G} (\nabla_{\bar{\mathbf{z}}} \bar{m} - \overline{\nabla_{\bar{\mathbf{z}}} m}) dt + \sqrt{\frac{v}{n}} \mathbf{G} dB_{\bar{\mathbf{z}}}, \quad (19)$$

with $\mathbf{B}_{\bar{\mathbf{z}}}$ a d -dimensional vector of independent standard Brownian motions, and the matrix square root $\sqrt{\mathbf{G}}$ can be any $d \times d$ matrix \mathbf{A} that satisfies $\mathbf{A}\mathbf{A}^\top = \mathbf{G}$ (for examples, see Kessy et al., 2018). Equation (19) provides a continuous-time extension to the framework of multivariate evolution introduced by Lande (1979). Alternatively, the stochastic dynamics of mean traits can be expressed using index notation as

$$d\bar{z}_i = \sum_{j=1}^d G_{ij} (\partial_j \bar{m} + \overline{\partial_j m}) dt + \sqrt{\frac{v}{n}} G_{ii} dB_{\bar{z}_i}, \quad (20)$$

where $\partial_i = \partial/\partial \bar{z}_i$ and, for each i , $B_{\bar{z}_i}$ is a standard Brownian motion and non-independence for each ij is encoded by the heuristic $dB_{\bar{z}_i} dB_{\bar{z}_j} = \rho_{ij} dt$, with $\rho_{ij} = G_{ij}/\sqrt{G_{ii} G_{jj}}$ the genetic correlation between traits i and j . This heuristic is particularly useful when applying Itô's formula (the stochastic analog of the chain rule, see Klebaner, 1998; Øksendal, 2013; or Evans, 2012) to derive dynamics for functions of mean trait values. This approach may be used, for example, to track the dynamics of interaction coefficients for coevolving species as a multivariate generalization of the approach taken by Week and Nuismer (2021).

In equation (19) the correlated effects of genetic drift on mean trait evolution are encoded by the product $(\sqrt{\mathbf{G}} dB_{\bar{\mathbf{z}}})_i = \sum_j (\sqrt{\mathbf{G}})_{ij} (dB_{\bar{\mathbf{z}}})_j$. In contrast, for equation (20), these correlated effects are encoded directly by the non-independence of the Brownian motions $B_{\bar{z}_1}, \dots, B_{\bar{z}_d}$. In particular, because $\mathbf{B}_{\bar{\mathbf{z}}}$ is defined such that its entries are independent, the i -th entry of $d\mathbf{B}_{\bar{\mathbf{z}}}$ cannot be equal to $dB_{\bar{z}_i}$ as $(d\mathbf{B}_{\bar{\mathbf{z}}})_i (d\mathbf{B}_{\bar{\mathbf{z}}})_j = \delta_{ij} dt$, where $\delta_{ij} = 1$ when $i = j$ and zero otherwise. These heuristics work in the absence of multivariate normality, but the deterministic component of $d\bar{\mathbf{z}}$ expressed above does depend on multivariate normality.

The expression of $d\bar{z}_i$ in equation (20) using non-independent Brownian motions is not standard. However, this form will be useful below to illustrate the relationship between the *BM* (equations (18), (20) and (23)) and *MG* (equations (26)) versions of the framework.

Supplementary Material, Section 5.4 shows that the application of the multivariate normal approximation to the stochastic dynamics of the \mathbf{G} -matrix leads to the matrix equation

$$d\mathbf{G} = \left(\mathbf{M} + 2 \mathbf{G} (\nabla_{\mathbf{G}} \bar{m} - \overline{\nabla_{\mathbf{G}} m}) \mathbf{G} - \frac{v}{n} \mathbf{G} \right) dt + \sqrt{\frac{v}{n}} \boldsymbol{\Gamma} : d\mathbf{B}_{\mathbf{G}}, \quad (21)$$

where $\boldsymbol{\Gamma}$ is a fourth-order tensor describing the covariance structure for the response of \mathbf{G} to drift. Defining products of fourth-order tensors \mathbf{X} and \mathbf{Y} by $(\mathbf{X} : \mathbf{Y})_{ijkl} = \sum_{mn} \mathbf{X}_{ijmn} \mathbf{Y}_{mnkl}$, we can write $\boldsymbol{\Gamma} = \sqrt{\boldsymbol{\Gamma}} : \sqrt{\boldsymbol{\Gamma}}$. Furthermore, we have $\sqrt{\boldsymbol{\Gamma}}_{ijkl} = (\sqrt{\mathbf{G}}_{ik} \sqrt{\mathbf{G}}_{jl} + \sqrt{\mathbf{G}}_{il} \sqrt{\mathbf{G}}_{jk})/\sqrt{2}$, and $\sqrt{\mathbf{G}}_{ij}$ is the ij -th entry of $\sqrt{\mathbf{G}}$ (as mentioned beneath equation (19) above, $\sqrt{\mathbf{G}}$ is any $d \times d$ matrix such that $\sqrt{\mathbf{G}} \sqrt{\mathbf{G}}^\top = \mathbf{G}$). Importantly, this implies $\sqrt{\mathbf{G}}_{ij}$ is not equal to $\sqrt{G_{ij}}$.

The product $\sqrt{\boldsymbol{\Gamma}} : d\mathbf{B}_{\mathbf{G}}$ returns a $d \times d$ matrix with ij -th entry $(\sqrt{\boldsymbol{\Gamma}} : d\mathbf{B}_{\mathbf{G}})_{ij} = \sum_{kl} \sqrt{\boldsymbol{\Gamma}}_{ijkl} (d\mathbf{B}_{\mathbf{G}})_{kl}$. The $d \times d$

matrix-valued Brownian motion $\mathbf{B}_{\mathbf{G}}$ has independent standard Brownian motions along its diagonal entries. The off-diagonal entries are standard Brownian motions scaled by one-half (i.e., the variance of $(\mathbf{B}_{\mathbf{G}})_{ij}(t)$ is $t/2$ when $i \neq j$), and symmetric entries are equivalent so that $(\mathbf{B}_{\mathbf{G}})_{ij} = (\mathbf{B}_{\mathbf{G}})_{ji}$. The covariance structure of $\mathbf{B}_{\mathbf{G}}$ is summarized by the heuristic

$$(d\mathbf{B}_{\mathbf{G}})_{ij} (d\mathbf{B}_{\mathbf{G}})_{kl} = \frac{\delta_{ik} \delta_{jl} + \delta_{il} \delta_{jk}}{2} dt. \quad (22)$$

Scaling the off-diagonals by one-half ensures variances and covariances are correctly propagated while calculating the stochastic consequences of drift for \mathbf{G} -matrix dynamics. Further information on symmetric normal matrices can be found in Gupta and Nagar (2018), particularly Theorem 2.5.1.

