Imperfect strategy transmission can reverse the role of population viscosity on the evolution of altruism

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Abstract

Population viscosity, *i.e.*, low emigration out of the natal deme, leads to high within-deme relatedness, which is beneficial to the evolution of altruistic behavior when social interactions take place among deme-mates. However, a detrimental side-effect of low emigration is the increase in competition among related individuals. The evolution of altruism depends on the balance between these opposite effects. This balance is already known to be affected by details of the life-cycle; we show here that it further depends on the fidelity of strategy transmission from parents to their offspring. We consider different life-cycles and identify thresholds of parent-offspring strategy transmission inaccuracy, above which higher emigration can increase the frequency of altruists maintained in the population. EXPLAIN RESULT Predictions were first obtained analytically assuming weak selection and equal deme sizes, then confirmed with stochastic simulations relaxing these assumptions. This result challenges the notion that the evolution of altruism REMOVE REQUIRE requires limited dispersal.

6 Introduction

In his pioneering work on the evolution of social behavior, Hamilton suggested that altruistic behavior would be associated to limited dispersal (Hamilton, 1964, p. 10). This notion, that tighter links between individuals are beneficial to the evolution of altruism, has been shown to hold in a number of population structures (see *e.g.* Ohtsuki et al., 2006; Taylor et al., 2007a; Lehmann et al., 2007; Allen et al., 2017). The rationale is that altruism is favored when altruists interact more with altruists than defectors do (Hamilton, 1975, p. 141; Fletcher & Doebeli, 2009), a condition that is met in viscous populations, *i.e.*, populations with limited dispersal.

Yet, living next to your kin also implies competing against them (West et al., 2002), which is detrimental to the evolution of altruism. The evolution of so-

cial traits hence depends on the balance between the positive effects of interactions with related individuals and the detrimental consequences of kin com-29 petition. Under specific conditions, the two effects can even compensate each other, thereby annihilating the impact of population viscosity on the evolution of altruism. First identified with computer simulations (Wilson et al., 1992), this cancellation result was analyzed by Taylor (1992a) in a model with synchronous generations (i.e., Wright-Fisher model) and a subdivided population of constant, infinite size. The cancellation result was later extended to heterogeneous populations (Rodrigues & Gardner, 2012, with synchronous generations and infinite population size), and other life-cycles, with generic regular population structures (Taylor et al., 2011, with synchronous generations but also with continuous generations and Birth-Death updating). However, small changes in the model's assumptions, such as overlapping generations (Taylor & Irwin, 2000) or the presence of empty sites (Alizon & Taylor, 2008) can tip the balance in the favor of altruism. This high dependence on life-cycle specificities highlights the difficulty of making general statements about the role of spatial structure on the evolution

of altruism. In this study, we will consider three different life-cycles: Wright-Fisher, where the whole population is renewed at each time step, and two Moran life-cycles (Birth-Death and Death-Birth), where a single individual dies and is replaced at each time step. These life-cycles are classically used in studies on altruism in structured populations. Even though they differ by seemingly minor details, they are known to have very different outcomes in models with perfect parent-offspring transmission (*e.g.*, Taylor, 1992a; Rousset, 2004; Ohtsuki et al., 2006; Lehmann et al., 2007; Taylor, 2010).

A large number of studies on the evolution of social behavior consider simple 52 population structures (typically, homogeneous populations sensu Taylor et al. 53 (2007a)) and often also infinite population sizes (but see Allen et al., 2017, for results on any structure). These studies also make use of weak selection approximations, and commonly assume rare (e.g., Leturque & Rousset, 2002; Taylor et al., 2007b; Tarnita & Taylor, 2014) or absent mutation (for models assuming 57 infinite population sizes, or models concentrating on fixation probabilities; see Lehmann & Rousset, 2014; Van Cleve, 2015, for recent reviews). These simplifying assumptions are often a necessary step towards obtaining explicit analytical results. Simple population structures (e.g., regular graphs, or subdivided populations with demes of equal sizes) help reduce the dimensionality of the system under study, in particular when the structure of the population displays symmetries such that all sites behave the same way in expectation. Weak selection approximations are crucial for disentangling spatial moments (Lion, 2016), that is, changes in global vs. local frequencies (though they can in some cases be relaxed, as in Mullon & Lehmann, 2014). Mutation, however, is usually ignored by 67 classical models of inclusive fitness because these models assume infinite population sizes, so that there is no need to add mechanisms that restore genetic diversity (Tarnita & Taylor, 2014). In populations of finite size, this diversifying effect can be obtained thanks to mutation.

When strategy transmission is purely genetic, it makes sense to assume that 72 mutation is relatively infrequent. Even in this case, though, mutations from "social" to "non-social" types cannot always be neglected. For instance, experiments with the bacteria Pseudomonas fluorescens have identified transitions between populations dominated by the ancestral "solitary" Smooth Morph type and mat-forming "social" Wrinkly Spreaders, that can be re-invaded by Smooth Morphs not contributing to the formation of the mat (hence described as "cheaters"). The transitions between the different types are due to spontaneous mutations occurring over the timescale of the experiment (Hammerschmidt et al., 2014). In addition to genetic transmission, a social strategy can also be culturally trans-81 mitted from parent to offspring. In this case, "rebellion" (as in Frank's Rebellious Child Model (Frank, 1997)), i.e., adopting a social strategy different from one's parents, does not have to be infrequent. Since it is known that imperfect strategy transmission can alter the evolutionary dynamics of social traits, in particular in spatially structured populations (see e.g., Allen et al., 2012; Débarre, 2017, for graph-structured populations), it is therefore important to understand the impact of imperfect strategy transmission on the evolution of social behavior. Here, we want to explore the consequences of imperfect strategy transmission from parents to their offspring on the evolution of altruistic behavior in subdivided populations¹. The question was tackled by Frank (1997), but with a non 91 "fully dynamic model" (Frank, 1997, legend of Fig.7). Relatedness was treated like a parameter, which precluded the exploration of the effects of population viscosity on the evolution altruism. For each of the three life-cycles that we consider, we compute the expected 95 (i.e., long-term) frequency of altruists maintained in a subdivided population, and investigate how this frequency is affected by mutation and emigration. We find that, contrary to what happens with perfect strategy transmission, higher $^{
m 1}$ Note that for the sake of concision, we use the word "mutation" throughout the paper, keeping

¹Note that for the sake of concision, we use the word "mutation" throughout the paper, keeping in mind that strategy transmission does not have to be genetic.

emigration can increase the expected frequency of altruists in the population.

Model and methods

101 Assumptions

We consider a population of total size N, subdivided into N_D demes connected 102 by dispersal, each deme hosting exactly n individuals (i.e., each deme contains 103 n sites, each of which is occupied by exactly one individual; $nN_D = N$). Each 104 site has a unique label $i, 1 \le i \le N$. There are two types of individuals in the 105 population, altruists and defectors. The type of the individual living at site i $(1 \le i \le N)$ is given by an indicator variable X_i , equal to 1 if the individual is an 107 altruist, and to 0 if it is a defector. The state of the entire population is given by 108 a vector $\mathbf{X} = \{X_i\}_{1 \le i \le N}$. For a given population state \mathbf{X} , the proportion of altruists is $\overline{X} = \sum_{i=1}^{N} X_i / N$. All symbols are summarized in table A1. 110 Reproduction is asexual. The offspring of altruists are altruists themselves 111 with probability $1-\mu_{1\to 0}$, and are defectors otherwise $(0 < \mu_{1\to 0} \le 1/2)$. Similarly, 112 the offspring of defectors are defectors with probability $1-\mu_{0\rightarrow1}$, and are altruists 113 otherwise $(0 < \mu_{0\rightarrow 1} \le 1/2)$. Our calculations will be simpler if we introduce the 114 following change of parameters: {eq:changemut}

$$v = \frac{\mu_{0 \to 1}}{\mu_{1 \to 0} + \mu_{0 \to 1}}$$
 (0 < v < 1), and (1a) {eq:nu}

$$\mu = \mu_{1 \to 0} + \mu_{0 \to 1} \quad (0 < \mu \le 1).$$
 (1b) {eq:mu}

The composite parameter v corresponds to the expected frequency of altruists in the population at the mutation-drift balance (*i.e.*, in the absence of selection; see Appendix A for details). We call v the "mutation bias" parameter. Parameter μ is the sum of the two mutation probabilities. In the absence of selection, at the mutation-drift equilibrium, the correlation between offspring type and their parent's type is $1-\mu$ (see Appendix A for details for the calculation). We call μ the

122 mutation intensity.

An individual of type X_k expresses a social phenotype $\phi_k = \delta X_k$, where δ is 123 assumed to be small ($\delta \ll 1$). Social interactions take place within each deme; a 124 focal individual interacts with its n-1 other deme-mates. We assume that social 125 interactions affect individual fecundity; f_k denotes the fecundity of the individ-126 ual at site k ($1 \le k \le N$), which depends on deme composition. We denote by b 127 the sum of the marginal effects of deme-mates' phenotypes on the fecundity of a 128 focal individual, and by -c the marginal effect of a focal individual's phenotype 129 on its own fecundity ($c \le b$; see system (A22) for formal definitions). 130

Offspring remain in the parental deme with probability 1-m and land on any site of the parental deme with equal probability (including the very site of their parent). With probability m, offspring emigrate to a different deme, chosen uniformly at random among the N_D-1 other demes. Denoting by d_{ij} the probability of moving from site i to site j, we have

$$d_{ij} = \begin{cases} d_{\text{in}} = \frac{1-m}{n} & \text{if sites } i \text{ and } j \text{ are in the same deme;} \\ d_{\text{out}} = \frac{m}{(N_D - 1)n} & \text{if they are in different demes,} \end{cases}$$
 (2) {eq:defD}

with $0 < m < 1 - \frac{1}{N_D}$. This upper bound is here to ensure that within-deme relatedness R, which will be defined later in the article, remains positive. When the emigration probability m is equal to the upper bound $1 - \frac{1}{N_D}$, the population is effectively well-mixed ($d_{\rm in} = d_{\rm out}$). We denote by $B_i = B_i(\mathbf{X}, \delta)$ the expected number of successful offspring of the

We denote by $B_i = B_i(\mathbf{X}, \delta)$ the expected number of successful offspring of the individual living at site i (successful means alive at the next time step), and by $D_i = D_i(\mathbf{X}, \delta)$ the probability that the individual living at site i dies. Both depend on the state of the population \mathbf{X} , but also on the way the population is updated from one time step to the next, i.e., on the chosen life-cycle (also called updating rule). We also define

$$W_i := (1 - \mu)B_i + 1 - D_i;$$
 (3) {eq:defW}

this is a particular definition of fitness, where the number of offspring produced (B_i) is scaled by the parent-offspring type correlation $(1 - \mu)$.

We will specifically explore three different life-cycles. At the beginning of
each step of each life-cycle, all individuals produce offspring, that can be mutated; then these juveniles move, within the parental deme or outside of it, and
land on a site. The next events occurring during the time step depend on the
life-cycle:

Moran Birth-Death: One of the newly created juveniles is chosen at random; it kills the adult who was living at the site, and replaces it; all other juveniles die.

Moran Death-Birth: One of the adults is chosen to die (uniformly at random among all adults). It is replaced by one of the juveniles who had landed in its site. All other juveniles die.

Wright-Fisher: All the adults die. At each site of the entire population, one of the juveniles that landed there is chosen and establishes at the site.

Previous studies have shown that, when social interactions affect fecundity, altruism is disfavored under the Moran Birth-Death and Wright-Fisher life-cycles, because the expected frequency of altruists under these life-cycles is lower than what it would be in the absence of selection (*e.g.*, Taylor, 1992a, 2010; Taylor et al., 2011; Débarre, 2017). However, we are interested in the actual value of the expected proportion of altruists in the population, not just whether it is higher or lower than the neutral expectation. This is why we are still considering the Moran Birth-Death and Wright-Fisher life-cycles in this study.

