Background selection and population differentiation

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Abstract

A general analytical formula is derived, which predicts the effects of background selection on population differentiation at a neutral locus as a result of its linkage with selected loci of deleterious mutations. The theory is based on the assumptions of random mating, multiplicative fitness, and weak selection in hermaphrodite plants in the island model of population structure. The analytical results show that $F_{st}$ at the neutral locus increases as a result of the effects of background selection, regardless of the dependence or independence among linked background selective loci. The increment in $F_{st}$ is closely related to the magnitude of linkage disequilibrium between the neutral locus and selected loci, and can be estimated by the ratio of $F_{st}$ with background selection to $F_{st}$ without background selection minus one. The steady-state linkage disequilibrium between a neutral locus and a selected locus in subpopulations, primarily attained by gene flow, decreases with the recombination rate, and can be enhanced when there are dependence among linked selected loci. Monte Carlo computer simulations with two- and three-locus models show that the analytical formulae perform well under general conditions. Application of the present theory may aid in analyzing the genome-wide mapping of the effect of background selection in terms of $F_{st}$.

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1. Introduction

Like the selectively favored mutations that cause hitchhiking effects on linked neutral loci (Maynard Smith and Haigh, 1974), selectively disfavored mutations can also change gene frequencies and reduce genetic diversities at linked neutral loci (“background selection”, Charlesworth et al., 1993). Early studies showed that a substantial reduction in genetic diversity at a neutral locus can result from its linkage to deleterious mutations (e.g., Charlesworth et al., 1993; Hudson and Kaplan, 1995; Nordborg et al., 1996). The genetic basis for maintaining both kinds of effects is the persistence of the linkage disequilibrium (LD) between neutral and selected loci (see the review by Barton, 2000). When LD equals zero, both kinds of effects disappear.

In a natural population without subdivision the LD between two linked neutral loci dissipates with generation as the consequence of recombination, and eventually approaches zero (e.g., Bennett, 1954; Hill and Robertson, 1968; Hill, 1974). In the population with subdivision the dissipation of global LD with generation is enhanced since the inter-subpopulation gene flow can reduce the effective size of the whole population and hence increase the drift speed (Wright, 1943). However, a certain amount of LD in local subpopulations can be attained owing to the effects of inter-subpopulation gene flow that counteracts genetic drift. Stable LD between selected nuclear loci without epistasis can be maintained in subdivided populations (Li and Nee, 1974). When the recombination fraction between selected nuclear loci is of the same order or smaller than the selection coefficient, a substantial amount of LD can be present in a cline (Slatkin, 1975). The LD between selected
nuclear and cytoplasmic loci that are physically unlinked can even be generated when inter-population gene flow (seed and pollen flow) takes place (e.g., Hu and Li, 2002).

Similarly, the persistence of the LD between one neutral locus and another selected locus is expected in a local population owing to the inter-subpopulation gene flow. As long as a certain amount of LD between neutral and selected loci is preserved, the effect of background selection should be present. The persistence of LD can cause an increase in variance of neutral allele frequencies and hence increase its population differentiation (Barton, 2000).

Previous LD studies are often examined in terms of genes or molecular markers as an “observation unit” at the equilibrium between gene flow and genetic drift. At a fine scale, the length of genomes for maintaining a certain amount of LD can be long in terms of the number of base pairs and so is the length within which background selection has a significant effect. It is meaningful to examine the effect of background selection in terms of single nucleotide polymorphisms (SNP) as an observation unit-marker. For example, 1 percent of recombination fraction (1 centiMorgan or cM) is equal to 1 million base pairs on the physical map in human genomes and contains about 1000 SNP (e.g., Wang et al., 1998). Within a few cMs of genetic distance the LD between one selected nucleotide site and another neutral site is likely substantial, and the effect of background selection on population differentiation at the individual neutral sites can be significant. Evidence indicates that high LD may extend over several centiMorgans in cattle and human genomes (e.g., Farnir et al., 2000; Abecasis et al., 2001). Needless to say, LD distribution along genomes varies with populations (e.g., Goddard et al., 2000; Shifman et al., 2003).

SNP are abundant in various organisms, such as in Arabidopsis thaliana and the rice genome (see the review by Rafalski, 2002). The genetic diversities of SNP within either the coding or non-coding regions of a gene are affected by their physical distances from the selected sites that may be located within the same gene or in the regions of other genes. For multiple linked genes with unequal numbers of SNP, the spatial pattern of genetic diversity across SNP could exhibit a patchy pattern along chromosomes. These naturally occurring patterns of SNP diversity provide a tool for mapping the effect of background selection in terms of population differentiation.