Unlike the expression for the stochastic component of $d\bar{\mathbf{z}}$, the stochastic component for $d\mathbf{G}$ does depend on the assumption of a multivariate normal trait distribution.

Using index notation, the expression (21) simplifies to a $d(d+1)/2$ -dimensional system of equations summarized by

$$\begin{aligned} dG_{ij} = & \left(M_{ij} + 2 \sum_{kl} G_{ik} (\partial_{kl} \bar{m} - \overline{\partial_{kl} m}) G_{lj} - \frac{v}{n} G_{ij} \right) dt \\ & + \sqrt{\frac{v}{n}} \sqrt{G_{ii} G_{jj} + G_{ij}^2} dB_{G_{ij}}, \end{aligned} \quad (23)$$

for $1 \leq i \leq j \leq d$ where $\partial_{ij} = \partial/\partial G_{ij}$ and for each ij we have $B_{G_{ij}}$ is a standard Brownian motion with $B_{G_{ji}} = B_{G_{ji}}$. The covariance between $B_{G_{ij}}$ and $B_{G_{kl}}$ is encoded by the heuristic

$$dB_{G_{ij}} dB_{G_{kl}} = \frac{G_{ik} G_{jl} + G_{il} G_{jk}}{\sqrt{(G_{ii} G_{jj} + G_{ij}^2)(G_{kk} G_{ll} + G_{kl}^2)}} dt. \quad (24)$$

Importantly, this implies that $dB_{G_{ij}}$ is not the ij -th entry of $d\mathbf{B}_{\mathbf{G}}$.

Finally, the noise process driving abundance in general does not covary with the noise processes driving mean traits and trait covariances (i.e., $dB_n dB_{\bar{z}_i} = dB_n dB_{G_{ki}} = 0$), and, under the assumption of multivariate normality, the same holds for trait means and trait covariances (i.e., $dB_{\bar{z}_i} dB_{G_{kl}} = 0$). This agrees with the off-diagonal entries in equation (7b) of Barton (1989), but my results for the diagonal entries disagree. See Supplementary Material, Section 5.4 for more details.

The vector-matrix expression of *BG* (equations (19) and (21)) is particularly well-suited for numerical exploration of models because the covariance structure of the noise processes associated with random genetic drift are written explicitly in terms of sums involving the entries of the matrix square root $\sqrt{\mathbf{G}}$. Hence, this version of the framework is easily interfaced with a common algorithm to numerically integrate systems of stochastic differential equations, the Euler-Maruyama method (Bayram et al., 2018). The numerical benefits of the *BG* version of the framework are highlighted in Section 3 below to study \mathbf{G} -matrix evolution.

The expression of *BG* in index notation (equations 20 and 23), while arguably more complex in appearance, are useful for deriving analytical results, and especially when applying Itô's formula to derive the dynamics of a quantity depending on $\bar{\mathbf{z}}$ and/or \mathbf{G} (such as the dynamics of growth rate, $d\bar{m}$, for instance). However, there are limitations with this application of *BG*, particularly for dealing with sums of stochastic differentials such as $U = a dB_{G_{ij}} + b dB_{G_{kl}}$. If the Brownian motions $B_{G_{ij}}$ and $B_{G_{kl}}$ were independent, then we

can write $U = \sqrt{a^2 + b^2} dB$ with B being another standard Brownian motion. Because $B_{G_{ij}}$ and $B_{G_{kl}}$ are not independent, which is captured by equation (24) above, it is not obvious how to properly express U in terms of a, b and a single standard Brownian motion B .

To overcome these challenges, and also to simplify typical calculations, it is useful to rewrite the stochastic components of equations (20) and (23) in terms of an underlying stochastic process \mathcal{M} . Because \mathcal{M} satisfies a martingale property (roughly, this means $\mathbb{E}[\mathcal{M}(t+s) | \mathcal{M}(t)] = \mathcal{M}(t)$ for all $s > 0$), I refer to this version of the framework as the Martingale Gradient version (or MG for short). In the following subsection, I present this version of the framework in a way that minimizes technical pre-requisites, while maintaining the same scope as equations (20) and (23) above. Additionally, heuristics are offered in Table 1 for making calculations, and these are demonstrated by deriving a stochastic differential equation for the correlation between two traits driven purely by random genetic drift.

2.2.2. The Martingale Gradient Version (MG)

In Supplementary Material, Section 4, I show that this framework is justified based on a martingale process \mathcal{M} . Brownian motion is a special case of a martingale process, and stochastic equations are often expressed in terms of a Brownian stochastic differential dB . However, in this framework, stochastic equations can also be expressed in terms of the stochastic differential with respect to \mathcal{M} , denoted by $d\mathcal{M}$. This establishes a powerful approach for deriving fundamental insights into evolutionary processes. Further mathematical details are provided in Supplementary Material, Sections 4 and 5, but here I focus on pragmatic aspects regarding calculations that involve $d\mathcal{M}$.

I begin by artificially defining symbols related to $d\mathcal{M}$ in terms of the Brownian motions that appear in equations (18), (19), and (21), and use these definitions to express the framework in terms of $d\mathcal{M}$. This is done purely for the sake of motivating the material that follows. After this, I introduce some properties of $d\mathcal{M}$ and show how these can be used to recover the Brownian motions initially used in the artificial definitions mentioned above. I then provide general heuristics summarized in Table 1 for working with $d\mathcal{M}$, and illustrate these heuristics by deriving the response of trait correlations to random genetic drift.

For now, define the symbols $d\mathcal{M}(1)$, $d\mathcal{M}(g_i - \bar{g}_i)$ and $d\mathcal{M}((g_i - \bar{g}_i)(g_j - \bar{g}_j) - G_{ij})$ as follows:

$$d\mathcal{M}(1) := \sqrt{v n} dB_n, \quad (25a)$$

$$d\mathcal{M}(g_i - \bar{g}_i) := \sqrt{v n G_{ii}} dB_{\bar{z}_i}, \quad (25b)$$

$$d\mathcal{M}((g_i - \bar{g}_i)(g_j - \bar{g}_j) - G_{ij}) := \sqrt{v n} \sqrt{G_{ii} G_{jj} + G_{ij}^2} dB_{G_{ij}}. \quad (25c)$$

Using this notation, the MG version of the framework can be written as

$$dn = \bar{m} n dt + d\mathcal{M}(1), \quad (26a)$$

$$d\bar{z}_i = \sum_{j=1}^d G_{ij} (\partial_j \bar{m} + \overline{\partial_j m}) dt + \frac{1}{n} d\mathcal{M}(g_i - \bar{g}_i), \quad (26b)$$

$$\begin{aligned} dG_{ij} = & \left(M_{ij} + 2 \sum_{kl} G_{ik} (\partial_{kl} \bar{m} - \overline{\partial_{kl} m}) G_{lj} - \frac{v}{n} G_{ij} \right) dt \\ & + \frac{1}{n} d\mathcal{M}((g_i - \bar{g}_i)(g_j - \bar{g}_j) - G_{ij}), \end{aligned} \quad (26c)$$

where recall that $\partial_i = \partial/\partial \bar{z}_i$ and $\partial_{ij} = \partial/\partial G_{ij}$.