69 Methods

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170 Analytical part

truists are given in Appendix B. They go as follows: first, we write an equation for the expected frequency of altruists in the population at time t+1, conditional 173 on the composition of the population at time t; we then take the expectation of 174 this quantity and consider large times t. After this, we write a first order expan-175 sion for phenotypic differences δ close to 0 (this corresponds to a weak selection 176 approximation). 177 The formula involves quantities that can be identified as neutral probabil-178 ities of identity by descent Q_{ij} . These quantities correspond to the probability 179 that individuals living at site i and j share a common ancestor and that no muta-180 tion occurred on either lineage since that ancestor, in a model with no selection 181 $(\delta = 0)$ and with mutation intensity μ ; this is the "mutation definition" of identity by descent (Rousset & Billiard, 2000). In a subdivided population like the one we 183

consider, there are only three possible values of Q_{ij} :

The calculation steps to obtain the expected (i.e., long-term) proportion of al-

$$Q_{ij} = \begin{cases} 1 & \text{when } i = j, \\ Q_{\text{in}} & \text{when } i \neq j \text{ and both sites are in the same deme,} \\ Q_{\text{out}} & \text{when both sites are in different demes.} \end{cases}$$
 (4) {eq:Q3}

These neutral probabilities of identity by descent depend on the chosen lifecycle, and are also computed by taking the long-term expectation of conditional expectations after one time step (see Appendix C.1 and C.2 and supplementary Mathematica file (Wolfram Research, Inc., 2017).)

189 Stochastic simulations

190 To check our results and also relax some key assumptions, we ran stochastic sim-

ulations (coded in C). The simulations were run for 10⁸ generations (one gener-

ation is one time step for the Wright-Fisher life-cycle, and N time steps for the

Moran life-cycles). For each set of parameters and life-cycle, using R (R Core

Team, 2015), we estimated the long-term frequency of altruists by sampling the

population every 10³ generations and computing the average frequency of altru-

ists. All scripts are available at

https://flodebarre.github.io/SocEvolSubdivPop/_

change address

198 Results

199 Expected frequencies of altruists for each life-cycle

200 For each of the life-cycles that we consider, the expected frequency of altruists in

the population, $\mathbb{E}[\overline{X}]$, can be approximated as

$$\mathbb{E}\left[\overline{X}\right] \approx v + \frac{\delta}{\mu B^*} v(1-v)(1-Q_{\text{out}}) \times \left[\underbrace{\frac{\partial W}{\partial f_{\bullet}}(-c) + \frac{\partial W}{\partial f_{\text{in}}}b}_{-C} + \underbrace{\left(\frac{\partial W}{\partial f_{\bullet}}b + (n-1)\frac{\partial W}{\partial f_{\text{in}}}(-c) + (n-2)\frac{\partial W}{\partial f_{\text{in}}}b\right)}_{\mathcal{B}}\underbrace{\frac{Q_{\text{in}} - Q_{\text{out}}}{1 - Q_{\text{out}}}}_{R}\right], \quad (5) \quad \{\text{eq:EXapprox}\}$$

with W as defined in eq. (3). Calculations leading to eq. (5) are presented in

Appendix B; notations are recapitulated in table A1. In particular, B^* is the ex-

204 pected number of offspring produced by an adult, in the absence of selection

(when $\delta = 0$; $B^* = 1$ for the Wright-Fisher life-cycle and $B^* = 1/N$ for the Moran

206 life-cycles). Subscript "•" denotes a focal individual itself, and "in" a deme-mate.

Partial derivatives are evaluated for $\delta = 0$.

The expected frequency of altruists in the population is approximated, under

weak selection ($\delta \ll 1$), by the sum of what it would be in the absence of selec-

tion ($\mathbb{E}_0[\overline{X}] = \nu$, first term in eq. (5)), plus a deviation from this value, scaled by

 δ . The $-\mathcal{C}$ term corresponds to the effects of a change of a focal individual's phenotype on its own fitness (with the fitness definition given in eq. (3)). The \mathcal{B} term corresponds to the sum of the effects of the change of deme-mates' phenotypes on an individual's fitness. It is multiplied by R, which is relatedness.

The parametrization proposed in eq. (1) allows us to decouple the effects of the two new mutation parameters, ν and μ . The mutation bias ν , which was defined in eq. (1a), does not affect the sign of the second ("deviation") term in eq. (5); it only appears in the $\nu(1-\nu)$ product. The mutation intensity μ , however, affects the values of W, $Q_{\rm in}$ and $Q_{\rm out}$. The presence of μ at the denominator in eq. (5) may look ominous; however, both R and $(1-Q_{\rm out})/\mu$ have a finite limit when $\mu \to 0$.

The different terms depend on the chosen life-cycle. We first focus on related edness R.

224 **Relatedness** R

Within-deme relatedness depends on the number of individuals that are born 225 at each time step, and hence on the chosen life-cycle. In a Moran life-cycle (de-226 noted by M), one individual is updated at each time step, while under a Wright-227 Fisher life-cycle (denoted by WF), N individuals – the whole population – are up-228 dated at each time step. The formulas for relatedness for any number of demes N_D and mutation intensity μ are presented in Appendix C.2 (eq. (A44) and eq. (A50)). 230 When we let the number of demes go to infinity $(N_D \to \infty)$ and the intensity of 231 mutation be vanishingly small ($\mu \rightarrow 0$), we recover the classical formulas for re-232 latedness as limit cases (eq. (A45) and eq. (A51)). 233 The effects of emigration m and mutation intensity μ on relatedness are rep-234 resented in figure 1. For $0 < m < 1 - 1/N_D$, within-deme relatedness is positive, 235 and it decreases with m and with μ (the mutation bias v has no effect). The effect 236

of the mutation intensity μ on relatedness is strongest at low emigration prob-

abilities m. As m increases, the relatedness values for different mutation intensities get closer, until they all hit zero for $m = 1 - 1/N_D$ (which is the emigration probability such that an offspring is equally likely to land in its parent's deme or in any other deme, *i.e.*, such that there is no proper population subdivision anymore).

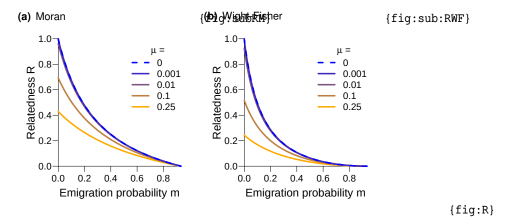


Figure 1: Within-deme relatedness of pairs of individuals R, as a function of the emigration probability m, for different values of the mutation probability μ (from 0 [blue] to 0.25 [orange]), and for the two types of life-cycles ((a): Moran, (b): Wright-Fisher). Other parameters: n=4 individuals per deme, $N_D=15$ demes.

43 Primary and secondary effects

We now turn to the \mathcal{B} and $-\mathcal{C}$ terms of eq. (5), which also depend on the chosen life-cycle. We further decompose these terms into primary (subscript P) and secondary (subscript S) effects (West & Gardner, 2010):

$$\mathcal{B} = \mathcal{B}_{P} + \mathcal{B}_{S},$$

$$-\mathcal{C} = \underbrace{-\mathcal{C}_{P}}_{Primary effect} + \underbrace{-\mathcal{C}_{S}}_{Secondary effect}$$
(6)

Primary effects correspond to unmediated consequences of interactions (they are included in $\frac{\partial W}{\partial f_{\bullet}}$). Secondary effects correspond to consequences of interactions mediated by other individuals, including competition.

50 Primary effects

251 Primary effects are the same for all the life-cycles that we consider:

$$\mathcal{B}_{\mathbf{P}}^{\mathrm{BD}} = \mathcal{B}_{\mathbf{P}}^{\mathrm{DB}} = \mathcal{B}_{\mathbf{P}}^{\mathrm{WF}} = (1 - \mu)\mathsf{b},\tag{7a}$$

$$-C_{\rm p}^{\rm BD} = -C_{\rm p}^{\rm DB} = -C_{\rm p}^{\rm WF} = (1 - \mu)(-c),$$
 (7b)

and they do not depend on the emigration probability m (see Appendix B.2 for details of the calculations).

As we have seen above, the relatedness terms $R^{\rm M}$ and $R^{\rm WF}$ decrease with m (keeping $m < 1 - 1/N_D$; see figure 1). Consequently, if we ignored secondary effects, we would conclude that the expected frequency of altruists in the population $\mathbb{E}[\overline{X}]$ decreases as the emigration probability m increases. However, secondary effects play a role as well.

259 Secondary effects

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obtain the following secondary effects:

Secondary effects take competition into account, that is, how the change in the 260 fecundity of an individual affects the fitness of another one. As shown already 261 in models with nearly perfect strategy transmission (Grafen & Archetti, 2008), 262 competition terms depend on the chosen life-cycle, because life-cycle details 263 affect the distance at which competitive effects are felt. Given the way the model 264 is formulated, $-C_S = B_S/(n-1)$ holds for all the life-cycles that we consider (see 265 Appendix B.2 for details of the calculations). 266 Under the Moran Birth-Death life-cycle, both the probability of reproducing 267 and the probability of dying depend on the composition of the population. We 268

$$-C_{\rm S}^{\rm BD} = \frac{\mathcal{B}_{\rm S}^{\rm BD}}{n-1} = -(\mathsf{b}-\mathsf{c})\left(-\frac{\mu}{N} + \frac{1-m}{n}\right). \tag{8a} \quad \{\mathsf{eq:BDsec}\}$$

{eq:secondary}

The competitive effects are the same for the Moran Death-Birth and Wright-

Fisher life-cycles. In both cases, the probabilities of dying are constant, so we can factor $(1 - \mu)$ in the equations:

$$-\mathcal{C}_{\rm S}^{\rm DB} = \frac{\mathcal{B}_{\rm S}^{\rm DB}}{n-1} = -\mathcal{C}_{\rm S}^{\rm WF} = \frac{\mathcal{B}_{\rm S}^{\rm WF}}{n-1} = -({\sf b}-{\sf c})(1-\mu) \left(\frac{(1-m)^2}{n} + \frac{m^2}{N-n}\right). \tag{8b} \quad \{\sf eq:DBsec}\}$$

These secondary effects (eq. (8a) and eq. (8b)) remain negative for the range 273 of emigration values that we consider $(0 < m < 1 - 1/N_D)$, and increase with m. In 274 other words, the intensity of competition decreases as emigration m increases. 275 While the value of these secondary effects increases with emigration m, re-276 latedness R, by which they are eventually multiplied in eq. (5), decreases with 277 m. We therefore cannot determine the overall effect of emigration m on the ex-278 pected frequency of altruists in the population by inspecting the different terms 279 of eq. (5) in isolation. For each life-cycle, we need to consider the entire equa-280 tions to know the overall effect of the emigration probability m on the expected 281 frequency of altruists $\mathbb{E}[\overline{X}]$ and on how it is affected by the (in)fidelity of parent-282 offspring transmission μ . 283

Changes of the expected frequency of altruists with the emigration probability m

The rather lengthy formulas that we obtain are relegated to the Appendix and supplementary Mathematica file, and we concentrate here on the results.

288 Moran Birth-Death

For the Moran Birth-Death life-cycle, we find that the expected frequency of altruists $\mathbb{E}[\overline{X}]$ is a monotonic function of the emigration probability m. The direction of the change depends on the value of the mutation probability μ compared to a threshold value μ_c^{BD} . When $\mu < \mu_c^{\mathrm{BD}}$, $\mathbb{E}[\overline{X}]$ decreases with m, while when

293 $\mu > \mu_c^{\mathrm{BD}}$, $\mathbb{E}[\overline{X}]$ increases with m. The critical value μ_c^{BD} is given by

$$\mu_c^{\rm BD} = 1 - \frac{b - c + \sqrt{(b - c)(4bN^2 + b - c)}}{2bN}$$
 (9) {eq:mucBD}

(recall that N is the total size of the population, $N=nN_D$.) This result is illustrated in figure 2(b); with the parameters of the figure, $\mu_c^{\rm BD} \approx 0.026$. The threshold value increases with both deme size n and number of demes N_D , up to a maximum value $1-\sqrt{1-c/b}$ (equal to 0.034 with our parameters.)

With this life-cycle however, the expected frequency of altruists $\mathbb{E}[\overline{X}]$ remains lower than v, its value in the absence of selection (*i.e.*, when $\delta=0$).

