

How single-locus F_{ST} varies with migration and selection for two populations in migration-selection balance

Abstract

A complete understanding of the role of natural selection in driving evolutionary change requires accurate estimates of the strength of selection acting at the genetic level in the ! This is challenging to achieve, but may be easier in the case of populations in migration-selection balance. When two populations are at equilibrium under migration-selection balance, there exist alleles selected different ways in the two populations. Such alleles can be identified from genome sequencing by their high values of single-locus F_{ST} . Intuitively it is clear that F_{ST} is increased by the strength of selection in each population but eroded by migration between them. Here we analyse a deterministic 1-locus 2-allele model of a large population distributed between two niches, and use this model to show graphically how F_{ST} increases with selection coefficients for given migration rates and dominance relationships between the alleles. At higher values of F_{ST} , possible values of selection coefficients are restricted to lie on contours on the $F_{ST}(s_1, s_2)$ surface. F_{ST} is here calculated from equilibrium allele proportions, and we also provide equations showing the general relationships between equilibrium allele proportions, selection coefficients, migration rates and dominance. We hope our methods may help those studying migration-selection balance to quantify the advantages conferred by adaptive genes.

| —
| the loci are deleted by F_{ST}

Keywords: migration-selection balance; dispersal; population genetics; ecological genomics; F_{ST} ;

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Introduction

A complete understanding of the role of natural selection in driving evolutionary change requires accurate estimates of the strength of selection acting at the genetic level in the wild, but until recent advances in molecular population genetics, measuring natural selection at the genetic level has been challenging (Linnen and Hoekstra 2009; Thurman and Barrett 2016). (Thurman and Barrett 2016) located 79 papers that used molecular techniques to study natural selection acting at the genetic level in natural populations, from which the importance of genomic data is clear, though variation in time and space can complicate tracking the strength of selection (Rudman et al. 2018). Several methods for inferring the strength of selection from ~~gen~~ frequency data have been developed in recent years (Tataru et al. 2017). Starting with the Wright–Fisher model of the effects of random genetic drift in a randomly mating population of finite size, several approaches have used the diffusion approximation to estimate the effects of various combinations of mutation, migration and selection on how allele frequencies change over the generations. For example (Vitalis et al. 2014) introduced a method extending the diffusion approximation of genetic drift in the migration-drift equilibrium island model to allow for the effects of selection. When applied to analysis of selection on the lactase-producing gene LCT (Vitalis et al. 2014)'s method showed that the strongest selection coefficients occurred in Europe and the Indus Valley, where scaled selection coefficients ranged up to 100. Scaled selection coefficients are defined as the product of selection coefficient and deme population size, but their effects can generally not be separated: to estimate the selection coefficient further information is needed on deme population size. Allele frequencies under the Wright–Fisher model can be approximated as time goes to infinity, but this requires that the scaled parameters relating to migration, mutation and selection are ~~in~~ on the order of 1 (Tataru et al. 2017).

Studies such as these start with a model of genetic drift and require complex mathematical development. Some simplification can be achieved if the starting point is instead a large population at equilibrium, so that drift can be ignored (Haldane 1930; Wright 1931). Starting from the models of (Wright 1931) and (Haldane 1930) of populations in migration-selection balance, (Hoekstra et al. 2004) derived for a simple 2-allele 2-population ~~model~~ an equation that allows estimation of the selection coefficient against a deleterious allele from its equilibrium frequency in each of two populations, the level of dominance and the migration rates between the populations. Using the equation (Hoekstra et al. 2004) estimated the selection coefficient acting on the *Mc1r* gene, which codes for coat colour, in populations of pocket mice living on black lava and on neighbouring light rocks. Migration rate was not estimated independent of population size, but was believed to be between 9.6×10^{-4} and 9.6×10^{-3} , giving the selection coefficient against the light-coat allele on the black larva between 0.039 and 0.39. Here we use a similar approach to study locally adapted populations at equilibrium held in a balance, with selection acting in different directions in different populations. This results in genetic differentiation between the populations if migration rates between the populations are sufficiently low (the term 'migration' is here used as a synonym of 'dispersal'). We show how if drift can be ignored, genomic studies of local adaptation can be used to estimate selection coefficients acting on the individual genes which together confer local adaptation.