The covariance structure of the system is maintained through heuristics (summarized in Table 1) for computing products of the above stochastic differentials $d\mathcal{M}(x)$. To understand the heuristics, I introduce some useful notation for keeping track of averages across the distribution of traits in the population. Specifically, for functions $x(\mathbf{g})$ and $y(\mathbf{g})$, I define the symbols $\|x\|$ and $\langle x, y \rangle$ as follows:

$$\|x\| = \sqrt{v n} \sqrt{\overline{x^2}} = \sqrt{v n} \sqrt{\int_{\mathbb{R}^d} x^2(\mathbf{g}) \tilde{p}(\mathbf{g}) d\mathbf{g}}, \quad (27a)$$

$$\langle x, y \rangle = v n \overline{xy} = v n \int_{\mathbb{R}^d} x(\mathbf{g}) y(\mathbf{g}) \tilde{p}(\mathbf{g}) d\mathbf{g}, \quad (27b)$$

where $\tilde{p}(\mathbf{g}) = \gamma(\mathbf{g})/n$ is the distribution of additive genetic values \mathbf{g} in the population (assumed to be multivariate normal). To provide a few examples, one can calculate $\|1\| = \sqrt{v n} \sqrt{\int 1^2 \tilde{p}(\mathbf{g}) d\mathbf{g}} = \sqrt{v n}$ along with

$$\|g_i - \bar{g}_i\| = \sqrt{v n} \sqrt{\int_{\mathbb{R}^d} (g_i - \bar{g}_i)^2 \tilde{p}(\mathbf{g}) d\mathbf{g}} = \sqrt{v n G_{ii}}, \quad (28a)$$

$$\langle g_i - \bar{g}_i, g_j - \bar{g}_j \rangle = v n \int_{\mathbb{R}^d} (g_i - \bar{g}_i)(g_j - \bar{g}_j) \tilde{p}(\mathbf{g}) d\mathbf{g} = v n G_{ij}. \quad (28b)$$

The martingale process \mathcal{M} mentioned above can be thought of as a mapping that associates functions with stochastic processes. This idea is formalized in Section 4 of the Supplement. In general, the process that \mathcal{M} associates with x , denoted $\mathcal{M}(x)$, will not be a Brownian motion. However, Supplementary Material, Section 4.2 provides justification for what I call the *scaling property*, which states that

$$d\mathcal{M}(\hat{x}) := \frac{d\mathcal{M}(x)}{\|x\|} \quad (29)$$

is the stochastic differential of a standard Brownian motion.

Using this heuristic, we can calculate

$$d\mathcal{M}(1) = \|1\| d\mathcal{M}(\hat{1}) = \sqrt{v n} d\mathcal{M}(\hat{1}), \quad (30a)$$

$$d\mathcal{M}(g_i - \bar{g}_i) = \|g_i - \bar{g}_i\| d\mathcal{M}(\widehat{g_i - \bar{g}_i}) = \sqrt{v n G_{ii}} d\mathcal{M}(\widehat{g_i - \bar{g}_i}). \quad (30b)$$

Comparing with equations (18) and (20), these calculations highlight the fact that $dB_n = d\mathcal{M}(\hat{1})$ and $dB_{\bar{z}_i} = d\mathcal{M}(\widehat{g_i - \bar{g}_i})$. A similar equality holds for $dB_{G_{ij}}$, but for the sake of brevity this expression is omitted (see Supplementary Material, Section 5.4 for details).

Informally speaking, this scaling property allows us to “factor out” the standard deviation from the noise process driving the dynamics of a univariate function of the population (such as n , \bar{z}_i , or G_{ij}), and to replace that noise process with the product of the resulting standard deviation with a standard Brownian motion. This is the crucial step for obtaining equations (18), (20), and (23) from the underlying martingale process \mathcal{M} , as detailed in Supplementary Material, Section 5 where the heuristics summarized in Table 1 are applied to derive MG and BG. The calculations shown in equation (30) are heuristic because they lead to accurate results without a rigorous understanding of their justification.

Just as $B_{\bar{z}_i}$ and $B_{\bar{z}_j}$ have a covariance encoded by the heuristic $dB_{\bar{z}_i} dB_{\bar{z}_j} = (G_{ij} / \sqrt{G_{ii} G_{jj}}) dt$, the processes $\mathcal{M}(x)$ and $\mathcal{M}(y)$ will also covary following a more general heuristic. More precisely, Supplementary Material, Section 4.2 justifies

the *multiplicative property* which states that, for functions x, y , we have

$$d\mathcal{M}(x) d\mathcal{M}(y) = \langle x, y \rangle dt, \quad (31)$$

where $\langle x, x \rangle = \|x\|^2$. In addition, we have $dt^2 = 0$ and $dt d\mathcal{M}(x) = d\mathcal{M}(x) dt = 0$ for any function x , which are exact equalities in standard approaches to stochastic differential equations (see Øksendal, 2013; Evans, 2012; Klebaner, 1998). In this context, phenotypic moments μ_1, μ_2 (such as \bar{z}_i or G_{ij}) behave as constants such that $\langle \mu_1 x, \mu_2 y \rangle = \mu_1 \mu_2 \langle x, y \rangle$ (but see Supplementary Material, Section 4.2 for justification).

Finally, we also have the *additive property*: $d\mathcal{M}(ax + by) = a d\mathcal{M}(x) + b d\mathcal{M}(y)$, for functions x, y and constants a, b (more generally, a, b can also be phenotypic moments such as \bar{z}_i and G_{ij} , see Section 4.2 of the Supplement). This property is particularly useful for computing the correct stochastic term for an equation resulting from the application of Itô's formula, as illustrated in Section 3 below.

Table 1. Heuristics summarizing the scaling, multiplicative, and additive properties of the stochastic differentials $d\mathcal{M}(x)$.