300 Moran Death-Birth

The relationship between $\mathbb{E}[\overline{X}]$ and m is a bit more complicated for the Moran Death-Birth life-cycle. For simplicity, we concentrate on what happens starting from low emigration probabilities (*i.e.*, on the sign of the slope of $\mathbb{E}[\overline{X}]$ as a function of m when $m \to 0$). If the benefits b provided by altruists are relatively low (b < c(n+1)), $\mathbb{E}[\overline{X}]$ initially increases with m provided the mutation probability μ is greater than a threshold value μ_c^{DB} given in eq. (10) below; otherwise, when the benefits are high enough, $\mathbb{E}[\overline{X}]$ initially increases with m for any value of μ . Combining these results, we write

$$\mu_c^{\rm DB} = \begin{cases} \frac{(n+1)c - b}{(2n-1)b - (n-1)c} & \text{if } b < c(n+1), \\ 0 & \text{otherwise.} \end{cases}$$
 (10) {eq:mucDB}

When b < c(n+1), the mutation threshold does not depend on the number of demes N_D , but increases with deme size n. In figure 2(a), the parameters are such that $\mu_c^{\mathrm{DB}}=0$.

When $\mu>\mu_c^{\mathrm{DB}}$, the expected frequency of altruists $\mathbb{E}[\overline{X}]$ reaches a maximum at an emigration probability m_c^{DB} (whose complicated equation is given in the

supplementary Mathematica file), as can be seen in figure 2(a). When the mutation probability gets close to 0 ($\mu \to 0$), $m_c^{\rm DB}$ also gets close to 0.

With the Death-Birth life-cycle, the expected frequency of altruists is higher than its neutral value v for intermediate values of the emigration probability m (unless $\mu \to 0$, in which case the lower bound tends to 0).

319 Wright-Fisher

Under a Wright-Fisher updating, the expected frequency of altruists in the population reaches an extremum at the highest admissible emigration value $m=1-\frac{1}{N_D}$. This extremum is a maximum when the mutation probability is higher than a threshold value $\mu_c^{\rm WF}$ given by

$$\mu_c^{\text{WF}} = 1 - \sqrt{1 - \frac{\mathsf{c}}{\mathsf{b}}},\tag{11}$$

and it is a minimum otherwise. With the parameters of figure 2(c), $\mu_c^{\rm WF}$ = 0.034.

With the Wright-Fisher life-cycle however, the expected frequency of altruists

remains below its value in the absence of selection, v.

Relaxing key assumptions

To derive our analytical results, we had to make a number of simplifying assumptions, such as the fact that selection is weak ($\delta \ll 1$), and the fact that the structure of the population is regular (all demes have the same size n). We checked with numerical simulations the robustness of our results when these key assumptions are relaxed.

Strong selection When selection is strong, the patterns that we identified not only still hold but are even more marked, as shown on figure A1.

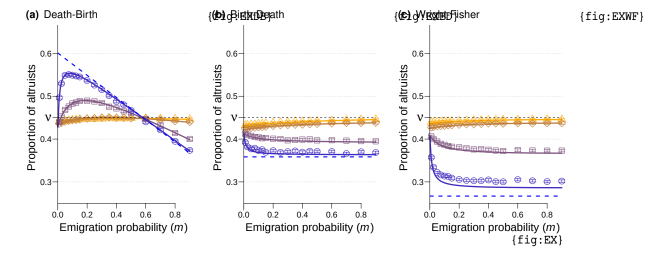


Figure 2: Expected proportion of altruists under weak selection, as a function of the emigration probability m, for different mutation values ($\mu=0.001$ (blue, dots), 0.01 (purple, squares), 0.1 (brown, diamonds), 0.25 (orange, triangles); the dashed blue lines correspond to $\mu=0$) and different life-cycles ((a) Moran Death-Birth, (b) Moran Birth Death, (c) Wright-Fisher). The curves are the analytical results, the points are the output of numerical simulations. Parameters: $\delta=0.005$, $\nu=0.45$, b=15, c=1, n=4 individuals per deme, $N_D=15$ demes.

Heterogeneity in deme sizes To relax the assumption of equal deme sizes, we randomly drew deme sizes at the beginning of simulations, with sizes ranging from 2 to 6 individuals and on average $\overline{n} = 4$ individuals per deme as previously. As shown in figure A2, the patterns initially obtained with a homogeneous population structure are robust when the structure is heterogeneous.

No self-replacement For the Moran model, it may seem odd that an offspring can replace its own parent (which can occur since $d_{ii} \neq 0$). Figure A3, plotted with dispersal probabilities preventing immediate replacement of one's own parent (for all sites i, $d_{ii} = d_{\text{self}} = 0$; $d_{\text{in}} = (1 - m)/(n - 1)$ for two different sites in the same deme, d_{out} remaining unchanged), confirms that this does affect our conclusions.

Infinite number of demes Our results are obtained in a population of finite size (the figures are drawn with $N_D=15$ demes), but still hold when the size of the population is larger. Figure 3(b) shows the range of emigration and mutation values such that altruism is favored, plotted also for $N_D \rightarrow \infty$.

Same graphs for dispersal and social interactions Compared to graphs classically used in evolutionary graph theory (e.g., regular random graphs, grids), the 351 island model is particular because the interaction graph and the dispersal graph 352 are different: interactions take place only within demes ($e_{out} = 0$), while offspring 353 can disperse out of their natal deme ($d_{\text{out}} > 0$). One may wonder whether our re-354 sult depends on this difference between the two graphs. Figure A4 shows that the 355 result still holds when the dispersal and interaction graphs are the same. In this 356 figure indeed, we let a proportion m (equal to the dispersal probability) of inter-357 actions occur outside of the deme where the individuals live, and set d_{self} , the 358 probability of self replacement, equal to 0, so that the dispersal and interactions 359 graphs are the same. Our conclusions remain unchanged.

61 Discussion

The expected frequency of altruists in a subdivided population can increase with the probability of emigration

Assuming that the transmission of a social strategy (being an altruist or a defec-364 tor) from a parent to its offspring could be imperfect, we found that the expected 365 frequency of altruists maintained in a population could increase with the prob-366 ability *m* of emigration out of the parental deme, a parameter tuning population 367 viscosity. This result can seem surprising, because it contradicts the conclusions 368 obtained under the assumption of nearly perfect strategy transmission (i.e., in 369 the case of genetic transmission, when mutation is very weak or absent). Under 370 nearly perfect strategy transmission indeed, increased population viscosity (i.e., 371 decreased emigration probability) is either neutral (Taylor, 1992a, and dashed 372 lines in figures 2(b)-(c)) or favorable (Taylor et al., 2007a, and dashed lines in 373 figure 2(a)) to the evolution of altruistic behavior. 374

Quantitative vs. qualitative measures

Often, evolutionary success is measured qualitatively, by comparing a quantity 376 (an expected frequency, or, in models with no mutation, a probability of fixa-377 tion) to the value it would have in the absence of selection. In our model, this 378 amounts to saying that altruism is favored whenever $\mathbb{E}[\overline{X}] > v$ (v is plotted as a 379 horizontal dashed line in figure 2). Some of our conclusions change if we use this 380 qualitative measure of evolutionary success: Under the Moran Birth-Death and 381 Wright-Fisher life-cycles, population viscosity does not promote the evolution of 382 altruism – actually, these two life-cycles cannot ever promote altruistic behavior 383 for any regular population structure (Taylor et al., 2011), whichever the probabil-384 ity of mutation (Débarre, 2017). However, under a Moran Death-Birth life-cycle 385 (figure 2(a)), altruism can be favored only at intermediate emigration probabil-

ities. Starting for initially low values of m, increasing the emigration probability can still favor the evolution of altruism under this qualitative criterion (see figure 3(b).)

Interpreting the effect of m on $\mathbb{E}[\overline{X}]$

To better understand the role played by the mutation intensity μ , we focus on the qualitative condition for the evolution of altruism ($\mathbb{E}[\overline{X}] > v$); and on the Death-Birth life-cycle, since this qualitative condition is not satisfied in the two other life-cycles. Having made sure that $\mathcal{B}^{\mathrm{DB}} > 0$ (as shown in the supplementary Mathematical file), the qualitative condition for altruism to be favored is given by

$$\mathbb{E}[\overline{X}] > \nu \Leftrightarrow R^{M} > \frac{\mathcal{C}^{DB}}{\mathcal{B}^{DB}}.$$
 (12) {eq:BCcond}

With the Death-Birth life-cycle, the $\mathcal{C}^{DB}/\mathcal{B}^{DB}$ ratio does not change with the mutation probability μ (the $(1-\mu)$ factors simplified out), but the ratio decreases 398 with the emigration probability m (with $0 < m < 1 - 1/N_D$; see the thick black 399 curve in figure 3(a)). This decrease of the C^{DB}/B^{DB} ratio is due to secondary 400 effects (competition) diminishing as emigration increases. Relatedness, on the 401 other hand, decreases with both μ and m (see figure 3(a)). We need to explain 402 the effect of the emigration probability m on condition (12) for different values 403 of mutation intensity μ . 404 When the emigration probability m is high, relatedness gets closer to zero 405 for all values of mutation intensity μ , while the $\mathcal{C}^{DB}/\mathcal{B}^{DB}$ remains positive; con-406

for all values of mutation intensity μ , while the $\mathcal{C}^{\mathrm{DB}}/\mathcal{B}^{\mathrm{DB}}$ remains positive; condition (12) is not satisfied. On the other hand, when the emigration probability m is vanishingly small, $\lim_{m\to 0} R^{\mathrm{M}} \leq \lim_{m\to 0} \frac{\mathcal{C}^{\mathrm{DB}}}{\mathcal{B}^{\mathrm{DB}}}$, the two only being equal when $\mu=0$. Hence, condition (12) is satisfied for vanishingly low m only when strategy transmission is perfect. Finally, as m increases to intermediate values, the $\frac{\mathcal{C}^{\mathrm{DB}}}{\mathcal{B}^{\mathrm{DB}}}$ ratio decreases with a steeper slope than relatedness R, so that the curves can cross provided the mutation probability μ is not too high, i.e., that R was not ini-

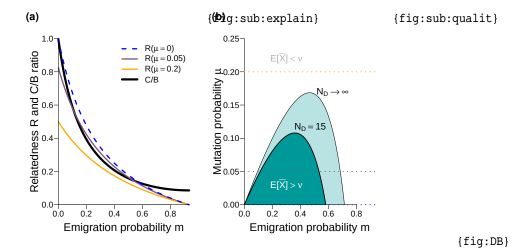


Figure 3: Understanding the effect of emigration m on whether altruism is favored in the Death-Birth life-cycle. (a) Comparison of the \mathcal{C}/\mathcal{B} ratio (thick black curve) and relatedness R (thin curves) for different values of the mutation probability μ (same color code as previously). (b) (m, μ) combinations for which $\mathbb{E}[\overline{X}] > v$. The dotted horizontal lines correspond to the mutation values used in panel (a). Unless specified, all other parameters are the same as in figure 2.

tially too low already. Hence, for no too high mutation intensity, there is a range of emigration values m such that condition (12) is satisfied.

The result is due to secondary effects

The result, that frequency of altruists can increase with the emigration probabil-416 ity m, may seem counterintuitive. It is the case because verbal explanations for the evolution of altruism often rely on primary effects only. Relatedness R de-418 creases with m, so it may be tempting to conclude that increases in the emigra-419 tion probability m are necessarily detrimental to the evolution of altruism. How-420 ever, secondary effects play an opposite role, as competition decreases with m, 421 and the effect is strongest at low values of m (see the black curve on figure 3(a); 422 in the absence of secondary effects, it would just be a horizontal line). 423 Secondary effects are less straightforward to understand than primary ef-424

fects, and yet they play a crucial role for social evolution in spatially structured

populations. Competition among relatives is for instance the reason for Taylor (1992b)'s cancellation result. Similarly, the qualitative differences between the Moran Birth-Death and Moran Death-Birth life-cycles is explained by the different scales of competition that the two life-cycle produce (Grafen & Archetti, 2008; Débarre et al., 2014). Secondary effects are also behind the evolution of social behaviors such as spite (West & Gardner, 2010).

432 Other model

433 How small is small and how large is large?

Our results were derived under the assumption of weak selection, assuming that 434 the phenotypic difference between altruists and defectors is small ($\delta \ll 1$). We 435 considered any fidelity of transmission (any μ between 0 and 1) and population 436 size. However, most models considering subdivided populations assume nearly 437 perfect strategy transmission ($\mu \rightarrow 0$) and infinite population sizes (number of 438 demes $N_D \to \infty$). The point is technical, but it is important to know that the or-439 der in which these limits are taken matters, i.e., one needs to specify how small 440 μ and δ are compared to the inverse size of the population 1/N. This remark complements findings by Sample & Allen (2017), who highlighted the quantitative differences between different orders of weak selection and large population limits.