Genetic differentiation between populations is generally summarised by F_{ST} (Whitlock 2011), and (Beaumont and Nichols 1996) showed how F_{ST} can be used in genomic studies to identify loci responsible for local adaptations by their relatively high values of F_{ST} (for recent examples see, e.g., (Flanagan and Jones 2017; Graham et al. 2018; Savolainen et al. 2013)). Loci not under selection have relatively low values of F_{ST} and information from these loci can provide estimates of migration rates between niches. As an example consider (Graham et al. 2018)'s study of high and low altitude populations of the speckled teal (*Anas flavirostris*) in South America, which used genome sequencing and F_{ST} analysis to identify 'outlier' genes selected in opposite ways in the two populations. The outlier genes had F_{ST} values in the range 0.44 – 0.77, and in some cases it was possible to identify the functions of the outlier genes that are adaptive at high altitude. The remaining non-outlier genes had F_{ST} values around 0.05 which, when further analysed suggested < 3% migration from low to high



altitude, and less from high to low. This genomic study of local adaptation identified loci responsible for local adaptations by their high values of F_{ST} , and genomic variation in unselected regions of the genome to estimate migration rates between niches. High values of F_{ST} occur in genes responsible for local adaptation in large populations at equilibrium, but F_{ST} may also be high in populations not at equilibrium, in which evolution is still under way, in ways that depend on the spatial structure of the environment (e.g., island model, isolation-by-distance, range expansion) (Lotterhos and Whitlock 2014; Lotterhos and Whitlock 2015). The methods developed in the present paper assume that populations are at equilibrium.

F_{ST} is calculated from measurements of the proportion of alleles of each type that are present in each population. Intuitively, it is clear that F_{ST} is increased by the strength of selection in each population but eroded by migration between them, but a method is needed to quantify the relationship. As a first step we show here how (Hoekstra et al. 2004)'s equation for a deterministic 1-locus 2-allele model of a large population distributed between two niches can be corrected and extended to show how selection coefficients can be obtained for both niches. The population is in migration-selection balance. Account is taken of migration rates and levels of dominance, and mating takes place within niches as in (Bulmer 1972). The development assumes equal population sizes in the two niches, but is easily modified for the general case. We show graphically how F_{ST} increases with selection coefficients for given migration rates and dominance relationships between the alleles. At higher values of F_{ST} possible values of selection coefficients are restricted to lie on contours on the $F_{ST}(s_1, s_2)$ surface. F_{ST} is here calculated from equilibrium allele proportions, and we also provide equations for the general relationships between equilibrium allele proportions, selection coefficients, migration rates and dominance. Without these equations equilibrium proportions can only be ascertained by simulation of the evolutionary process, as in (Sibly and Curnow submitted).



Theory and methods

Population genetic model

Our model has one locus with two alleles P and Q in an environment consisting of two niches with some migration between niches prior to mating, as depicted in Fig. 1. The locus determines ecological adaptation to one niche or the other. The fitness of the three genotypes in each niche is shown in Table 1.

Genotype	Niche 1			Niche 2		
	PP	PQ	QQ	PP	PQ	QQ
Fitness	1	$1+hs_1$	$1+s_1$	1	$1+hs_2$	$1+s_2$
Relative frequency	p_1^2	$2p_1q_1$	q_1^2	p_2^2	$2p_2q_2$	q_2^2

} a symmetric parameterisation might be preferable

Table 1. The fitness of the three genotypes in each niche. p_1 and q_1 represent the relative frequencies of the P and Q alleles at the start of a generation in niche 1, their frequencies in niche 2 are p_2 and q_2 . Carriers of the QQ genotype obtain fitnesses $1 + s_1$ and $1 + s_2$ in niches 1 and 2 respectively, PP homozygotes have fitness 1 in both niches. Parameter h indicates the level of dominance of the Q allele.

We assume a large population so that the dynamics are deterministic. Generations are discrete and individuals die after mating. The life histories occur in the following order. At the start of each generation individuals in each niche mate at random, and all mating individuals obtain the same number of offspring. The number of offspring of each genotype that survive in each niche is the product of its initial frequency and its fitness. Population regulation then returns population numbers to their initial values so that both niches have the same number of individuals. Finally some individuals migrate between niches, as shown in Fig. 1, leading to the start of the next generation.