Property	Heuristic
Scaling	$d\mathcal{M}(x) = \ x\ d\mathcal{M}(\hat{x}), \mathcal{M}(\hat{x})$ a BM
Multiplicative	$d\mathcal{M}(x) d\mathcal{M}(y) = \langle x, y \rangle dt$
Additive	$d\mathcal{M}(ax + by) = a d\mathcal{M}(x) + b d\mathcal{M}(y)$

The heuristics associated with the *MG* version of the framework are summarized in Table 1. Example usage of these heuristics are provided through the derivation of this framework in Supplementary Material, Section 5. With these heuristics, it is straightforward to work with equations (26) to rigorously obtain the dynamics of quantities related to the population. For instance, the heuristics summarized in Table 1 may be used to formally derive a stochastic differential equation for the evolution of genetic correlations in response to drift. Indeed, I do this now.

3. Genetic Correlations and G-matrix Dynamics

In the following subsection I illustrate how to apply the above framework to obtain new models of evolutionary phenomena. Specifically, I apply the heuristics introduced in the *MG* version of the framework to derive the dynamics of genetic correlations responding only to random genetic drift. Using this derivation, I then provide biological insights into the consequences of drift for genetic correlations in asexually reproducing populations. In section 3.2, I then discuss the significance of these results in the context of empirical research on G-matrix dynamics.

3.1. Consequences of Drift for Trait Correlations

The additive genetic correlation between traits z_i and z_j is given by $\rho_{ij} = G_{ij} / \sqrt{G_{ii} G_{jj}}$. For this subsection, I focus on the single correlation between z_i and z_j , and simply write this as ρ . To focus on the effects of random genetic drift, and for the sake of simplicity, I ignore mutation, selection, and abundance dynamics in this analysis. For the sake of illustration, I outline only the steps needed to demonstrate how the framework is applied and to obtain the biological result. The full derivation is provided in Supplementary Material, Section 6.1.

Because we have the stochastic equations for dG_{ij} , dG_{ii} , and dG_{jj} (equation 26c), and because ρ can be thought of as a function $\rho(G_{ij}, G_{ii}, G_{jj}) = G_{ij} / \sqrt{G_{ii} G_{jj}}$, we can apply Itô's

formula, which, in this context, states the following stochastic analog of the chain-rule: $d\rho = K + L$, with $K = (\partial_{ij}\rho) dG_{ij} + (\partial_{ii}\rho) dG_{ii} + (\partial_{jj}\rho) dG_{jj}$ and

$$\begin{aligned} L = \frac{1}{2} & \left[(\partial_{ij}^2 \rho) (dG_{ij})^2 + (\partial_{ii}^2 \rho) (dG_{ii})^2 + (\partial_{jj}^2 \rho) (dG_{jj})^2 \right] \\ & + (\partial_{ij} \partial_{ii} \rho) (dG_{ij}) (dG_{ii}) + (\partial_{ij} \partial_{jj} \rho) (dG_{ij}) (dG_{jj}) \\ & + (\partial_{ii} \partial_{jj} \rho) (dG_{ii}) (dG_{jj}), \end{aligned} \quad (32)$$

where $\partial_{ij}\rho$ is the partial derivative of ρ with respect to G_{ij} .

With this formula in hand, all that is left is to carry out calculations based on the heuristics from the *MG* version of the framework. First off, we can directly compute the *deterministic* component of the sum K and find that this cancels to zero. This part of the calculation does not require the heuristics for $d\mathcal{M}$ introduced above. However, by using equation (26c) in the expression for K , and by leveraging the additive property of $d\mathcal{M}$, the *stochastic* component of K can be rewritten as

$$\begin{aligned} & \frac{1}{n} (\partial_{ij} \rho) d\mathcal{M}((g_i - \bar{g}_i)(g_j - \bar{g}_j) - G_{ij}) \\ & + \frac{1}{n} (\partial_{ii} \rho) d\mathcal{M}((g_i - \bar{g}_i)(g_i - \bar{g}_i) - G_{ii}) \\ & + \frac{1}{n} (\partial_{jj} \rho) d\mathcal{M}((g_j - \bar{g}_j)(g_j - \bar{g}_j) - G_{jj}) \\ & = \frac{1}{n} d\mathcal{M} \left((\partial_{ij} \rho)((g_i - \bar{g}_i)(g_j - \bar{g}_j) - G_{ij}) \right. \\ & \quad \left. + (\partial_{ii} \rho)((g_i - \bar{g}_i)(g_i - \bar{g}_i) - G_{ii}) \right. \\ & \quad \left. + (\partial_{jj} \rho)((g_j - \bar{g}_j)(g_j - \bar{g}_j) - G_{jj}) \right). \end{aligned} \quad (33)$$

Writing the argument to $d\mathcal{M}$ on the right-hand-side of equation (33) as H_{ij} , the scaling property shows the stochastic component of K can be written as

$$\frac{1}{n} \|H_{ij}\| d\mathcal{M}(\hat{H}_{ij}). \quad (34)$$

Calculation of $\|H_{ij}\|$ only requires computation of the first order derivatives $\partial_{ij}\rho, \partial_{ii}\rho, \partial_{jj}\rho$ and the fact from multivariate normal distributions that

$$\begin{aligned} & \int_{\mathbb{R}^d} (g_i - \bar{g}_i)(g_j - \bar{g}_j)(g_k - \bar{g}_k)(g_l - \bar{g}_l) \tilde{p}(g) dg \\ & = G_{ij} G_{kl} + G_{ik} G_{jl} + G_{il} G_{jk}. \end{aligned} \quad (35)$$

Working through these calculations provides

$$K = \sqrt{\frac{v}{n}} (1 - \rho^2) dB_\rho, \quad (36)$$

where $B_\rho := \mathcal{M}(\hat{H}_{ij})$ is a scalar-valued standard Brownian motion.

To compute L , I first rewrite the product $(dG_{ij})(dG_{kl})$ using the notation introduced in the *MG* version of the framework above (i.e., using equations (26)), which provides

$$\begin{aligned} (dG_{ij})(dG_{kl}) &= \frac{1}{n^2} d\mathcal{M}[(g_i - \bar{g}_i)(g_j - \bar{g}_j) - G_{ij}] \\ & \times d\mathcal{M}[(g_k - \bar{g}_k)(g_l - \bar{g}_l) - G_{kl}]. \end{aligned} \quad (37)$$

Applying the multiplicative property for $d\mathcal{M}$, the product simplifies to

$$\begin{aligned} (dG_{ij})(dG_{kl}) &= \frac{1}{n^2} [\langle (g_i - \bar{g}_i)(g_j - \bar{g}_j), (g_k - \bar{g}_k)(g_l - \bar{g}_l) \rangle \\ & \quad - v n G_{ij} G_{kl}] dt, \end{aligned} \quad (38)$$