Imperfect transmission and Rebellious Children

Our model bears resemblance to the Rebellious Child Model by Frank (1997), who studied the evolution of a vertically transmitted cultural trait in an asexually reproducing population. In Frank's model, however, relatedness r is treated as a fixed parameter (Frank, 1997, legend of Figure 7). Our model is mechanistic; relatedness r necessarily depends on the mutation probability μ , because probabilities of identity by descent do.

Mutation was also previously included in models investigating the maintenance of cooperative microorganisms in the presence of cheaters (Brockhurst et al., 2007; Frank, 2010). In both of these models however, only loss-of-function mutation was considered, which corresponds to setting the mutation bias at v = 0 in our model. This means that the all-cheaters state is absorbing; no matter how favored cooperators may otherwise be, in the long run, a finite population will only consist of cheaters.

459 Cultural transmission

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476

Strategy transmission does not have to be genetic: it can be cultural. In our model, strategy transmission occurs upon reproduction, so this is a case of vertical cultural transmission.

The model could nevertheless be interpreted as a representation of horizon-463 tal transmission, if we described reproduction as an instance of an individual 464 convincing another one to update its strategy. The Moran Death-Birth model 465 can be interpreted as a modified imitation scheme (Boyd & Richerson, 2002; Oht-466 suki et al., 2006) - with a specific function specifying who is imitated -, with mu-467 tation (Kandori et al., 1993). First, we choose uniformly at random an individual 468 who may change its strategy; with probability μ the individual chooses a random 469 strategy (altruistic with probability ν), and with probability $1-\mu$ it imitates another individual. Who is imitated depends on the distance to the focal individual 471 (with probability m it is a random individual in another deme) and on the "fe-472 cundities" of those individuals (as shown in table A2). With this interpretation of 473 the updating rule however, there is not reproduction nor death anymore. 474

It remains to be investigated how imperfect strategy transmission would affect the effect of population viscosity on the evolution of altruism in a model implementing both reproduction and horizontal cultural transmission (as in Lehmann et al., 2008). Such a model could then contrast the effects of impecfect genetic

transmission and imperfect horizontal cultural transmission.

480 Coevolution of dispersal and social behavior

This work also raises the question of what would happen if dispersal (e.g., the 481 emigration probability m) could evolve as well. Recent work on the topic has 482 shown that under some conditions disruptive selection could take place, lead-483 ing to a polymorphism between sessile altruists and mobile defectors (Parvinen, 484 2013; Mullon et al., 2017). The assumptions of these studies however differ from 485 ours in important ways, in that they consider continuous traits and use an adap-486 tive dynamics framework, where, notably, mutations are assumed to be very 487 rare. It remains to be investigated how non-rare and potentially large mutations 488 would affect their result. 489

490 Acknowledgements

491 [redacted]

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596 Figures

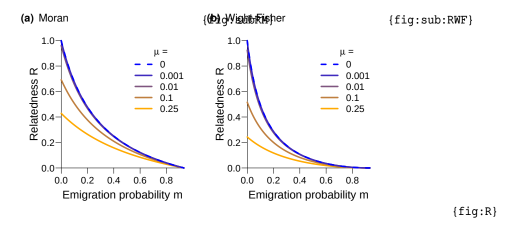


Figure 1: Within-deme relatedness of pairs of individuals R, as a function of the emigration probability m, for different values of the mutation probability μ (from 0 [blue] to 0.25 [orange]), and for the two types of life-cycles ((a): Moran, (b): Wright-Fisher). Other parameters: n=4 individuals per deme, $N_D=15$ demes.

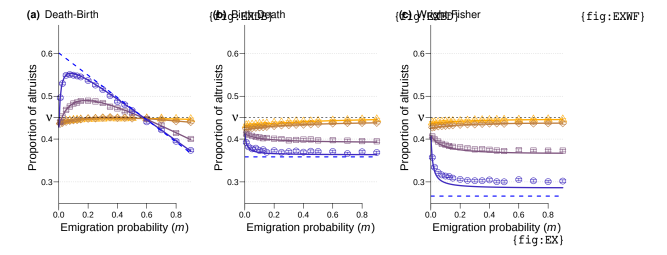


Figure 2: Expected proportion of altruists under weak selection, as a function of the emigration probability m, for different mutation values ($\mu=0.001$ (blue, dots), 0.01 (purple, squares), 0.1 (brown, diamonds), 0.25 (orange, triangles); the dashed blue lines correspond to $\mu=0$) and different life-cycles ((a) Moran Death-Birth, (b) Moran Birth Death, (c) Wright-Fisher). The curves are the analytical results, the points are the output of numerical simulations. Parameters: $\delta=0.005$, $\nu=0.45$, b=15, c=1, n=4 individuals per deme, $N_D=15$ demes.

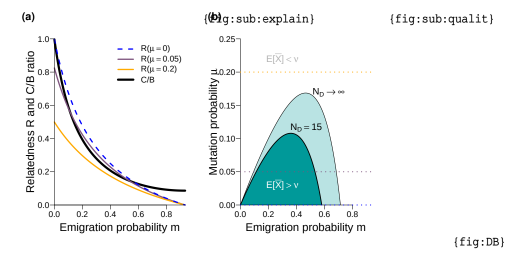


Figure 3: Understanding the effect of emigration m on whether altruism is favored in the Death-Birth life-cycle. (a) Comparison of the \mathcal{C}/\mathcal{B} ratio (thick black curve) and relatedness R (thin curves) for different values of the mutation probability μ (same color code as previously). (b) (m, μ) combinations for which $\mathbb{E}[\overline{X}] > v$. The dotted horizontal lines correspond to the mutation values used in panel (a). Unless specified, all other parameters are the same as in figure 2.

597 Supplementary figures

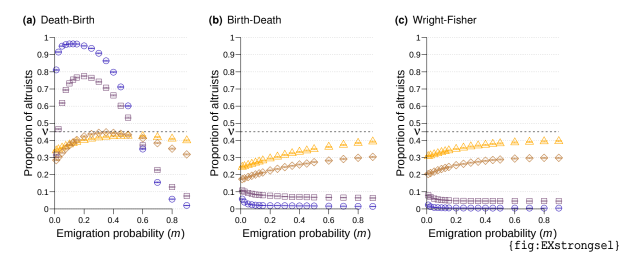


Figure A1: Equivalent of figure 2 (simulations only) but with strong selection (δ = 0.1); please note the change of scale on the vertical axis. All other parameters and legends are identical to those of figure 2 (increasing mutation probabilities from blue dots to orange triangles).

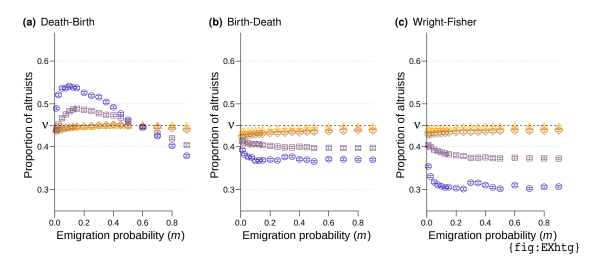


Figure A2: Equivalent of figure 2 (simulations only) but with a heterogeneous population structure: deme sizes range from 1 to 5 individuals per deme, the average deme size is 4 as in figure 2; all other parameters and legend are identical to those of figure 2.

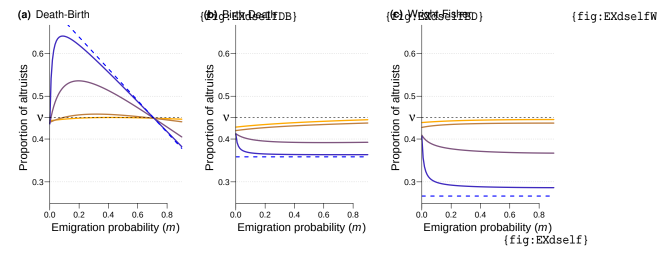


Figure A3: Equivalent of figure 2 (analysis only), with no self-replacement ($d_{ii} = d_{self} = 0$ for all sites).

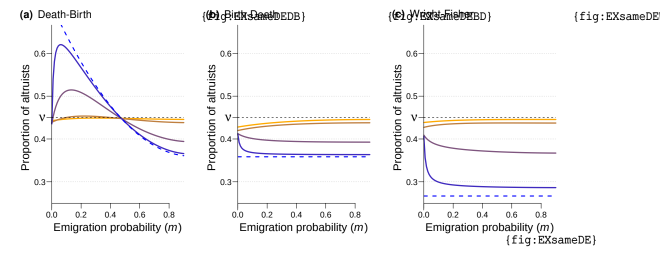


Figure A4: Equivalent of figure 2 (analysis only), with equal dispersal and interaction graphs (*i.e.*, no self-replacement [$d_{ii} = d_{\text{self}} = 0$ for all sites], and a proportion m of the interactions occurring outside of the home deme).

Supplementary Table

- b Sum of the marginal effects of deme-mates' phenotypes on focal individual's fecundity (benefit) \mathcal{B}
- Sum of the marginal effects of deme-mates' phenotypes on the fitness W of a focal individual
- B_i Expected number of successful offspring of the individual living at site i (r.v.)
- B^* Value of B_i for all sites, in the absence of selection ($\delta = 0$)
- С Marginal effect of a focal individual's phenotype on its own fecundity (cost)
- \mathcal{C} Marginal effect of an individual's phenotype on its own fitness W
- d_{ii} Dispersal probability from site i to site j
- D_i Probability that the individual currently living at site i is dead at the end of the time step (r.v.)
- Interaction probability from site *i* to site *j* e_{ii}
- Fecundity of the individual currently living at site i (r.v.) f_i
- nDeme size
- Number of demes N_D
- NTotal population size $(N = N_D n)$
- **Emigration probability** m
- P_{ii} (Long-term) Expected state of the pair of sites (i, j)
- (Long-term) Probability of identity by descent of individuals at sites i and j Q_{ij}
- Pairwise within-deme relatedness (see eq. (5))
- W_i Measure of fitness, counting offspring only when unmutated (see eq. (3))
- Indicator variable, equal to 1 if site i is occupied by an altruist, to 0 otherwise (r.v.) X_i
- \overline{X} Frequency of altruists in the population (r.v.)
- δ Phenotypic distance between altruists and defectors; strength of selection
- Phenotype of the individual living at site i; $\phi_i = \delta X_i$ (r.v.) ϕ_i
- Mutation probability μ
- Mutation bias: probability that mutant is altruist
- Subscript corresponding to primary effects P
- S Subscript corresponding to secondary effects
- Subscript used to denote a focal individual
- Subscript used when $i \neq j$ and the two sites are in the same deme in
- Subscript used when the two sites *i* and *j* are in different demes out
- Subscript used when i = iself
- 0 Sub- or superscript meaning that a quantity is evaluated at $\delta = 0$
- BDSuperscript corresponding to the Moran Birth-Death model
- Superscript corresponding to the Moran Death-Birth model DB
- Superscript corresponding to a Moran model Μ
- WF Superscript corresponding to the Wright-Fisher model

{tab:symbols}

Table A1: List of symbols. "r.v." means random variable.