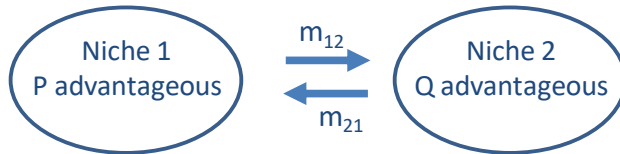


Fig. 1. Conceptual overview of the model. For clarity, the equal-size niches are shown distinct, but in nature may be contiguous or overlap. m_{12} and m_{21} specify the proportion of individuals in one niche that migrate to the other each generation after viability selection and population regulation have occurred. When analysing the model we suppose that Q is disadvantageous in niche 1 (i.e., s_1 is negative) but advantageous in niche 2 (i.e., s_2 is positive), while PP homozygotes have fitness 1 in both niches. There are no sex differences in fitnesses or migration rates.

The convention is to denote by m_{ij} the proportion of individuals in popul. i replaced by immigrants from popul. j, i.e. the backward migr. rate.

Recurrence equations giving the frequencies of the P allele in successive generations

We now derive recurrence equations for the above model that give the frequencies of the P allele in successive generations in each niche. The relative frequencies of P and Q at the start of a generation in the two niches are given in Table 1. The relative frequencies of P in the two niches after selection are:

$$p_1' = (p_1^2 + (1+hs_1)p_1q_1)/(1+2hp_1q_1s_1+s_1q_1^2) \quad (1a)$$

$$p_2' = (p_2^2 + (1+hs_2)p_2q_2)/(1+2hp_2q_2s_2+s_2q_2^2) \quad (1b)$$

If you defined m_{ij} as the backward migration rate, rather than the forward migration rate (see Fig. 1), then I think you can use a simpler equation:

The relative frequencies of P after migration are: $p_1'' = (1 - m_{12}) p_1' + m_{12} p_2'$

$$p_1'' = \frac{(1-m_{12})p_1' + m_{21}p_2'}{1+m_{21}-m_{12}} = ((1-m_{12})p_1' + m_{21}p_2')/c_1 \quad (2a)$$

$$p_2'' = \frac{(1-m_{21})p_2' + m_{12}p_1'}{1+m_{12}-m_{21}} = ((1-m_{21})p_2' + m_{12}p_1')/c_2 \quad (2b)$$

where p_1' and p_2' are obtained from equations 1, $c_1 = 1 + m_{21} - m_{12}$ and $c_2 = 1 + m_{12} - m_{21}$

The equations show how the frequencies of P in the two niches in the next generation, p_1'' and p_2'' , can be derived from the frequencies in the present generation, p_1 and p_2 .

Calculation of selection coefficients from equilibrium values of p_1 and p_2 and migration rates

The recurrence equations derived in the previous section can be used to analyse what happens when the population reaches equilibrium. At equilibrium $p_1''=p_1$ and $p_2''=p_2$, so from equations 2:

$$p_1' = \frac{c_1 p_1 - m_{21} p_2'}{1-m_{12}} = \frac{c_2 p_2 - (1-m_{21}) p_2'}{m_{12}} \Rightarrow m_{12} [c_1 p_1 - m_{21} p_2'] = (1-m_{12}) [c_2 p_2 - (1-m_{21}) p_2'] \quad (3)$$

$$\Rightarrow m_{12} m_{21} p_2' - (1-m_{12})(1-m_{21}) p_2' = m_{12} c_1 p_1 - (1-m_{12}) c_2 p_2$$

If m_{12} , m_{21} and equilibrium values of p_1 and p_2 are known, then equation 3 can be rearranged to obtain:

$$p_2' = \frac{m_{12}(c_1 p_1 + c_2 p_2) - c_2 p_2}{m_{12} + m_{21} - 1}, = K_2, \text{ say.} \quad (4a)$$

$$\text{Similarly } p_1' = \frac{m_{21}(c_1 p_1 + c_2 p_2) - c_1 p_1}{m_{12} + m_{21} - 1}, = K_1, \text{ say.} \quad (4b)$$

Setting this equal to p_2' as given by equation 1b we obtain:

$$s_2 = \frac{p_2 - K_2}{K_2 q_2^2 + h p_2 q_2 (2K_2 - 1)} \quad (5a)$$

$$s_1 = \frac{p_1 - K_1}{K_1 q_1^2 + h p_1 q_1 (2K_1 - 1)} \quad (5b)$$

where K_1 and K_2 are given by equation 4. When m_{12} , m_{21} and h are known, uncertainty in the estimates of s_1 and s_2 can be quantified because the estimates of p_1 and p_2 are binomially distributed.