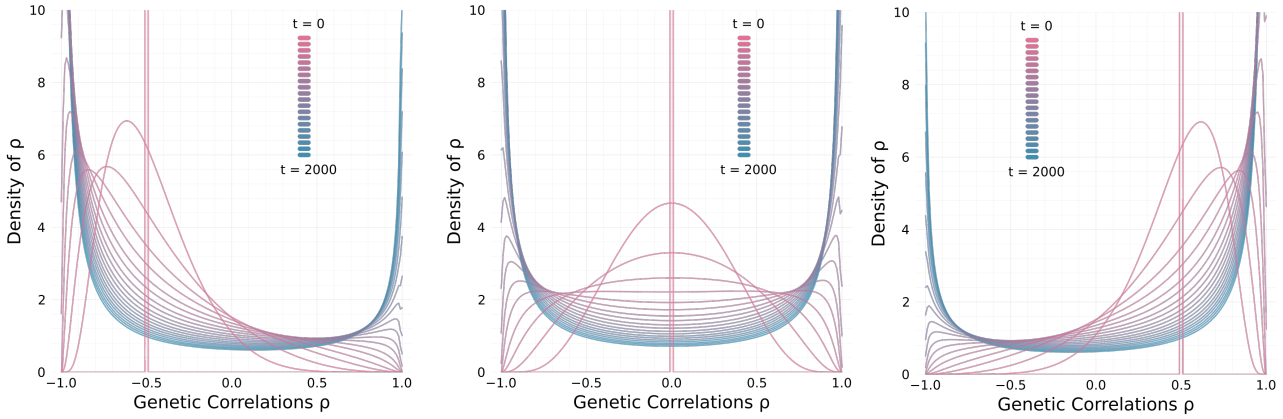


Fig. 1. The distribution of genetic correlations evolving under drift alone converges to a non-integrable stationary solution. Shown here are dynamics for initial distributions taking approximate point masses at $\rho_0 = -0.5$ (left panel), $\rho_0 = 0.0$ (middle panel), and $\rho_0 = 0.5$ (right panel). Solutions at earlier times are colored green, and later times are purple which run until $t = 2000$. The rate of drift is set to $v/n = 0.001$.

where I made use of $\langle (g_i - \bar{g}_i)(g_j - \bar{g}_j), G_{kl} \rangle = v n G_{ij} G_{kl}$. Again applying property (35) from above, I obtain the further simplification $(dG_{ij})(dG_{kl}) = \frac{v}{n}(G_{ik}G_{jl} + G_{il}G_{jk})$. Interfacing this result with derivatives of $\rho(G_{ij}, G_{ii}, G_{jj})$, I arrive at

$$L = -\frac{1}{2} \frac{v}{n} \rho(1 - \rho^2) dt. \quad (39)$$

Hence, by applying the MG version of the framework, I find that trait correlations driven entirely by drift follow the ordinary stochastic differential equation

$$d\rho = -\frac{1}{2} \frac{v}{n} \rho(1 - \rho^2) dt + \sqrt{\frac{v}{n}}(1 - \rho^2) dB_\rho. \quad (40)$$

Equation (40) shows that the effect of drift is mediated by the ratio of the variance of lifetime reproductive output v (typically set to $v = 1$ in classical quantitative genetic models such as Lande, 1976) to the effective population size n . Additionally, one can check that (because mutation and selection are absent) $\rho = \pm 1$ are stationary points. Furthermore, leveraging the fact that equation (40) defines a one-dimensional diffusion, we can in principle solve for its stationary distribution $\hat{q}(\rho)$ (Karlin and Taylor, 1981; Etheridge, 2010), which is done in Supplementary Material, Section 6.2. However, in attempt to do so we arrive at the non-integrable function:

$$\hat{q}(\rho) \propto \frac{1}{(1 - \rho^2)^{3/2}}. \quad (41)$$

The lack of a formal stationary distribution creates a challenge for understanding the limiting behavior of genetic correlations evolving solely in response to drift. This can be partially overcome by again leveraging the theory of one-dimensional diffusions to demonstrate that the boundaries ± 1 are both attracting and unattainable (Karlin and Taylor, 1981). That is, genetic correlations tend towards their extremes, but never fix at ± 1 . This is visualized in Figure 1 by plotting the distribution $q(\rho)$ of genetic correlations as numerical solutions to the forwards Kolmogorov (i.e., Fokker-Planck) equation associated with (40). Biologically, $|\rho|$ increases because drift randomly samples finite numbers of individuals, thereby causing transient correlations among the additive genetic values

these individuals carry. Further biological implications are discussed in the following section.

Another way to view this result is by applying Itô's formula to $u = \tanh^{-1}(\rho)$ (detailed calculations given in Supplementary Material, Section 6.3). Doing so returns

$$du = \frac{1}{2} \frac{v}{n} \tanh(u) dt + \sqrt{\frac{v}{n}} dB_u. \quad (42)$$

Because $\rho = \tanh(u)$ is a monotone increasing function of u , statements about u map directly to ρ . In particular, $u = \rho = 0$ is unstable because $\tanh(u)$ is positive for positive u , and negative for negative u . Additionally, if u is much greater than 1, then $du \approx v dt/2n + \sqrt{v/n} dB_u$ and, writing this point as u_0 , we have $\mathbb{E}[u_t] \approx u_0 + v t/2n$. The analogous approximation holds when u is much less than -1 . Justification for these claims is made in Supplementary Material, Section 6.3. This agrees with the boundary classification result above that demonstrates drift has an overall tendency to drive trait correlations towards ± 1 . Numerical estimates for sample paths of the solution to (40), illustrated by Figure 2, support this conclusion.

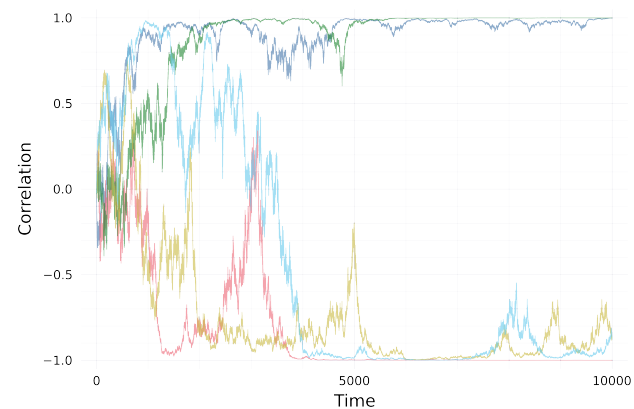


Fig. 2. Drift drives trait correlations towards ± 1 . Shown here are five replicates illustrating the path-behavior of trait correlations following equation (40). Each replicate is initiated with $\rho_0 = 0$ and the rate of drift is set to $v/n = 0.001$.