The purpose of this study is to develop further population genetic theory required for understanding the effect of background selection on population differentiation at a neutral locus. Although the effect of background selection on genetic diversity of a neutral gene has widely been appreciated, the theoretical investigation of such effect on population differentiation is much less fully explored (Charlesworth et al., 1997; Nordborg, 1997; Barton, 2000). In this study, we analytically derive the population differentiation at a neutral site due to its linkage with the sites that are subject to unfavorable mutations. Computer simulations are conducted to validate the analytical results that demonstrate the increase in $F_{st}$ owing to the effect of background selection.

2. Assumptions

Based on the classical island model of population structure (Wright, 1969), here we consider diallelic selected nuclear loci (diploid) that are linked with a neutral locus in a hermaphrodite population of plants. For simplicity the selected loci addressed throughout this study refer to those with selectively disfavored mutation. Weak selection is considered in modeling so that all terms containing the second or higher order of selection coefficient are neglected. Like Nordborg et al. (1996), the selected loci are subject to a balance of mutation–selection–migration, and genetic drift effects are assumed negligible. The dependence among selected loci, caused by gene flow, is considered, relaxing the independence assumption made by Nordborg et al. (1996) and Hudson and Kaplan (1995).

The modeling procedure is based on a sequence of events in the life cycle of hermaphrodite plants: pollen flow, random combination between pollen and ovules (random mating), seed flow, mutation, natural selection, genetic drift, and next adults. This procedure is similar to Hu and Ennos (1999) except that mutation and background selection are included and also similar to Nordborg et al. (1996) except that migration is considered. The gene frequencies in migrants of pollen grains or seeds are equal to the average of gene frequencies over all subpopulations. The gene frequency in ovules before random combination with pollen grains is assumed to be the same as that in the preceding generation.

In the following we first derive the change of gene frequency at a neutral locus as a result of linkage to one and two selected loci, and then give a general expression for the change due to the background selection from an arbitrary number of selected loci. Wright’s $F_{st}$ is then employed to describe the population differentiation.

3. Allele frequency

3.1. Two-locus case

Consider a selected locus $A$ that is linked to a neutral locus $C$ in the $i$th subpopulation. The wild-type allele at the $A$ locus is denoted by $A_i$, and its mutant allele by $a_i$;
their frequencies are \( p_{A_i} \) and \( p_{a_i} \) \((p_{A_i} + p_{a_i} = 1)\), respectively. Let the mutation rate from the wild-type allele to the mutant allele be \( u_i \) at the \( A \) locus. The fitness of genotypes is assumed to be 1, 1–\( s_{i1} \), and 1–2\( x_{i1} \), for the genotypes of \( AA_C_i \), \( Aa_C_i \), and \( aa_C_i \), respectively. The migration rates of pollen and seeds into each subpopulation are denoted by \( m_p \) and \( m_s \), respectively. According to the life cycle mentioned in the assumptions, the change in the allele frequency at the \( A \) locus is given by

\[
\Delta p_{A_i} = p_{A_i}p_{a_i}s_{i1} - u_ip_{A_i} - \bar{m}(p_{A_i} - \bar{p}_{A_i}).
\]

where \( \bar{m} = m_s + m_p/2 \), \( \bar{p}_{A_i} \) is the frequency of the allele \( A_i \) in migrants (seeds and pollen grains). This equation can also be implied from Wright’s general expression (Wright 1969, p. 474). The first term on the right-hand side represents the decrease in frequency due to mutation and selection, and the second term is the reduction due to migration. At steady state, \( \Delta p_{A_i} = 0 \), the allele frequencies at the \( A \) locus can be analytically solved from Eq. (1).

\[
p_{A_i} = \frac{(s_{i1} - u_i - \bar{m}) \pm \sqrt{(s_{i1} - u_i - \bar{m})^2 + 4s_{i1}\bar{m}\bar{p}_{A_i}}}{2s_{i1}} \quad \text{with the condition of } 0 \leq p_{A_i} \leq 1.
\]

Consider the neutral locus that has alleles \( C_i \) and \( c_i \) in the \( i \)th subpopulation. Let the mutation rate from \( C_i \) to \( c_i \) be \( v_i \). There are four types of two-locus gametes: \( A_iC_i, A_iC_i, a_iC_i, \) and \( a_iC_i \), with frequencies of \( P_{A_iC_i}, P_{A_iC_i}, P_{a_iC_i}, \) and \( P_{a_iC_i} \), respectively. According to Eqs. (A.3) and (A.4) in Appendix A, the steady state is the change due to migration (seed and pollen flow), the second term is the change due to the mutation of the allele \( C_i \) to other alleles, and the third term is the change due to the linkage to the selected \( A \) locus. If the linkage disequilibrium is of the order similar to the selection coefficient \( (s_{i1}) \), the third term on the right-hand side of Eq. (2) is negligible. Since LD is primarily generated by the inter-subpopulation gene flow, its magnitude can be much greater than the order of selection coefficient when the recombination fraction is very small, say within a few cMs of genome.