Equations 5 show how selection coefficients s_1 and s_2 can be calculated from measurements of m_{12} , m_{21} , h and equilibrium values of p_1 and p_2 .

Calculating equilibrium proportions of alleles in each population when selection coefficients are known

We now turn to the inverse problem of calculating equilibrium proportions of alleles in each population when selection coefficients are known. From equation 3:

$$p_1 = \frac{m_{12} + m_{21} - 1}{c_1 m_{12}} p_2' + \frac{1 - m_{12}}{c_1 m_{12}} c_2 p_2 \quad (6)$$

Substituting for p_2' from equation 1b gives an equation for p_1 in terms of p_2 , m_{12} , m_{21} , h , s_1 and s_2 :

$$p_1 = \frac{m_{12} + m_{21} - 1}{c_1 m_{12}} \left(\frac{p_2 (1 + h s_2 q_2)}{1 + 2 h s_2 p_2 q_2 + s_2 q_2^2} \right) + \frac{1 - m_{12}}{c_1 m_{12}} c_2 p_2 \quad (7a)$$

The analogous equation for p_2 is:

$$p_2 = \frac{m_{12} + m_{21} - 1}{c_2 m_{21}} \left(\frac{p_1 (1 + h s_1 q_1)}{1 + 2 h s_1 p_1 q_1 + s_1 q_1^2} \right) + \frac{1 - m_{21}}{c_2 m_{21}} c_1 p_1 \quad (7b)$$

Substituting p_2 from equation 7b into equation 7a yields an equation in p_1 , m_{12} , m_{21} , h , s_1 and s_2 which can be solved to obtain equilibrium values of p_1 for given values of m_{12} , m_{21} , h , s_1 and s_2 .

Perhaps show figure illustrating the convergence.

Equilibrium values of p_2 can be obtained similarly. Equations 7 show how equilibrium proportions of alleles in each population can be calculated when selection coefficients are known.

Finding evolutionary outcomes by simulation of the evolutionary process

Equations 7 do not provide explicit expressions for the equilibrium values of p_1 and p_2 . An alternative to using an equation solver to obtain equilibrium values is to simulate the evolutionary process for specified values of h , m_{12} , m_{21} , s_1 and s_2 using the recurrence equations 1 and 2. These relate allele frequencies in one generation to their frequencies in the preceding generation. The simulations whose equilibrium values are presented here were started at an initial frequency of Q of 0.01 in niche 2 and continued to equilibrium as determined by visual inspection of trajectories of allele frequencies. Equilibrium was generally reached within a few hundred generations but took over 10,000 generations for $(s_1, s_2) = (0.01, 0.01)$, $(0.01, 0.1)$ and $(0.1, 0.01)$. Checks showed that the simulated evolutionary outcomes satisfied equations 7.

Calculation of F_{ST}

F_{ST} was calculated from allele frequencies using the equation $F_{ST} = \sigma_S^2 / \sigma_T^2$, where σ_S^2 and σ_T^2 represent the variances of an allele's frequency between subpopulations, and in the total population, respectively (Holsinger and Weir 2009). The variance of p , the frequency of an allele in a population of size n , is given by the binomial distribution as npq . So $F_{ST} = (p_T q_T - \frac{1}{2} p_1 q_1 - \frac{1}{2} p_2 q_2) / p_T q_T$, where subscripts T, 1 and 2 refer to the total population and the populations in niches 1 and 2 respectively. F_{ST} can therefore alternatively be thought of in terms of the average frequency of heterozygotes in the two populations compared with the frequency of heterozygotes if there was random mating between all the individuals in the two populations. F_{ST} was assigned the value 0 when one allele became fixed in both niches because F_{ST} tends to 0 as any one allele tends to fixation.