1004 To confirm the heuristics return correct expressions, I also
 1005 simulated the evolution of the \mathbf{G} -matrix driven solely by
 1006 random genetic drift for a 2-dimensional trait by applying
 1007 the Euler-Maruyama method (Bayram et al., 2018) to the
 1008 BG version of the framework. I then back-calculated the
 1009 genetic correlation ρ based on the simulated time-series of \mathbf{G} .
 1010 Figure 1 in Supplementary Material, Section 1 demonstrates
 1011 this approach agrees with equation (40). Further information
 1012 on the numerical implementation is given in Supplementary
 1013 Material, Section 1, and associated code is available at the
 1014 github repository github.com/bobweek/multi-mtgc.

1015 3.2. Evolution of The \mathbf{G} -Matrix in Response to Drift

1016 The above result demonstrates that drift increases correlations
 1017 between traits, at least for asexually reproducing populations
 1018 where recombination is absent. This perspective offers
 1019 an important refinement of the conventional wisdom that
 1020 \mathbf{G} -matrices responding to drift should merely contract
 1021 proportionally (Lande, 1979; Phillips and McGuigan, 2006;
 1022 Dugand et al., 2021; Mallard et al., 2023a). In particular, it is
 1023 classically thought that the orientation of the \mathbf{G} -matrix should
 1024 not change due to drift on average, and hence any changes
 1025 in orientation may be a sign of selection (Roff, 2000; Steppan
 1026 et al., 2002; Cano et al., 2004; Chapuis et al., 2008; Aguirre
 1027 et al., 2013).

1028 This idea can be obtained from equation (21) if the
 1029 stochastic component is neglected. Then, the deterministic
 1030 response due to drift is described by $\dot{\mathbf{G}} = -v\mathbf{G}/n$, which
 1031 has the solution $\mathbf{G}_t = \mathbf{G}_0 e^{-vt/n}$, and is a continuous-
 1032 time equivalent of the result found on page 409 in a paper
 1033 by Lande (1979). Numerical results displayed in Figure 2 of
 1034 Supplementary Material, Section 1 indicate agreement with this
 1035 classical scaling result in expectation. However, when stochastic
 1036 fluctuations dominate the dynamics over the timescale of
 1037 interest (so that individual sample paths diverge strongly
 1038 despite a smooth deterministic trend) the average response
 1039 to drift may provide little information for within-population
 1040 dynamics.

1041 Taking an experimental approach, Phillips et al., (2001)
 1042 established isolated populations of *Drosophila melanogaster*
 1043 from a common base population and found significant variation
 1044 across replicates for the response of \mathbf{G} to drift. This occurs
 1045 in spite of the fact that *Drosophila melanogaster* is a
 1046 sexually reproducing organism, and recombination should
 1047 act to break up correlations caused by linkage (discussed
 1048 further below) and thereby reduce variation across replicates.
 1049 Figure 3 emphasizes these variable outcomes theoretically by
 1050 superimposing expected trait correlations over a collection
 1051 of individual outcomes. The expected correlations are
 1052 approximated by averaging over replicates, but it took a very
 1053 large number of replicates (>1000) to obtain a satisfactory
 1054 deterministic trend. Because population size enters only
 1055 through the ratio v/n , increasing n rescales the time axis but
 1056 does not reduce pathwise variability or improve the predictive
 1057 power of the average trend. The averages shift towards zero,
 1058 in the sense of a slow directional trend that is not resolved to
 1059 convergence over the finite time window shown, but we can see
 1060 individual replicates are not predicted by this trend. Hence, to
 1061 gain a more accurate picture of \mathbf{G} -matrix evolution, there is a
 1062 need to understand the path behavior of individual outcomes.

1063 One approach to gain insight for the trait correlation of
 1064 a given replicate, as opposed to the average, is to analyze
 1065 the proportion of time that correlations spend near ± 1 over

1066 the total time the process is running. Figure 4 plots the
 1067 proportion of time that the magnitude of the trait correlation
 1068 ρ spends above the threshold $1 - \varepsilon$, i.e. with $|\rho| > 1 - \varepsilon$,
 1069 averaged over 10 replicate trajectories with initial value $\rho_0 = 0$
 1070 and $v/n = 0.001$, for $\varepsilon \in \{0.20, 0.10, 0.05, 0.02\}$. Smaller ε
 1071 corresponds to increasingly strict neighborhoods of ± 1 . This
 1072 result shows that trait correlations typically aggregate near ± 1
 1073 in drifting, isolated, asexual populations with no mutational
 1074 input. Hence, interpreting this as a signature of drift, we can
 1075 say that if trait correlations in a population exhibit significant
 1076 variation away from ± 1 , then either the population has been
 1077 drifting for only a short time relative to its effective population
 1078 size, or other processes not captured by this model must be
 1079 at play, such as ongoing mutation or recombination, which
 1080 are explicitly excluded here. I therefore discuss mechanisms
 1081 maintaining genetic correlations before concluding this section.

1082 Pleiotropic loci and genetic linkage are two mechanisms that
 1083 maintain genetic correlations (Lande, 1980). In this framework,
 1084 pleiotropic mutations would arise from non-zero off-diagonals of
 1085 the mutation matrix \mathbf{M} . In contrast, drift facilitates chance
 1086 correlations of additive genetic values between traits across
 1087 individuals. Hence, this second mode of genetic correlation
 1088 accumulation is directly analogous to the build-up of linkage
 1089 by drift (Hill and Robertson, 1966; Ohta and Kimura, 1969;
 1090 Lucek and Willi, 2021). For sexually reproducing populations,
 1091 it is expected that genetic correlations should be maintained
 1092 by pleiotropic loci as recombination breaks up linkage (Lande,
 1093 1980; Jones et al., 2003; Phillips and McGuigan, 2006).
 1094 However, the results in this section show that drift alone drives
 1095 genetic correlations toward extreme values. This raises the
 1096 natural question of how this tendency would be counteracted
 1097 by recombination, and whether a drift-recombination balance
 1098 analogous to that studied in classical population genetics (e.g.,
 1099 Ohta and Kimura, 1969) would emerge.

1100 The drift-only example considered here is intended as
 1101 an illustrative application of the framework rather than a
 1102 quantitative fit to existing experiments. In a clonal, mutation-
 1103 free setting, the model predicts that the stochastic dynamics
 1104 of genetic correlations depend on time only through the
 1105 rescaled variable $\tau = (v/n)t$. Equivalently, for populations
 1106 with different effective sizes n , the distributions of correlation
 1107 trajectories become comparable after rescaling time by n/v ,
 1108 so that increasing population size slows the dynamics without
 1109 altering their qualitative path behavior. This prediction can
 1110 be tested empirically using pathwise statistics such as the
 1111 distribution of first-passage times $\tau_c = \inf\{t : |\rho(t)| > c\}$ to
 1112 a high-correlation threshold c , or the long-run fraction of time
 1113 that correlations satisfy $|\rho| > c$; both quantities are predicted
 1114 to scale with n/v . Such tests would, in principle, be feasible
 1115 in replicated clonal systems with controlled demography and
 1116 minimal mutational input, including bottlenecked microbial
 1117 populations (e.g., *Escherichia coli* or asexually propagated
 1118 *Saccharomyces cerevisiae*) or clonal macrophytes such as
 1119 duckweed (*Lemna* or *Spirodela*) maintained in controlled
 1120 microcosms. Extending the framework to incorporate sexual
 1121 reproduction and recombination would be a natural next step
 1122 for making quantitative predictions about drift-recombination
 1123 balance. Nevertheless, the present results highlight that
 1124 focusing solely on average, deterministic responses can obscure
 1125 substantial pathwise structure, suggesting that conventional
 1126 perspectives on \mathbf{G} -matrix evolution merit closer examination.