Appendix

A Mutation parameters

599

{sec:app:mutation}

- In the main text, we first introduce effective mutation parameters: $\mu_{1\to 0}$, the probability that an altruist has defector offspring, and $\mu_{0\to 1}$, the probability that a defector has altruist offspring.
- ₆₀₄ A.1 Expected frequency of altruists at the mutation-drift balance

{sec:app:defnu}

- We assume that there is no selection acting ($\delta = 0$), but that there still are two types of individuals in the population.
- Let Y be the type of a randomly chosen individual in the population, given a proportion y of altruists in the population. In expectation, we have

$$\mathbb{E}[Y] = y. \tag{A1a}$$

Let Y' be the type of a randomly chosen individual at the next time step, given the frequency y at the previous time step. This randomly chosen individual is altruist if its parent was (y) and it did not mutate $(\mu_{1\rightarrow 0})$, or if its parent was not altruist (1-y), but the offspring mutated into one $(\mu_{0\rightarrow 1})$. We obtain

$$\mathbb{E}[Y'] = y(1 - \mu_{1 \to 0}) + (1 - y)\mu_{0 \to 1}. \tag{A1b}$$

The expected frequency of altruists, denoted by v, is found by solving $\mathbb{E}[Y] = \mathbb{E}[Y']$. We obtain

$$v = \frac{\mu_{0 \to 1}}{\mu_{1 \to 0} + \mu_{0 \to 1}}.$$
 (A2) {eq:app:nuformula}

A.2 Parent-offspring correlation at the mutation drift balance

We can then compute the parent-offspring type correlation at the mutation-drift balance. First, let us compute parent-offspring covariance:

$$\begin{aligned} \text{Cov} \big[YY' \big] &= \mathbb{E} \big[YY' \big] - \mathbb{E} \big[Y' \big] \mathbb{E} \big[Y \big] \\ &= \nu (1 - \mu_{1 \to 0}) - \left(\nu (1 - \mu_{1 \to 0}) + (1 - \nu) \mu_{0 \to 1} \right) \nu \\ &= \nu (1 - \nu) (1 - \mu_{1 \to 0} - \mu_{0 \to 1}). \end{aligned} \tag{A3} \quad \{ \text{eq:app:Cov} \}$$

Then, the standard deviations are given by

$$\sigma_{Y} = \sqrt{\mathbb{E}[Y^{2}] - \mathbb{E}[Y]^{2}} = \sqrt{\mathbb{E}[Y] - \mathbb{E}[Y]^{2}}$$

$$= \sqrt{v(1-v)},$$
(A4) {eq:app:SD1}

619 and

$$\begin{split} \sigma_{Y'} &= \sqrt{\mathbb{E}\big[Y'^2\big] - \mathbb{E}\big[Y'\big]^2} = \sqrt{\mathbb{E}\big[Y'\big] - \mathbb{E}\big[Y'\big]^2} \\ &= \sqrt{\nu(1-\nu)(1-\mu_{1\to 0}-\mu_{0\to 1}) - (\nu(1-\nu)(1-\mu_{1\to 0}-\mu_{0\to 1}))^2}. \end{split} \tag{A5)} \quad \{\text{eq:app:SD2}\}$$

Finally, the parent-offspring correlation is given by

$$\operatorname{Corr}[YY'] = \frac{\operatorname{Cov}[YY']}{\sigma_Y \sigma_{Y'}};$$

using the formulas eq. (A3)–(A5), and replacing ν by its value (mutation-drift equilibrium, eq. (A2)), we obtain

Corr
$$[YY'] = 1 - (\mu_{1\to 0} + \mu_{0\to 1}) = 1 - \mu.$$
 (A6)

A.3 Redefining the mutation scheme

{sec:app:mutnew}

With the new mutation parameters μ and ν , we can describe the mutation scheme differently.

If we denote by X_i the type of a given parent, then the expected type of one of its offspring is

$$\mathbb{E}[X_i'|X_i] = X_i(1 - \mu_{1 \to 0}) + (1 - X_i)\mu_{0 \to 1}. \tag{A7a} \quad \{eq:app:expoff\}$$

Replacing $\mu_{1\to 0}$ and $\mu_{0\to 1}$ by equivalent combinations of μ and ν , *i.e.*,

$$\mu_{1\to 0} = \mu(1-\nu) \text{ and } \mu_{0\to 1} = \mu\nu,$$
 (A7b)

629 then eq. (A7a) becomes

620

$$\mathbb{E}[X_i'|X_i] = X_i(1-\mu) + \mu\nu. \tag{A7c} \qquad \text{eq:app:expoff2}$$

We can redefine the mutation scheme and interpret eq. (A7c) as follows. Parents transmit their strategy to their offspring with probability $1 - \mu$; with probability μ , offspring do not inherit their strategy from their parent but instead get one randomly: with probability ν , they become altruists, with probability $1 - \nu$ they become defectors. With this alternative description, we can call "mutants" individuals who have the same type as their parent.

B Expected frequency of altruists

{sec:app:EX}

B.1 For a generic life-cycle

{sec:app:generic}

We want to compute the expected proportion of altruists in the population. We 638 represent the state of the population at a given time t using indicator variables 639 $X_i(t), 1 \le i \le N$, equal to 1 if the individual living at site i at time t is an altru-640 ist, and equal to 0 if it is a defector; these indicator variables are gathered in a 641 *N*-long vector $\mathbf{X}(t)$. The set of all possible population states is $\Omega = \{0,1\}^N$. The 642 proportion of altruists in the population is written $\overline{X}(t) = \sum_{i=1}^{N} X_i(t)$. We denote 643 by $B_{ii}(X(t),\delta)$, written B_{ii} for simplicity, the probability that the individual at site j at time t+1 is the newly established offspring of the individual living at site i at time t. The expected number of successful offspring produced by the indi-646 vidual living at site *i* at time *t* is given by $B_i = \sum_{i=1}^{N} B_{ji}$. We denote by $D_i(X(t), \delta)$ 647 (D_i for simplicity) the probability that the individual living at site i at time t has 648 been replaced (i.e., died) at time t + 1. These quantities depend on the chosen 649 life-cycle and on the state of the population; they are given in table A2 for each 650 of the life-cycles that we consider. 651

Life-cycle	B_{ji}	D_i
Moran Birth-Death	$d_{ij}\frac{f_i}{\sum_{k=1}^N f_k}$	$\frac{\sum_{j=1}^{N} d_{ji} f_j}{\sum_{k=1}^{N} f_k}$
Moran Death-Birth	$\frac{1}{N} \frac{d_{ij} f_i}{\sum_{k=1}^N d_{kj} f_k}$	$\frac{1}{N}$
Wright-Fisher	$\frac{d_{ij}f_i}{\sum_{k=1}^N d_{kj}f_k}$	1

{tab:BD}

Table A2: Formulas of B_{ji} and D_i for each of the life-cycles that we consider; f_i (shorthand notation for $f_i(X, \delta)$) is the fecundity of the individual living at site i, and d_{ji} is a dispersal probability, given in eq. (2).

Since a dead individual is immediately replaced by one new individual (i.e., population size remains constant and equal to N),

$$D_i = \sum_{j=1}^{N} B_{ij}$$
 (A8a) {eq:DBequiv}

holds for all sites i and all life-cycles.

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The structure of the population is also such that in the absence of selection $(\delta = 0$, so that $f_i = 1$ for all sites $1 \le i \le N$), all individuals have the same probability of dying and the same probability of having successful offspring (*i.e.*, of

having offspring that become adults at the next time step), so that

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$$D_i^0 = \sum_{j=1}^N B_{ji}^0 = B_i^0 =: B^*, \tag{A8b}$$
 {eq:DBRV}

where the 0 subscript means that the quantities are evaluated for $\delta=0$. This also implies that B_{ij}^0 and D_i^0 do not depend on the state **X** of the population. For the Moran life-cycles, $B^* = 1/N$, while for the Wright-Fisher life-cycle, $B^* = 1$. 661 (The difference between eq. (A8b) and eq. (A8a) is that we are now considering 662 offspring produced by i landing on i). 663

Given that the population is in state $\mathbf{X}(t)$ at time t, the expected frequency of altruists at time t + 1 is given by

$$\mathbb{E}\big[\overline{X}(t+1)|\mathbf{X}(t)\big] = \frac{1}{N}\sum_{i=1}^{N} \left[B_i(1-\mu)X_i + (1-D_i)X_i + B_i\mu\nu\right]. \tag{A9a} \quad \{\text{eq:conditionalchange}\}$$

The first term within the brackets corresponds to births of unmutated offspring from parents who are altruists (X_i) . The second term corresponds to the survival of altruists. The third term corresponds to the births of mutants who became altruists (which occurs with probability ν), whichever the type of the parent. 669

Given that there is no absorbing population state (a lost strategy can always 670 be recreated by mutation), there is a stationary distribution of population states; 671 the expected frequency of altruists does not change anymore for large times t 672 (realized frequencies of course keep changing). We denote by $\xi(\mathbf{X}, \delta, \mu)$ the prob-673 ability that the population is in state **X**, given the strength of selection δ and the 674 mutation probability μ . Taking the expectation of eq. (A9a) ($\mathbb{E}[\overline{X}] = \sum_{X \in \Omega} \overline{X} \xi(\mathbf{X}, \delta, \mu)$), we obtain, after reorganizing:

$$0 = \frac{1}{N} \sum_{X \in \Omega} \left[\sum_{i=1}^{N} \left(B_i (1 - \mu) X_i - D_i X_i \right) + \sum_{i=1}^{N} B_i \mu \nu \right] \xi(\mathbf{X}, \delta, \mu). \tag{A10} \quad \{ \text{eq:statdist} \}$$

Now, we use the assumption of weak selection ($\delta \ll 1$) and consider the first-677 order expansion of eq. (A10) for δ close to 0.

$$\begin{split} 0 &= \frac{1}{N} \sum_{X \in \Omega} \left[\sum_{i=1}^{N} \left(B_i^0 (1 - \mu) X_i - D_i^0 X_i \right) + \sum_{i=1}^{N} B_i^0 \mu \nu \right] \xi(\mathbf{X}, 0, \mu) \\ &+ \frac{1}{N} \sum_{X \in \Omega} \left[\sum_{i=1}^{N} \left(\frac{\partial B_i (1 - \mu) - D_i}{\partial \delta} X_i \right) + \sum_{i=1}^{N} \frac{\partial B_i}{\partial \delta} \mu \nu \right] \xi(\mathbf{X}, 0, \mu) \\ &+ \frac{1}{N} \sum_{X \in \Omega} \left[\sum_{i=1}^{N} \left(B_i^0 (1 - \mu) X_i - D_i^0 X_i \right) + \sum_{i=1}^{N} B_i^0 \mu \nu \right] \frac{\partial \xi(\mathbf{X}, \delta, \mu)}{\partial \delta}, \end{split} \tag{A11}$$

where all the derivatives are evaluated for $\delta = 0$. The first line of eq. (A11) is equal to zero, because $B_i^0=D_i^0=B^*$ (eq. (A8b)), and because in the absence of selection $(\delta=0)$, the expected state of every site i is $\mathbb{E}_0\big[X_i\big]=\sum_{X\in\Omega}X_i\xi(X,0,\mu)=v$ (by definition of ν , see Appendix A.1). The second terms of the second and third lines are both zero, because for all the life-cycles that we consider, the total number of births in the population during one time step $(\sum_{i=1}^{N} B_i)$ does not depend on population phenotypic composition (it is exactly 1 death for the Moran lifecycles, and exactly N for the Wright-Fisher life-cycle). Eq. (A11) then becomes

$$0 = \frac{\delta}{N} \sum_{i=1}^{N} \left[\sum_{X \in \Omega} \left(\frac{\partial B_i}{\partial \delta} (1 - \mu) - \frac{\partial D_i}{\partial \delta} \right) X_i \xi(\mathbf{X}, 0, \mu) - \sum_{X \in \Omega} \mu B^* X_i \frac{\partial \xi}{\partial \delta} \right] + O\left(\delta^2\right), \quad (A12) \quad \{\text{eq:weaksel0} : \mathbf{X} \in \mathcal{A} : \mathbf{X} : \mathbf{X} \in \mathcal{A} : \mathbf{X} :$$

where the derivatives are evaluated at $\delta = 0$. For conciseness, we define

$$W_i = (1 - \mu)B_i + (1 - D_i),$$
 (A13) {eq:app:defW}

a measure of fitness counting offspring only when they are unmutated (in the sense of the alternate mutation scheme described in Appendix A.3). With this, using the expectation notation, and denoting by $\mathbb{E}_0[]$ expectations under $\delta = 0$, we can rewrite and reorganize eq. (A12) as

$$\delta \mu B^* \frac{\partial \mathbb{E}[\overline{X}]}{\partial \delta} = \frac{\delta}{N} \sum_{i=1}^{N} \mathbb{E}_0 \left[\frac{\partial W_i}{\partial \delta} X_i \right] + O(\delta^2). \tag{A14} \quad \{eq: weaksel0 reorg\}$$

Now, we use a first time the law of total probabilities, taking individual phenotypes ϕ_k are intermediate variables:

$$\begin{split} \frac{\partial W_i}{\partial \delta} &= \sum_{k=1}^N \frac{\partial W_i}{\partial \phi_k} \frac{\partial \phi_k}{\partial \delta} \\ &= \sum_{k=1}^N \frac{\partial W_i}{\partial \phi_k} X_k, \end{split} \tag{A15} \quad \{eq:totalprobal}$$

by definition of ϕ_k ($\phi_k = \delta X_k$), and where the derivatives are evaluated for all $\phi_i = 0$, $1 \le i \le N$. Introducing the notation $P_{ij} = \mathbb{E}_0[X_i X_j]$ (expected state of a pair of sites), eq. (A14) becomes

$$\delta \mu B^* \frac{\partial \mathbb{E}[\overline{X}]}{\partial \delta} = \frac{\delta}{N} \sum_{i=1}^{N} \sum_{k=1}^{N} \frac{\partial W_i}{\partial \phi_k} P_{ik} + O(\delta^2). \tag{A16}$$