Visualisation of the function $F_{ST} = F_{ST}(m, s_1, s_2)$ when $m_{12} = m_{21} = m$

In this section we show how equilibrium values of F_{ST} are related to the migration rates between the niches, and the strengths of selection s_1 and s_2 within them. We assume from now on that migration rates are the same in each direction, so that $m_{12} = m_{21} = m$. We begin by showing how genetic differentiation, measured by F_{ST} , is eroded by migration when selection is maximal, total selection one way in niche 1 and the other way in niche 2 (Fig. 2). Maximal selection produces maximum values of F_{ST} , and these decline as migration rates increase, from a maximum of 1 when populations are isolated, to around 0.4 when migration rates are 10% and down to zero when migration rate is 50% (Fig. 2). The maximum values of F_{ST} are a little higher for $h=0.5$ than for $h=1$ (dominance) when migration rates are less than 15%, but otherwise similar.

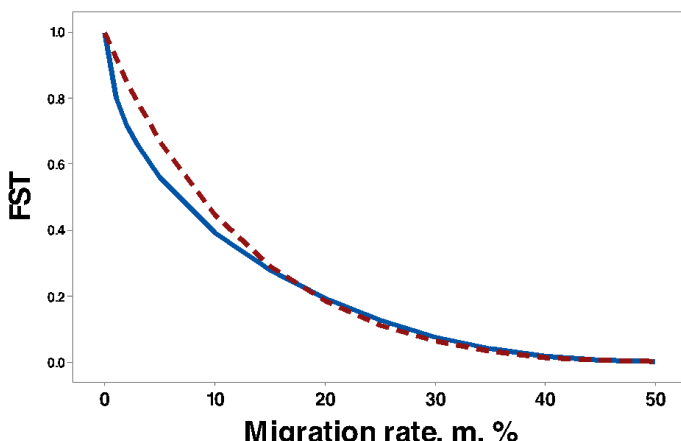


Fig. 2. The maximum values of F_{ST} in relation to migration rate, m . Maximum values of F_{ST} occur when selection on QQ is total one way in niche 1 and the other way in niche 2, and this was approximated by setting s_1 at -1 and s_2 at 10^9 . Solid blue curve is for $h=1$, dashed red curve $h = 0.5$. Values of F_{ST} were calculated for populations at equilibrium as determined by simulation of evolutionary trajectories of allele frequencies. Equilibrium judged by eye was generally achieved within 100 generations. The table of F_{ST} vs. m from which the figure was constructed is given in Supplementary Materials.

To visualise the effects of lower values of s_1 and s_2 on F_{ST} we calculated equilibrium values of F_{ST} over a grid of values of m , s_1 and s_2 to obtain visualisations of the function $F_{ST} = F_{ST}(m, s_1, s_2)$, and these are presented in Fig. 3 for two levels of dominance h . The panels in the rows of Fig. 3 represent visualisations of the function $F_{ST} = F_{ST}(m, s_1, s_2)$ for three values of migration rate m . The highest values of F_{ST} are achieved when migration rates are lowest (left-hand panels of Fig. 3). Within each panel the highest value of F_{ST} occurs when selection is at its strongest, for Q in niche 2 ($s_2 = 1$) and against Q in niche 1 ($s_1 = -1$). If selection is too low, then either P or Q go to fixation and F_{ST} goes to zero. At other values of s_1 and s_2 , the equilibrium is a polymorphism and $F_{ST} > 0$. The shapes of the F_{ST} surfaces for $h = 1$ (top row of Fig. 3) and $h = 0.5$ (bottom row) are qualitatively similar but they differ a little in quantitative detail. The F_{ST} surfaces shown in Fig. 3 are not perfectly symmetrical about the $s_2 = -s_1$ plane, because $s_1 = -1$ represents total selection against QQ in niche 1 but $s_2 = +1$ does not represent total selection for QQ in niche 2; that is achieved when $s_2 = \infty$.