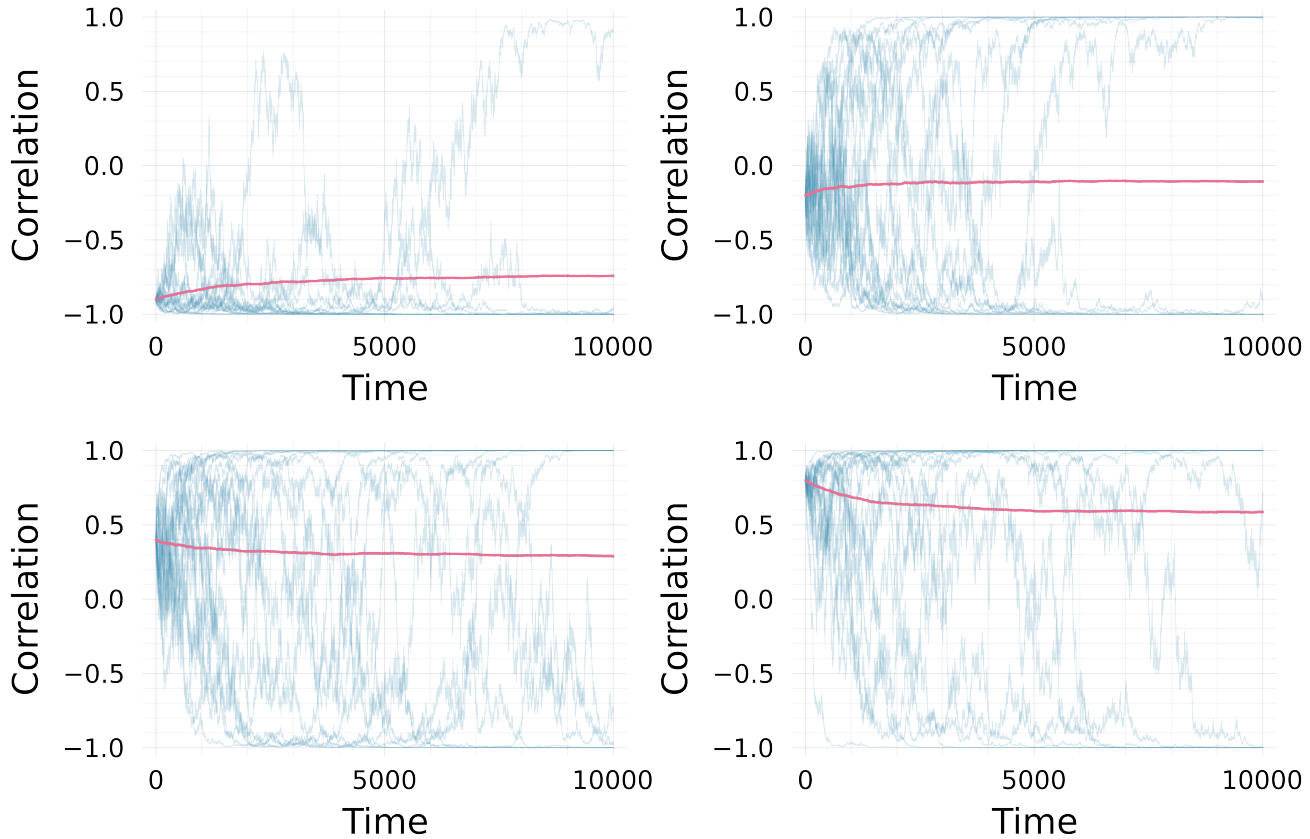


Fig. 3. Averages over replicated time-series of trait correlation do not predict within population trait correlations. Averages were computed using 3000 replicates with initial correlations $\rho_0 = -0.9$ (top-left), $\rho_0 = -0.2$ (top-right), $\rho_0 = 0.4$ (bottom-left), and $\rho_0 = 0.8$ (bottom-right). For the sake of clarity, just 20 random replicates are shown behind the average trends.

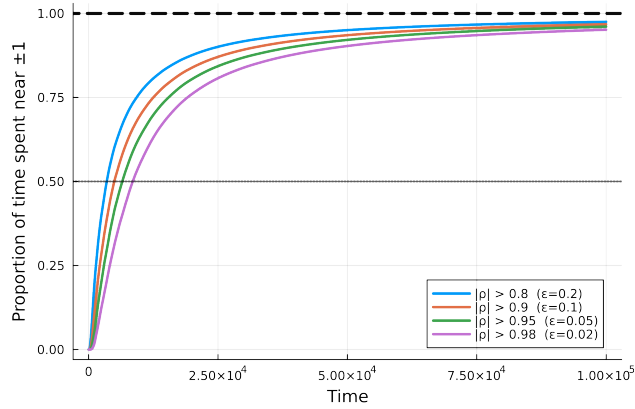


Fig. 4. Trait correlations (ρ) within populations rapidly evolve towards ± 1 under random genetic drift. Shown is the proportion of time that $|\rho|$ spends above a threshold $1 - \varepsilon$, averaged over 10 replicate trajectories, with $\varepsilon \in \{0.20, 0.10, 0.05, 0.02\}$. Smaller ε corresponds to increasingly strict neighborhoods of ± 1 . Simulations start from $\rho_0 = 0$ with $v/n = 0.001$.

particular emphasis on the use of multi-locus models (Bulmer, 1972; Chakraborty and Nei, 1982; Slatkin, 1987; Turelli and Barton, 1994; Barton and Turelli, 2004; Barton et al., 2017; Walsh and Lynch, 2018). The advantage of this approach is its clear connection with explicit genetic details. However, its primary challenge is the manipulation of complex expressions that emerge at such level of detail. To overcome this, and establish a flexible tool for modelling the integrated ecological and evolutionary dynamics of populations carrying multivariate traits, I struck a balance between incorporating genetic detail and analytical tractability. As a consequence, this framework has potential for wide-spread use across topics in evolution, ecology, and population biology.