So far, we have not used the specificities of the population structure that we consider. Once we have fixed a focal individual i, in expectation there are only three types of individuals: the focal itself (denoted by " \bullet "), n-1 other individuals in the focal's deme (denoted by "in"), and N-n individuals in other demes (denoted by "out"). We note that given that the size of the population is fixed ($\sum_{i=1}^{N} (B_i - D_i) = 0$), and given that the total number of births does not depend on population composition in the life-cycles that we consider,

$$\sum_{i=1}^{N} \frac{\partial W_i}{\partial \delta} = 0,$$

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which we can rewrite as (Rousset & Billiard, 2000, p.817–818)

$$\frac{\partial W_i}{\partial \phi_i} + (n-1)\frac{\partial W_i}{\partial \phi_{\text{in}}} + (N-n)\frac{\partial W_i}{\partial \phi_{\text{out}}} = 0. \tag{A17} \quad \{\text{eq:derivsumW}\}$$

With this, eq. (A16) becomes

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$$\delta \mu B^* \frac{\partial \mathbb{E}\left[\overline{X}\right]}{\partial \delta} = \frac{\delta}{N} \sum_{i=1}^{N} \left(\frac{\partial W_i}{\partial \phi_i} + (n-1) \frac{\partial W_i}{\partial \phi_{\text{in}}} \frac{P_{\text{in}} - P_{\text{out}}}{P_{ii} - P_{\text{out}}} \right) (P_{ii} - P_{\text{out}}) + O\left(\delta^2\right). \quad (A18) \quad \{\text{eq:weaksel1CBRP}\}$$

We can also replace the P terms by 706

$$P_{ij} = Q_{ij}v + (1 - Q_{ij})v^{2}$$

$$= v^{2} + v(1 - v)Q_{ij}.$$
(A19) {eq:QP}

In Appendix C.1, using recursions on P_{ij} , we will see that Q_{ij} can be interpreted as a probability of identity by descent, i.e., the probability that the individuals at sites i and j have a common ancestor and that no mutation (using the alternative mutation scheme described in Appendix A.3) has occurred on either lineage since the ancestor. Eq. (A18) becomes

$$\delta \mu B^* \frac{\partial \mathbb{E}\left[\overline{X}\right]}{\partial \delta} = \frac{\delta}{N} \sum_{i=1}^{N} \left(\underbrace{\frac{\partial W_i}{\partial \phi_i}}_{-\mathcal{C}} + \underbrace{(n-1)\frac{\partial W_i}{\partial \phi_{\text{in}}}}_{\mathcal{B}} \underbrace{\frac{Q_{\text{in}} - Q_{\text{out}}}{1 - Q_{\text{out}}}}_{R} \right) (1 - Q_{\text{out}}) v (1 - v) + O\left(\delta^2\right). \tag{A20} \quad \{\text{eq:weaksel1CBR}}$$

We can further decompose the derivatives, now using the fecundities f_{ℓ} as intermediate variables, i.e.,

$$\frac{\partial W_i}{\partial \phi_k} = \sum_{\ell=1}^N \frac{\partial W_i}{\partial f_\ell} \frac{\partial f_\ell}{\partial \phi_k}.$$
 (A21)

The term $\frac{\partial f_{\ell}}{\partial \phi_k}$ is the marginal effect of a change in the phenotype of the in-714 dividual living at site k on the fecundity of the individual living at site ℓ . By assumption, social interactions take place within demes only, so whenever ℓ and kare in different demes, we have $\frac{\partial f_\ell}{\partial \phi_k} = \frac{\partial f_\ell}{\partial \phi_{\text{out}}} = 0$. We then need to characterize the effect of one's own phenotype $(k = \ell)$ and of another deme-mate's phenotype $(k = \ell)$ and ℓ different sites in the same deme) on fecundity, and define

{eq:derivf}

$$\frac{\partial f_{\ell}}{\partial \phi_{\ell}}\Big|_{\delta=0} = -\mathsf{c},$$
 (A22a)

$$\frac{\partial f_{\ell}}{\partial \phi_{\ell}} \Big|_{\delta=0} = -c, \tag{A22a}$$

$$\frac{\partial f_{\ell}}{\partial \phi_{\text{in}}} \Big|_{\delta=0} = \frac{b}{n-1}. \tag{A22b}$$

Eq. (A20) then becomes (using notation • to refer to the focal individual itself,

and where $W = W_i$, since the derivatives are the same for all i):

$$\delta \mu B^* \frac{\partial \mathbb{E}\left[\overline{X}\right]}{\partial \delta} = \delta \nu (1 - \nu) (1 - Q_{\text{out}}) \times \left(\underbrace{\frac{\partial W}{\partial f_{\bullet}} (-c) + \frac{\partial W}{\partial f_{\text{in}}} b}_{-\mathcal{C}} + \underbrace{\left(\frac{\partial W}{\partial f_{\bullet}} b + (n - 1) \frac{\partial W}{\partial f_{\text{in}}} (-c) + (n - 2) \frac{\partial W}{\partial f_{\text{in}}} b\right)}_{\mathcal{B}} \underbrace{\frac{Q_{\text{in}} - Q_{\text{out}}}{1 - Q_{\text{out}}}}_{R}\right) + O\left(\delta^2\right). \tag{A23} \quad \{\text{eq:weaksel2}\}$$

(As previously, all derivatives are evaluated at $\delta = 0$.)

Finally, we obtain a first-order approximation of the expected frequency of altruists in the population with

$$\mathbb{E}\left[\overline{X}\right] = \nu + \delta \left. \frac{\partial \mathbb{E}\left[\overline{X}\right]}{\partial \delta} \right|_{\delta=0} + O\left(\delta^2\right), \tag{A24} \quad \{eq:app:EXgeneric\}$$

where $v = \mathbb{E}_0[\overline{X}]$ (expected frequency in the absence of selection), and where $\frac{\partial \mathbb{E}[\overline{X}]}{\partial \delta}\Big|_{\delta=0}$ is obtained from eq. (A23). We then need to replace the B_i and D_i terms by their formulas for each life-cycle; they are given in table A2.

B.2 Derivatives for the specific life-cycles

{sec:app:dW}

We use the formulas presented in table A2 and the definition of $W = W_i$ given in eq. (A13) for each life-cycle. In eq. (A26), eq. (A28) and eq. (A30), the first lines within parentheses correspond to primary effects, and the second line to secondary effects.

Moran Birth-Death Under this life-cycle, we obtain

{eq:dWBD}

$$\frac{\partial W^{\rm BD}}{\partial f_{\bullet}} \bigg|_{\delta=0} = (1-\mu) \left(\frac{1}{N} - \frac{1}{N^2} \right) - \left(\frac{1-m}{nN} - \frac{1}{N^2} \right) = \frac{1-\mu}{N} + \frac{\mu}{N^2} - \frac{1-m}{nN}, \quad (A25a)$$

$$\frac{\partial W^{\rm BD}}{\partial f_{\rm in}} \bigg|_{\delta=0} = (1-\mu) \left(-\frac{1}{N^2} \right) - \left(\frac{1-m}{nN} - \frac{1}{N^2} \right) = \frac{\mu}{N^2} - \frac{1-m}{nN}. \quad (A25b)$$

With these derivatives, eq. (5) becomes

$$\mathbb{E}\left[\overline{X}\right] \approx \nu + \frac{\delta}{\mu} \nu (1 - \nu) (1 - Q_{\text{out}}^{\text{M}}) \times \\ \left[\underbrace{\left(\frac{(1 - \mu)(-c)}{+(b - c)\left(\frac{\mu}{N} - \frac{1 - m}{n}\right)}\right)}_{-\mathcal{C}^{\text{BD}}} + \underbrace{\left(\frac{(1 - \mu)b}{+(b - c)(n - 1)\left(\frac{\mu}{N} - \frac{1 - m}{n}\right)}_{\mathcal{B}^{\text{BD}}} \underbrace{\frac{Q_{\text{in}}^{\text{M}} - Q_{\text{out}}^{\text{M}}}{1 - Q_{\text{out}}^{\text{M}}}\right]}_{\text{R}^{\text{M}}},$$

$$(A26) \quad \{\text{eq:EXBD}\}$$

In addition, for both Moran life-cycles, we have $B_{\rm M}^*=1/N$. The secondary effects (second line in the parentheses in eq. (A26)) include competitive effects

on the probability of reproducing, and consequences of social interactions on the probability that a given individual dies. Note that the secondary effects remain negative for the realistic range of emigration values that we consider (*i.e.*, $m < 1 - 1/N_D$).

Moran Death-Birth Under this life-cycle, we obtain

{eq:dWDB}

$$\left. \frac{\partial W^{\rm DB}}{\partial f_{\bullet}} \right|_{\delta=0} = \frac{1-\mu}{N} \left[1 - \left(\frac{(1-m)^2}{n} + \frac{m^2}{N-n} \right) \right],\tag{A27a}$$

$$\left. \frac{\partial W^{\rm DB}}{\partial f_{\rm in}} \right|_{\delta=0} = -\frac{1-\mu}{N} \left(\frac{(1-m)^2}{n} + \frac{m^2}{N-n} \right). \tag{A27b}$$

With the Death-Birth life-cycle, eq. (5) becomes

$$\begin{split} \mathbb{E}\left[\overline{X}\right] &\approx \nu + \frac{\delta}{\mu}\nu(1-\nu)(1-Q_{\text{out}}^{\text{M}}) \times \\ &\left[\underbrace{\begin{pmatrix} (1-\mu)(-c) \\ -(b-c)(1-\mu)\left(\frac{(1-m)^2}{n} + \frac{m^2}{N-n}\right) \end{pmatrix}}_{-\mathcal{C}^{\text{DB}}} + \underbrace{\begin{pmatrix} (1-\mu)b \\ -(b-c)(n-1)(1-\mu)\left(\frac{(1-m)^2}{n} + \frac{m^2}{N-n}\right) \right)}_{\mathcal{B}^{\text{DB}}} \underbrace{\begin{pmatrix} Q_{\text{in}}^{\text{M}} - Q_{\text{out}}^{\text{M}} \\ 1 - Q_{\text{out}}^{\text{M}} \end{pmatrix}}_{R^{\text{M}}}, \\ &\left(A28\right) \quad \{\text{eg:EXDB}} \end{split}$$

With this life-cycle, Death occurs first, and the probability of dying is independent from the state of the population (since we assume that social interactions affect fecundity. We can therefore factor $(1 - \mu)$ in all terms. The primary effects (first lines in the parentheses) remain the same as with the Birth-Death lifecycle. However, the Death-Birth life-cycle leads to different secondary effects compared to the Birth-Death life-cycle: competition occurs at a different scale (Grafen & Archetti, 2008). Finally, with this life-cycle as we defined it, the probabilities of identity by descent Q are the same as with the Birth-Death model.

Wright-Fisher Under this life-cycle, we obtain

{eq:dWWF}

$$\left. \frac{\partial W^{\text{WF}}}{\partial f_{\bullet}} \right|_{\delta=0} = (1-\mu) \left[1 - \left(\frac{(1-m)^2}{n} + \frac{m^2}{N-n} \right) \right],\tag{A29a}$$

$$\left. \frac{\partial W^{\text{WF}}}{\partial f_{\text{in}}} \right|_{\delta=0} = -(1-\mu) \left(\frac{(1-m)^2}{n} + \frac{m^2}{N-n} \right). \tag{A29b}$$

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For the Wright-Fisher life-cycle, we have $B_{\rm WF}^*=1$. Replacing the derivatives presented in eq. (A29) into eq. (5), we obtain

$$\begin{split} \mathbb{E}\left[\overline{X}\right] &\approx \nu + \frac{\delta}{\mu}\nu(1-\nu)(1-Q_{\text{out}}^{\text{WF}}) \times \\ &\left[\underbrace{\begin{pmatrix} (1-\mu)(-c) \\ -(b-c)(1-\mu)\left(\frac{(1-m)^2}{n} + \frac{m^2}{N-n}\right) \end{pmatrix}}_{-\mathcal{C}^{\text{WF}}} + \underbrace{\begin{pmatrix} (1-\mu)b \\ -(b-c)(n-1)(1-\mu)\left(\frac{(1-m)^2}{n} + \frac{m^2}{N-n}\right) \end{pmatrix}}_{\mathcal{B}^{\text{WF}}} \underbrace{\begin{pmatrix} Q_{\text{in}}^{\text{WF}} - Q_{\text{out}}^{\text{WF}} \\ 1 - Q_{\text{out}}^{\text{WF}} \end{pmatrix}}_{R^{\text{WF}}}\right], \\ &\left[(A30) \quad \{\text{eq:EXWF}}\right] \end{split}$$

- The only but important different between eq. (A30) and eq. (A28) is the value
- $_{755}$ of the probabilities of identity by descent Q, because the number of individuals
- that are updated at each time step differs.