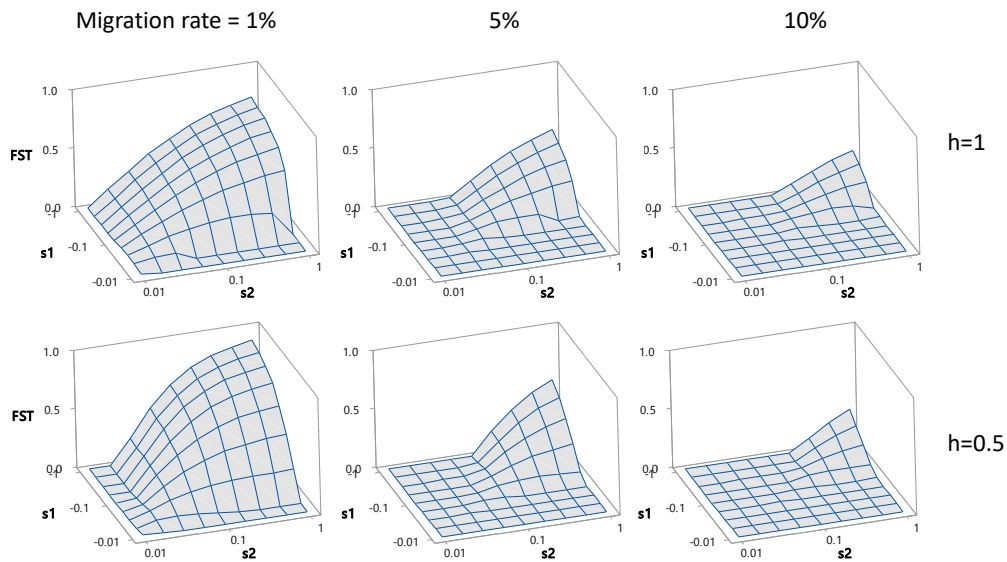


Fig. 3 Values of F_{ST} in relation to selection coefficients s_1 and s_2 for three rates of migration between niches m , and two levels of dominance h . Top row: Q is dominant (i.e. $h = 1$); bottom row $h = 0.5$. Values of F_{ST} were calculated for populations at equilibrium as determined by simulation of the evolutionary process. The tables of s_1 , s_2 and F_{ST} from which these figures were constructed are given in Supplementary Materials.

If F_{ST} and migration rates are known, Fig. 3 can be used to put bounds on selection coefficients. For a given value of migration rate, possible values of s_1 and s_2 for a given value of F_{ST} lie on the relevant contour in Fig. 3. Contours are shown in Fig. 4 for the case that migration rate = 1% and $h=0.5$. From the contour corresponding to a given value of F_{ST} , the possible determining values of s_1 and s_2 can now be identified. If $F_{ST} = 0.7$ and selection pressures are equal, for instance, then s_2 and $|s_1|$ must

be around 0.25, but if they are not equal then the value of one or other of them could be as low as 0.15.

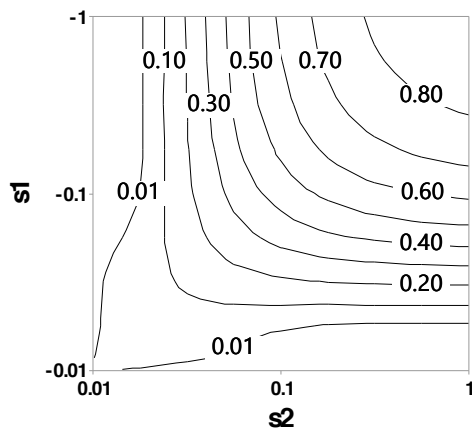


Fig. 4. Contours of F_{ST} in relation to selection coefficients s_1 and s_2 for the case that migration rate = 1% and $h=0.5$. These are the contours for the lower left panel in Fig. 3.

Discussion

Methods with which to estimate selection coefficients are needed to further understanding of evolutionary processes in wild populations. The required measurements can be made relatively easily in locally adapted populations if the populations can be assumed to be in equilibrium. Here, using a two-niche two-allele model, we show in equations 5 how selection coefficients s_1 and s_2 can be estimated from measurements of migration rates m_{12} and m_{21} , levels of dominance h and the equilibrium values of allele proportions in the two habitats, p_1 and p_2 . Because F_{ST} is generally reported rather than allele proportions, we analysed the relationship between F_{ST} and selection coefficients, and present the results graphically in Figs. 2 – 4 as visualisations of the function $F_{ST} = F_{ST}(m, s_1, s_2)$ when migration rates m between the two niches are equal. These visualisations show how equilibrium values of F_{ST} are related to the migration rates between the niches, and the strengths of selection s_1 and s_2 within them. F_{ST} declines as migration rates increase (Figs. 2 and 3). If F_{ST} and migration rates are known, Figs. 3 and 4 show how bounds can be put on possible values of selection coefficients. The analysis presented assumes that after population regulation the populations have equal size. If instead the sizes of populations 1 and 2 are M_1 and M_2 respectively then equations 2 have to be modified with m_{21} replaced by $m_{21}M_2/M_1$ in equation 2a, and m_{12} replaced by $m_{12}M_1/M_2$ in equation 2b, with knock-on modifications required in subsequent equations.