From the setting of the conditional probabilities of \( x_{0i} \) and \( x_{1i} \), \( D_{AC(0)} \) can be expressed by

\[
D_{AC(0)} = p_{A_i}p_{a_i}(x_{0i} - x_{1i}).
\]

When the neutral allele \( C_i \) is equally distributed under the mutant and mutant-free backgrounds of the \( A \) locus, i.e. \( x_{0i} = x_{1i} \), the effect of background selection equals zero \( (D_{AC(0)} = 0) \).

Let \( r_i \) be the recombination fraction between the \( A \) and \( C \) loci. Following the approach similar to Nordborg et al. (1996, p. 170), the changes in the conditional probabilities of \( x_{0i} \) and \( x_{1i} \) due to the joint effects of migration, selection, and mutation at the \( A \) locus are derived as Eqs. (A.3) and (A.4) in Appendix A. When there is no effect of migration, Eqs. (A.3) and (A.4) reduce to the previous results of Nordborg et al. (1996).

Let \( \Delta p_{C_i} \) be the change in the frequency of the neutral allele \( C_i \) due to the joint effects of background selection, mutation, and migration. According to Eqs. (A.3) and (A.4) in Appendix A and the relation of \( p_{C_i} = P_{A_iC_i} + P_{a_iC_i} \), the analytical expression for \( \Delta p_{C_i} \) is given by

\[
\Delta p_{C_i} = \Delta P_{A_iC_i} + \Delta P_{a_iC_i} = p_{A_i}\Delta x_{0i} + p_{a_i}\Delta x_{1i} = -\bar{m}(p_{C_i} - \bar{p}_C) - v_r p_{C_i} + x_{1i}D_{AC(0)},
\]

where \( D_{AC(0)} \) is the LD between the \( A \) and \( C \) loci in the \( i \)th subpopulation. The first term on the right-hand side of Eq. (2) is the change due to migration (seed and pollen flow), the second term is the change due to the mutation of the allele \( C_i \) to other alleles, and the third term is the change due to the linkage to the selected \( A \) locus. If the linkage disequilibrium is of the order similar to the selection coefficient \( (s_{i1}) \), the third term on the right-hand side of Eq. (2) is negligible. Since LD is primarily generated by the inter-subpopulation gene flow, its magnitude can be much greater than the order of selection coefficient when the recombination fraction is very small, say within a few cMs of genome.
3.2. Three-locus case

Assume that the neutral locus is linked on either side to a selected locus each with disfavored mutations. The difference between the two- and three-locus cases is that the effects of LD between selected loci generated by gene flow and the double crossover among the three loci are included. Assume that another diallelic selected locus \( B \) links to the neutral locus \( C \) at the opposite side to the \( A \) locus, i.e., the order of \( ACB \). The wild-type allele at the \( B \) locus in the \( i \)th subpopulation is denoted by \( B_i \), and its mutant allele is denoted by \( b_i \); their frequencies are \( p_{B_i} \) and \( p_{b_i} \), respectively. Let the mutation rate from the wild-type allele to the mutant be \( u_2 \) at the \( B \) locus. The fitness is assumed to be \( 1, 1 - s_2u_2, 1 - 2s_2u_2 \) for the genotypes of \( B_iB_i, B_ib_i, \) and \( b_ib_i \), respectively.

Using the assumption of multiplicative viability, the fitness for each two-locus genotype can be readily calculated. Following the life cycle the steady-state allele frequencies at the \( A \) and \( B \) loci at the balance of migration–selection–mutation are shown to have the following relations:

\[
p_{A_i} p_{a_i} s_{1i} - u_1 p_{A_i} - \hat{m}(p_{A_i} - p_A) - D_{AB(i)} s_{2i} = 0, \quad (7a)
\]

\[
p_{B_i} p_{B_i} s_{2i} - u_2 p_{B_i} - \hat{m}(p_{B_i} - p_B) - D_{AB(i)} s_{1i} = 0, \quad (7b)
\]

where \( \hat{m} \) is the frequency of the allele \( B_i \) in migrants (seeds and pollen grains), and \( D_{AB(i)} \) is the steady-state LD between the \( A \) and \( B \) loci.