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C Probabilities of identity by descent

C.1 Expected state of pairs of sites and probabilities of identity by descent

{sec:app:IBD}

Here we show the link between the expected state of a pair of sites P_{ij} and probabilities of identity by descent Q_{ij} . In our derivation of $\mathbb{E}[\overline{X}]$, P_{ij} is the quantity that appears, but most studies use Q_{ij} . Both are evaluated in the absence of selection ($\delta = 0$).

C.1.1 Moran model

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In a Moran model, exactly one individual dies and one individual reproduces during one time step. Given a state **X** at time t, at time t+1 both sites i and $j \neq i$ are occupied by altruists, if i) it was the case at time t and neither site was replaced by a non-altruist (first term in eq. (A31)), or ii) if exactly one of the two sites was occupied by a non-altruist at time t, but the site was replaced by an altruist (second and third terms of eq. (A31)):

$$\begin{split} \mathbb{E} \big[X_i X_j(t+1) | X(t) &= \mathbf{X} \big] = & X_i X_j \left(1 - \sum_{k=1}^N \frac{1}{N} \left(d_{ki} + d_{kj} \right) \left((1 - X_k) (1 - \mu) + \mu (1 - \nu) \right) \right) \\ &+ X_i (1 - X_j) \sum_{k=1}^N \frac{1}{N} d_{kj} \left(X_k (1 - \mu) + \mu \nu \right) \\ &+ X_j (1 - X_i) \sum_{k=1}^N \frac{1}{N} d_{ki} \left(X_k (1 - \mu) + \mu \nu \right). \end{split} \tag{A31} \quad \{ \text{eq:app:PijM1} \}$$

We take the expectation of this quantity, and consider that the stationary distribution is reached $(t \to \infty)$; then $\mathbb{E}[X_i X_j(t+1)] = \mathbb{E}[X_i X_j(t)]$, and we obtain

$$P_{ij} = \frac{1}{2} \left(\sum_{k=1}^{N} (1 - \mu) \left(d_{kj} P_{ki} + d_{ki} P_{kj} \right) \right) + \mu v^2 \qquad (i \neq j), \tag{A32} \quad \{eq:app:PijM\}$$

773 while $P_{ii} = v$.

Now we substitute $P_{ij} = v^2 + v(1 - v)Q_{ij}$ in eq. (A32), we obtain

$$Q_{ij} = \frac{1}{2} \sum_{k=1}^{N} (1 - \mu) \left(d_{ki} Q_{kj} + d_{kj} Q_{ki} \right), \tag{A33} \quad \{eq:app:QijM\}$$

and we realize that Q_{ij} is the probability that the individuals at sites i and $j \neq i$ are identical by descent. To compute it indeed, we need to pick which site was last updated (equal probabilities), then who was the parent (k); the other individual needs to be identical by descent to the parent, and no mutation should have occurred $(1 - \mu)$.

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780 C.1.2 Wright-Fisher model

In a Wright-Fisher model, all individuals are replaced at each time step, so we directly consider the state of the parents:

$$\begin{split} \mathbb{E} \big[X_i X_j(t+1) | X(t) &= \mathbf{X} \big] = \sum_{k,\ell=1}^N d_{ki} d_{\ell j} \bigg(X_k X_\ell (1-\mu+\mu\nu)^2 \\ &\quad + (X_k (1-X_\ell) + (1-X_k) X_\ell) \, (1-\mu+\mu\nu) (\mu\nu) \\ &\quad + (1-X_k) (1-X_\ell) (\mu\nu)^2 \bigg) \end{split} \tag{A34} \quad \{ \text{eq:app:PijWF1} \}$$

The first term of eq. (A34) corresponds to both parents being altruists, and hav-

ing altruist offspring; the second line corresponds to exactly one parent being

altruist, and the third line to both parents being non-altruists (in this latter case,

the two offspring have to be both mutants to be altruists).

787 Taking the expectation and simplifying, we obtain

$$P_{ij} = \sum_{k,\ell=1}^{N} \left(P_{kl} (1 - \mu)^2 \right) + (2 - \mu) \mu v^2. \tag{A35} \quad \{eq:app:PijWF\}$$

Replacing P_{ij} by $v^2 + v(1-v)Q_{ij}$, eq. (A35) becomes

$$Q_{ij} = \sum_{k,\ell=1}^{N} d_{ki} d_{\ell j} Q_{k\ell} (1-\mu)^{2}.$$
 (A36) {eq:app:QijWF}

Again, Q_{ij} corresponds to a probability of identity by descent: the individuals at sites i and j are identical by descent if their parents were and if neither mutated $((1-\mu)^2)$.

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C.2 Probabilities of identity by descent in a subdivided population

{sec:app:Qsubdiv}

Two individuals are said to be identical by descent if there has not been any mutation on either lineage since their common ancestor. Because of the structure of the population, there are only three types of pairs of individuals, and hence three different values of the probabilities of identity by descent of pairs of sites Q_{ij} :

$$Q_{ij} = \begin{cases} 1 & \text{when } i = j; \\ Q_{\text{in}} & \text{when } i \neq j \text{ and both sites are in the same deme;} \\ Q_{\text{out}} & \text{when sites } i \text{ and } j \text{ are in different demes.} \end{cases}$$
 (A37)

The values of $Q_{\rm in}$ and $Q_{\rm out}$ depend on the type of life-cycle that we consider.

Here, we will use formulas derived in Débarre (2017) for "two-dimensional 799 population structures". The name comes from the fact that we only need two 800 types of transformations to go from any site to any other site in the population: 801 permutations on the deme index, and permutations on the within-deme index. 802 We rewrite site labels $(1 \le i \le N)$ as (ℓ_1, ℓ_2) , where ℓ_1 is the index of the deme $(1 \le i \le N)$ 803 $\ell_1 \leq N_D$) and ℓ_2 the position of the site within the deme $(1 \leq \ell_2 \leq n)$. Then, we 804 introduce notations $ilde{d}_{i_1}$ and $ilde{Q}_{i_1}$, that correspond to the dispersal probability and 805 probability of identity by descent to a site at distances i_1 and i_2 in the amongdemes and within-deme dimensions (*e.g.*, $\tilde{d}_{i_1} = d_{j_1} \,_{j_1+i_1}$.)

Also, in this section, we distinguish between $d_{\text{self}} = d_{ii}$ and d_{in} (in the main 807 808

Also, in this section, we distinguish between $d_{\text{self}} = d_{ii}$ and d_{in} (in the main text, $d_{\text{self}} = d_{\text{in}}$).

10 C.2.1 Moran model

In Débarre (2017), it was shown that

$$\tilde{\mathcal{Q}}_{r_{2}}^{r_{1}} = \frac{1}{N} \sum_{q_{1}=0}^{N_{1}-1} \sum_{q_{2}=0}^{N_{2}-1} \frac{\mu \lambda_{M}'}{1 - (1 - \mu)\tilde{\mathcal{D}}_{q_{1}}^{q_{1}}} \exp\left(i\frac{2\pi q_{1}r_{1}}{N_{1}}\right) \exp\left(i\frac{2\pi q_{2}r_{2}}{N_{2}}\right) \tag{A38a} \quad \{\text{eq:app:Q2DM}}$$

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$$\tilde{\mathcal{D}}_{q_1} = \sum_{\ell_1 = 0}^{N_1 - 1} \sum_{\ell_2 = 0}^{N_2 - 1} \tilde{d}_{\ell_1} \exp\left(-i\frac{2\pi q_1 \ell_1}{N_1}\right) \exp\left(-i\frac{2\pi q_2 \ell_2}{N_2}\right), \tag{A38b} \quad \{\text{eq:app:D2D}\}$$

and λ_M' such that $\tilde{\mathcal{Q}}_0 = 1$. Let us first compute $\tilde{\mathcal{D}}_{q_1}^{q_1}$ in the case of a subdivided population, with $N_1 = N_D$ and $N_2 = n$:

$$\tilde{\mathcal{D}}_{q_{2}}^{q_{1}} = d_{\text{self}} + \sum_{\ell_{2}=1}^{N_{2}-1} d_{\text{in}} \exp\left(-i\frac{2\pi q_{2}\ell_{2}}{N_{2}}\right) + \sum_{\ell_{1}=1}^{N_{1}-1} \sum_{\ell_{2}=0}^{N_{2}-1} d_{\text{out}} \exp\left(-i\frac{2\pi q_{1}\ell_{1}}{N_{1}}\right) \exp\left(-i\frac{2\pi q_{2}\ell_{2}}{N_{2}}\right) \\
= d_{\text{self}} + \left(\delta_{q_{2}}(N_{2}-1) + (1-\delta_{q_{2}})(-1)\right) d_{\text{in}} + \left(\delta_{q_{1}}(N_{1}-1) + (1-\delta_{q_{1}})(-1)\right) \left(\delta_{q_{2}}N_{2}\right) d_{\text{out}} \\
= d_{\text{self}} + \left(\delta_{q_{2}}N_{2}-1\right) d_{\text{in}} + \left(\delta_{q_{1}}N_{1}-1\right) \delta_{q_{2}}N_{2} d_{\text{out}}. \tag{A39a}$$

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815 (δ_q is equal to 1 when q is equal to 0 modulo the relevant dimension, and to 0 816 otherwise). So for the three types of distances that we need to consider (distance 817 0, distance to another deme-mate, distance to individual in another deme), and 818 with $N_1 = N_D$ and $N_2 = n$, we obtain

{eq:app:Dsystem}

$$\tilde{\mathcal{D}}_0 = 1, \tag{A40a}$$

$$\tilde{\mathcal{D}}_{q_1} = 1 - m - \frac{m}{N_D - 1} \quad (q_1 \not\equiv 0 \pmod{N_1}),$$
 (A40b)

$$\tilde{\mathcal{D}}_{q_1} = d_{\text{self}} - d_{\text{in}} \quad (q_2 \not\equiv 0 \pmod{N_2}).$$
 (A40c)

So for $\tilde{\mathcal{Q}}$, using system (A40) in eq. (A38a),

$$\begin{split} \tilde{\mathcal{Q}}_{r_{2}^{1}} &= \frac{\mu \lambda_{M}^{\prime}}{N} \left[\frac{1}{1 - (1 - \mu)\tilde{\mathcal{D}}_{0}^{0}} + \sum_{q_{2}=1}^{N_{2}-1} \frac{1}{1 - (1 - \mu)\tilde{\mathcal{D}}_{0}^{0}} \exp\left(-i\frac{2\pi q_{2}r_{2}}{N_{2}}\right) \right. \\ &\quad + \sum_{q_{1}=1}^{N_{1}-1} \frac{1}{1 - (1 - \mu)\tilde{\mathcal{D}}_{q_{1}}^{0}} \exp\left(-i\frac{2\pi q_{1}r_{1}}{N_{1}}\right) \\ &\quad + \sum_{q_{1}=1}^{N_{1}-1} \sum_{q_{2}=1}^{N_{2}-1} \frac{1}{1 - (1 - \mu)\tilde{\mathcal{D}}_{q_{1}}^{0}} \exp\left(-i\frac{2\pi q_{1}r_{1}}{N_{1}}\right) \exp\left(-i\frac{2\pi q_{2}r_{2}}{N_{2}}\right) \right] \\ &\quad = \frac{\mu \lambda_{M}^{\prime}}{N} \left[\frac{1}{1 - (1 - \mu)} + \frac{1}{1 - (1 - \mu)(d_{\text{self}} - d_{\text{in}})} (\delta_{r_{2}}N_{2} - 1) \right. \\ &\quad + \frac{1}{1 - (1 - \mu)(1 - m - \frac{m}{N_{D} - 1})} (\delta_{r_{1}}N_{1} - 1) \\ &\quad + \frac{1}{1 - (1 - \mu)(d_{\text{self}} - d_{\text{in}})} (\delta_{r_{1}}N_{1} - 1)(\delta_{r_{2}}N_{2} - 1) \right]. \end{split} \tag{A41} \quad \{\text{eq:app:Q2DMsol}\}$$