Our model agrees with that of Hoekstra et al. (2004) in modelling selection and corrects an error in Hoekstra et al. (2004)'s model of migration. Applied to a single niche our model is identical to that of Hoekstra et al. (2004) in the way selection is modelled: the first term in Hoekstra et al. (2004)'s equation 1 is equivalent to our equation 1a. In modelling migration Hoekstra et al. (2004) assume that the frequency of migrants is the frequency before selection. Selection will act on migrants as well as residents, but this is not accounted for in Hoekstra et al. (2004)'s equation 1.

Inferences in practice will need to take account of several caveats. Our calculations are for large populations at equilibrium, but in the real world selection pressures, migration rates and population

This is a well-established fact in the field, right?

What's the error?

sizes may vary over time, so that F_{ST} and other variables vary too. The methods presented here need extension to take account of the effects of population size and drift, perhaps using approaches such as those reviewed by Tataru et al (2017). Furthermore, loci may become differentiated between populations not because they are themselves selected, but because they are physically close to loci that are selected. This phenomenon of linked selection is the subject of ongoing research (e.g., Burri 2017).

All the results presented here show some dependence on levels of dominance. Although there is interest in the evolution of dominance, little is known of values of levels of dominance in natural populations (Huber et al. 2018; Thurman and Barrett 2016) except that overdominance is infrequent (Thurman and Barrett 2016). Here in Figs. 2 and 3 we present evolutionary outcomes for what are, in the absence of overdominance, the extreme values 0.5 and 1. Figs. 2 and 3 suggest that in the absence of overdominance, levels of dominance have only small effects on equilibrium values of F_{ST} .

The method of calculating selection coefficients using equation 5 requires knowledge of migration rates, the allele proportions in the two habitats and the level of dominance. Migration rates and allele proportions are routinely measured in genomic studies. While levels of dominance likely have only small effects, it is worth considering whether selection coefficients and the level of dominance at individual loci could be measured directly in genomic field studies. If within each population individuals mate at random, as in the model presented here, then the genotypes of the offspring will be initially at each locus in Hardy-Weinberg proportions. The offspring are then subject to viability selection which changes the genotype proportions. Comparison of genotype proportions in offspring and adults could provide information about selection coefficients and levels of dominance directly. Suppose, for instance, that at a locus of interest an allele P initially has relative frequency p so that the initial proportions of offspring genotypes PP, PQ and QQ are as in Table 1, $p^2: 2pq: q^2$, then if there are no QQ genotypes in the adults and the ratio of PP to PQ is $p^2: 2pq$, it follows that P is dominant and there is complete selection against QQ, i.e., $h = 0$ and $s = -1$. In general, if the ratios of the genotypes in adults are $a: b: c$ then $s = \frac{cp^2}{aq^2} - 1$ and $h = (\frac{b}{2pq} - 1)/s$. In this way selection coefficients and levels of dominance could be measured directly in field studies.

If the strength of selection decreases or migration increases there comes a point at which both alleles are not retained, and one or the other goes to fixation, and F_{ST} declines to zero. This point represents the boundary of what Sibly and Curnow (submitted) termed the 'polymorphism set' within which stable genetic polymorphisms occur. The polymorphism set is equivalent to the set of non-zero values of F_{ST} , and its boundaries define the limits to local adaptation. The polymorphism set is of particular interest because it can be the starting point for speciation, given a mechanism for phenotype matching and the emergence of a matching cue (Sibly et al. 2019). The boundaries of the polymorphism set were found by (Sibly and Curnow submitted) by simulation of the evolutionary process, but could alternatively be found by solving the two non-linear simultaneous equations 7.

Given current increase in studying local adaptation using ecological genomics, we hope our methods may, when migration rates are known, help quantify the advantages conferred by adaptive genes.

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