A central feature of this framework is that a wide range of classical models can be obtained by choosing an appropriate fitness function m . For example, stochastic Lotka-Volterra dynamics for a community of species is obtained from this framework by applying the growth rate $m_i = r_i + \sum_j \alpha_{ij} n_j$ for species i , with n_j being the abundance of species j and α_{ij} the interaction coefficient. The long time behavior of the resulting stochastic Lotka-Volterra process for two species has been studied by Cattiaux and Méléard (2009). In addition, by making the interaction coefficients α_{ij} dependent on multivariate trait values \mathbf{z}_i , \mathbf{z}_j , models integrating coevolution with abundance feedbacks can be obtained similar to those studied by Gokhale et al., (2013); Cortez and Weitz (2014); and Patel et al., (2018). As a special case,

4. Discussion

The classical quantitative genetic approach has stressed the importance of considering the genetic architecture of traits for understanding evolutionary processes, and has placed

continuous time analogs of multivariate coevolution models are obtained by focusing on two species with fixed (or infinite) abundances, and assuming $\alpha_{ij}(\mathbf{z}_i, \mathbf{z}_j)$ depends on the Euclidean distance between \mathbf{z}_i and \mathbf{z}_j (Gilman et al., 2012; Débarre et al., 2014). Klausmeier et al. (2020) studied models of evolutionary rescue with univariate traits using growth rates of the forms $m(z, t) = r - \psi(\theta(t) - z)^2/2$ and $m(z, t) = r + r_0 e^{-\psi(\theta(t)-z)^2/2}$, where $\theta(t)$ is a dynamic phenotypic optimum and ψ is the strength of stabilizing selection. Applying multivariate generalizations of these growth rates to the above framework leads to extensions of an evolutionary rescue model involving demographic stochasticity studied by Xu et al. (2023). Additionally, Jones et al. (2012) also studied a model of phenotypic adaptation to a dynamic optimum, but in the context of **G**-matrix evolution. This framework can then be used as a bridge between research topics such as evolutionary rescue and **G**-matrix evolution. This list provides a small set of examples for how this framework can be used to derive new models across a broad range of topics in ecology, evolution, and population biology.

Further work is needed to extend this framework in several directions. For instance, it is possible to incorporate sexual reproduction and recombination by assuming each trait value is determined by the sum of additive genetic effects contributed by two gametes, rather than by a single genetic value. In this case, each gamete contributes an additive genetic effect g_i , and the trait is encoded as $g_i + g'_i$, where g_i and g'_i arise from convex combinations of the parental gametic values, similar to the approach taken by Lande (1980). How to formalize this extension using measure-valued processes is not obvious. In another direction, environmental stochasticity (i.e., stochastic growth rates) can be formally incorporated in a measure-valued context following the work of Mytnik (1996). Extending the framework in this direction may yield important additional insights complementing the work of Lande (2007; 2008), who studied the consequences of environmental stochasticity for long-run population growth rates. An extension in this direction may also be useful for studies investigating the consequences of environmental stochasticity on **G**-matrix evolution (e.g., Engen and Sæther, 2024).

I also point to two possible extensions that concern the role of development. The first accounts for the evolution of the mutational variance-covariance matrix **M** within the classical quantitative genetic framework. Allowing **M** to evolve enables analytical and simulation models for the evolution of evolvability, as studied by Jones et al. (2007), including numerical integration of the resulting stochastic differential equations (e.g. via Euler-Maruyama), rather than relying exclusively on individual-based models, and may yield more precise quantitative predictions amenable to experimental study (Mallard et al., 2023a). A second, conceptually distinct extension would be to replace the linear, regression-based genotype-phenotype map and residual model used here with nonlinear or mechanistic developmental maps, as in evo-devo frameworks (e.g., Milocco and Salazar-Ciudad, 2022; González-Forero, 2022, 2024b,a), in which genetic and phenotypic covariation emerge from explicit developmental dynamics rather than being parameterized through a mutational covariance matrix.

The application of this framework to study **G**-matrix evolution uncovered a more nuanced picture for the role of drift than what conventional wisdom suggests (Phillips and McGuigan, 2006). In particular, although drift indeed proportionally reduces **G**-matrices when averaged over many

replicated populations, it also drives trait correlations towards their extremes within populations. This agrees with the result that the expected eigenvalues of **G** differ from the eigenvalues of the expectation of **G** under drift (Griswold et al., 2007). In addition, the impact of drift on genetic correlations can be understood as a reduction in the effective dimensionality of **G**-matrices (Hine and Blows, 2006), reflecting an extension of the principle that drift erodes heritable variation. That is, because drift drives pairwise correlations to ± 1 , the distribution of multivariate traits in a population that has been evolving solely under drift may be well approximated after projecting onto a lower-dimensional trait space. Conceptually, this is similar to dimensionality reduction techniques such as principal components analysis (Kirkpatrick and Meyer, 2004). Importantly, this effect of drift on trait correlations calls into question comparative quantitative genetic methods utilizing **G**-matrices which routinely treat differences in **G**-orientation as a signal of selection in diverged populations (Roff, 2000; Steppan et al., 2002; Phillips and McGuigan, 2006; Cano et al., 2004; McGuigan, 2006; Chapuis et al., 2008; Aguirre et al., 2013; Dugand et al., 2021; Mallard et al., 2023a). These findings have broad implications for evolutionary biology, highlighting the need for revised theoretical perspectives and further empirical validation.

The study of genetic correlations and **G**-matrix dynamics using this framework can be extended by studying equation (21) with mutation and selection. In the special case of one-dimensional traits, stabilizing selection, and no abundance dynamics, the theory of one-dimensional diffusions (Etheridge, 2010) can be used to show that the stationary distribution of additive genetic variance follows a generalized inverse Gaussian distribution (Jørgensen, 2012). This suggests the stationary distribution of **G** may follow a matrix-variate generalization, such as the matrix generalized inverse Gaussian (MGIG) distribution (Fazayeli and Banerjee, 2016). Proposed distributions may be checked by evaluating the forwards Kolmogorov equation associated with equation (21) at equilibrium assuming a solution that follows the density of the proposal. Identification of the stationary distribution for (21) may then be used to study the distribution of genetic correlations maintained by interactions between mutation, selection, and random genetic drift.

In summary, this work introduces a versatile framework for modeling the stochastic eco-evolutionary dynamics of multivariate traits, providing a unifying approach that integrates mutation, selection, demographic stochasticity, and drift. By balancing mathematical rigor with accessibility, this framework enables the derivation of new models across a broad spectrum of population biology, making it a valuable tool for both theoretical and applied researchers. With its broad applicability, the framework presented here offers a foundation for future studies investigating the dynamics of populations in both theoretical and empirical contexts.

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