820 In particular,

$$\begin{split} \tilde{\mathcal{Q}}_{0}^{0} &= \frac{\mu \lambda_{M}^{\prime}}{N} \left[\frac{1}{\mu} + \frac{1}{1 - (1 - \mu)(d_{\text{self}} - d_{\text{in}})} (n - 1) + \frac{1}{1 - (1 - \mu)(1 - m - \frac{m}{N_{D} - 1})} (N_{D} - 1) \right. \\ &\quad + \frac{1}{1 - (1 - \mu)(d_{\text{self}} - d_{\text{in}})} (N_{D} - 1) (n - 1) \right] \\ &= 1. \end{split} \tag{A42a} \quad \{ \text{eq:app:Q2D1} \}$$

We find λ_M' using eq. (A42a). Let's now go back to eq. (A41): when $r_1=0$, the two individuals are in the same deme. They are different when $r_2\not\equiv 0$, and so:

$$Q_{\text{in}} = \frac{\mu \lambda_M'}{N} \left[\frac{1}{\mu} + \frac{1}{1 - (1 - \mu)(d_{\text{self}} - d_{\text{in}})} (-1) + \frac{1}{1 - (1 - \mu)(1 - m - \frac{m}{N_D - 1})} (D - 1) + \frac{1}{1 - (1 - \mu)(d_{\text{self}} - d_{\text{in}})} (D - 1) (-1) \right].$$
(A42b)

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And when $r_1 \not\equiv 0$, the two individuals are in different demes:

$$Q_{\text{out}} = \frac{\mu \lambda_M'}{N} \left[\frac{1}{\mu} + \frac{1}{1 - (1 - \mu)(d_{\text{self}} - d_{\text{in}})} (-1) + \frac{1}{1 - (1 - \mu)(1 - m - \frac{m}{N_D - 1})} (-1) + \frac{1}{1 - (1 - \mu)(d_{\text{self}} - d_{\text{in}})} \right].$$
(A42c)

With $d_{\text{self}} = d_{\text{in}} = (1 - m)/n$, we eventually obtain:

{eq:QM}

$$Q_{\text{in}}^{\text{M}} = \frac{(1-\mu)\left(m+\mu(N_D(1-m)-1)\right)}{(1-\mu)m(N_D\mu(n-1)+1)+(N_D-1)\mu(\mu(n-1)+1)},$$
 (A43a)

$$Q_{\text{out}}^{\text{M}} = \frac{(1-\mu)m}{(1-\mu)m(N_D\mu(n-1)+1) + (N_D-1)\mu(\mu(n-1)+1)}.$$
 (A43b)

The probability that two different deme-mates are identical by descent, $Q_{\rm in}^{\rm M}$, decreases monotonically with the emigration probability m, while $Q_{\rm out}^{\rm M}$ monotonically increases with m (see figure A5(a)).

When the mutation probability μ is vanishingly small ($\mu \to 0$), both $Q_{\rm in}^{\rm M}$ and $Q_{\rm out}^{\rm M}$ are equal to 1: in the absence of mutation indeed, the population ends up fixed for one of the two types, and all individuals are identical by descent. Note that we obtain a different result if we first assumed that the size of the population is infinite ($N_D \to \infty$), because the order of limits matters; for instance, $\lim_{N_D \to \infty} Q_{\rm out}^{\rm M} = 0$.

Using eq. (A43), relatedness under the Moran model is given by

$$R^{\rm M} = \frac{(1-\mu)(N_D(1-m)-1)}{N_D(1-\mu)m(n-1) + (N_D-1)(1+\mu(n-1))}. \tag{A44} \quad \{eq:app:RM\}$$

When there is an infinite number of demes $(N_D \to \infty)$ and mutation is vanishingly small $(\mu \to 0)$, we recover

$$\lim_{\mu \to 0} \lim_{N_D \to \infty} R^{M} = \lim_{N_D \to \infty} \lim_{\mu \to 0} R^{M} = \frac{1 - m}{1 + m(n - 1)}.$$
 (A45) {eq:app:RMlim}

37 C.2.2 Wright-Fisher

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For the Wright-Fisher updating, the equation for $ilde{Q}$ is different:

$$\tilde{Q}_{r_{2}}^{r_{1}} = \frac{1}{N} \sum_{q_{1}=0}^{N_{1}-1} \sum_{q_{2}=0}^{N_{2}-1} \frac{\mu \lambda'_{WF}}{1 - (1 - \mu)^{2} (\tilde{\mathcal{D}}_{q_{1}})^{2}} \exp\left(-\iota \frac{2\pi q_{1} r_{1}}{N_{1}}\right) \exp\left(-\iota \frac{2\pi q_{2} r_{2}}{N_{2}}\right), \quad (A46)$$

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with $\tilde{\mathcal{D}}$ given in eq. (A38b). In a subdivided population, with $N_1=N_D$ and $N_2=n$, this becomes

$$\begin{split} \tilde{\mathcal{Q}}_{r_{2}}^{r_{1}} &= \frac{1}{N} \bigg[\frac{\mu \lambda_{WF}'}{1 - (1 - \mu)^{2} (\tilde{\mathcal{D}}_{0})^{2}} + \sum_{q_{2}=1}^{N_{2}-1} \frac{\mu \lambda_{WF}'}{1 - (1 - \mu)^{2} (\tilde{\mathcal{D}}_{0})^{2}} \exp \bigg(- i \frac{2\pi q_{2} r_{2}}{N_{2}} \bigg) \\ &+ \sum_{q_{1}=1}^{N_{1}-1} \frac{\mu \lambda_{WF}'}{1 - (1 - \mu)^{2} (\tilde{\mathcal{D}}_{q_{1}})^{2}} \exp \bigg(- i \frac{2\pi q_{1} r_{1}}{N_{1}} \bigg) \\ &+ \sum_{q_{1}=1}^{N_{1}-1} \sum_{q_{2}=1}^{N_{2}-1} \frac{\mu \lambda_{WF}'}{1 - (1 - \mu)^{2} (\tilde{\mathcal{D}}_{q_{1}})^{2}} \exp \bigg(- i \frac{2\pi q_{1} r_{1}}{N_{1}} \bigg) \exp \bigg(- i \frac{2\pi q_{2} r_{2}}{N_{2}} \bigg) \bigg] \\ &= \frac{\mu \lambda_{WF}'}{N} \bigg[\frac{1}{1 - (1 - \mu)^{2}} + \frac{1}{1 - (1 - \mu)^{2} (d_{\text{self}} - d_{\text{in}})^{2}} (\delta_{q_{2}} N_{2} - 1) \\ &+ \frac{1}{1 - (1 - \mu)^{2} (1 - m - \frac{m}{N_{D} - 1})^{2}} (\delta_{q_{1}} N_{1} - 1) \\ &+ \frac{1}{1 - (1 - \mu)^{2} (d_{\text{self}} - d_{\text{in}})^{2}} (\delta_{q_{1}} N_{1} - 1) (\delta_{q_{2}} N_{2} - 1) \bigg] \\ &= \frac{\mu \lambda_{WF}'}{N} \bigg[\frac{1}{1 - (1 - \mu)^{2}} + \frac{1}{1 - (1 - \mu)^{2} (d_{\text{self}} - d_{\text{in}})^{2}} (\delta_{q_{1}} N_{1} - 1) \bigg] . \end{aligned} \tag{A47} \quad \{\text{eq:app:Q2DWFsol}\}$$

To find λ'_{WF} , we solve $\tilde{\mathcal{Q}}_0 = 1$, *i.e.*,

$$1 = \frac{\mu \lambda_{WF}'}{N} \left[\frac{1}{1 - (1 - \mu)^2} + \frac{1}{1 - (1 - \mu)^2 (d_{\text{self}} - d_{\text{in}})^2} (N_2 - 1) N_1 + \frac{1}{1 - (1 - \mu)^2 (1 - m - \frac{m}{N_D - 1})^2} (N_1 - 1) \right]. \tag{A48a}$$

Then from eq. (A47) we deduce

$$Q_{\rm in} = \frac{\mu \lambda_{WF}'}{N} \left[\frac{1}{1 - (1 - \mu)^2} - \frac{1}{1 - (1 - \mu)^2 (d_{\rm self} - d_{\rm in})^2} N_1 + \frac{1}{1 - (1 - \mu)^2 (1 - m - \frac{m}{N_D - 1})^2} (N_1 - 1) \right]. \tag{A48b}$$

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$$Q_{\text{out}} = \frac{\mu \lambda_{WF}'}{N} \left[\frac{1}{1 - (1 - \mu)^2} - \frac{1}{1 - (1 - \mu)^2 (1 - m - \frac{m}{d-1})^2} \right].$$
 (A48c)

With $d_{\text{self}} = d_{\text{in}} = (1 - m)/n$, we obtain:

{eq:QWF}

$$Q_{\rm in}^{\rm WF} = \frac{-N_D + M_1 + M_2}{(n-1)N_D + M_1 + M_2},\tag{A49a}$$

$$Q_{\text{out}}^{\text{WF}} = \frac{-\frac{1}{N_D - 1} M_1 + M_2}{(n - 1)N_D + M_1 + M_2},\tag{A49b}$$

845 with

$$M_1 = \frac{N_D - 1}{1 - \frac{(1 - \mu)^2 (N_D (1 - m) - 1)^2}{(N_D - 1)^2}}$$
 and $M_2 = \frac{1}{1 - (1 - \mu)^2}$.

(These formulas are compatible with, *e.g.*, results presented by Cockerham & Weir (1987), adapted for haploid individuals).

In the Wright-Fisher life-cycle, $Q_{\rm in}^{\rm WF}$ decreases until $m=m_c^{\rm WF}=\frac{N_D-1}{N_D}$, while $Q_{\rm out}^{\rm WF}$ follows the opposite pattern. The threshold value $m_c^{\rm WF}$ corresponds to an emigration probability so high that $d_{\rm in}=d_{\rm out}$.

The two probabilities of identity by descent go to 1 when the mutation probability μ is very small ($\mu \to 0$), except if we first assume that the number of demes is very large ($N_D \to \infty$); for instance, with this life-cycle as well, $\lim_{N_D \to \infty} Q_{\text{out}}^{\text{WF}} = 0$.

Also, because more sites (all of them, actually) are updated at each time step, $Q_{\rm in}$ is lower for the Wright-Fisher updating than for a Moran updating, under which only one site is updated at each time step (compare figure A5(a) and A5(b)).

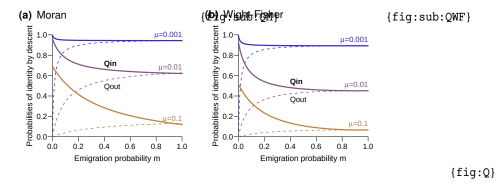


Figure A5: Probabilities of identity by descent, for two different individuals within the same deme $(Q_{\rm in},$ full curves) and two individuals in different demes $(Q_{\rm out},$ dashed curves), as a function of the emigration probability m, for different values of the mutation probability μ (0.001, 0.01, 0.1), and for the two types of life-cycles ((a): Moran, (b): Wright-Fisher). Other parameters: n=4 individuals per deme, $N_D=15$ demes.

Combining the formulas presented in eq. (A49), we obtain

$$R^{\text{WF}} = \frac{(1 - N_D(1 - m))^2 (1 - \mu)^2}{D^{\text{WF}}},$$
 (A50) {eq:app:RWF}

860 with

$$\begin{split} \mathbf{D}^{\mathrm{WF}} = & 1 - N_D(2(1+m(n-1)) - N_D(1+(2-m)m(n-1))) - 2\mu \\ & + 2(N_D(N_D(1-m)-2)(1-m)(n-1) + n)\mu - (1-N_D(1-m))^2(n-1)\mu^2. \end{split}$$

When the number of demes is very large and mutation is vanishingly small,

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$$\lim_{\mu \to 0} \lim_{N_D \to \infty} R^{\rm WF} = \lim_{N_D \to \infty} \lim_{\mu \to 0} R^{\rm WF} = \frac{(1-m)^2}{1 + (2-m)m(n-1)}. \tag{A51} \quad \{eq:app:RWFlim\}$$

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