The steady-state \( D_{AB(i)} \) can be obtained from Eq. (B.2) in Appendix B, that is

\[
D_{AB(i)} = \frac{\hat{m}(\hat{D}_{AB} + (p_{A_i} - \hat{p}_A)(p_{B_i} - \hat{p}_B))}{1 - (1 - \hat{m} - (p_{A_i} - p_A)s_{1i} - (p_{B_i} - p_B)s_{2i} - u_1 - u_2)(1 - r)},
\]

(8)

where \( r \) is the recombination rate between the \( A \) and the \( B \) loci. The analytical expressions for the allele frequencies at the \( A \) and \( B \) loci are hard to obtain using the joint Eqs. (7) and (8). In the specific case where allele frequencies at the \( A \) and \( B \) loci are coincident, i.e., \( s_{1i} = s_{2i} = s, u_1 = u_2 = u \), \( \hat{p}_A = \hat{p}_B = \hat{p} \), and \( p_{A_i} = p_{B_i} = p_i \), we obtained a cubic equation

\[
d_0 p_i^3 + d_1 p_i^2 - d_2 p_i - d_3 = 0,
\]

where

\[
d_0 = 4(1 - r)s^2,
\]

\[
d_1 = s(1 + \hat{m} - (1 - \hat{m} - 2u + 2s)(1 - r)
- 4(s - u - m)(1 - r)),
\]

\[
d_2 = (1 - (1 - \hat{m} - 2u + 2s)(1 - r))(s - u - \hat{m})
+ 4s\hat{m}(1 - r)\hat{p} + 2s\hat{m}\hat{p}
\]

and

\[
d_3 = \hat{m}((1 - (1 - \hat{m} - 2u + 2s)(1 - r))\hat{p} - s_i\hat{D}_{AB} - sp_i^2).
\]

Solutions to Eq. (9) can be calculated with the Mathematica tool.

There are eight types of three-locus gametes: \( A_iC_iB_i \), \( A_iC_iB_i \), \( A_iC_iB_i \), \( A_iC_iB_i \), \( A_iC_iB_i \), \( A_iC_iB_i \), \( A_iC_iB_i \), and \( A_iC_iB_i \). Let the probability that the allele \( C_i \) is linked with a mutant-free background of gametes with respect to the two selected loci, \( p(C_i|A_iB_i) \), be \( y_{0i} \). Similarly, let \( p(C_i|A_iB_i) = y_{1i} \), \( p(C_i|A_iB_i) = y_{2i} \), and \( p(C_i|A_iB_i) = y_{3i} \). All these conditional probabilities are in the range of 0 to 1 \((0 \leq y_{ji} \leq 1, j = 0, 1, 2, 3)\). The frequencies of the three-locus gametes \( A_iC_iB_i \) and \( A_iC_iB_i \) can be written as \( p_{A_iC_iB_i} = p_{A_iB_i} y_{0i} \), and \( p_{A_iC_iB_i} = p_{A_iB_i}(1 - y_{0i}) \), respectively, where \( p_{A_iB_i} \) is the frequency of gamete \( A_iB_i \). The expressions for the remaining six three-locus gametes \( A_iC_iB_i \), \( A_iC_iB_i \), \( A_iC_iB_i \), \( A_iC_iB_i \), and \( A_iC_iB_i \) can be written in a similar way. The conditional probabilities for \( y_{0i}, y_{1i}, y_{2i}, \) and \( y_{3i} \) in migrants are denoted as \( y_{0i}, y_{1i}, y_{2i}, \) and \( y_{3i} \), respectively.

Let \( r_2 \) be the recombination fraction between the \( B \) and \( C \) loci. The changes in the conditional probabilities of \( y_{0i}, y_{1i}, y_{2i}, \) and \( y_{3i} \) due to the joint effects of migration, selection, and mutation are given in Appendix C. According to Appendix C the analytical expression for \( \Delta p_{C_i} \) is

\[
\Delta p_{C_i} = \Delta P_{A_iC_iB_i} + \Delta P_{A_iC_iB_i} + \Delta P_{A_iC_iB_i} + \Delta P_{A_iC_iB_i}
= P_{A_iB_i}\Delta y_{0i} + P_{A_iB_i}\Delta y_{1i} + P_{A_iB_i}\Delta y_{2i} + P_{A_iB_i}\Delta y_{3i}
= -\hat{m}(p_{C_i} - \hat{p}_C) - y_{0i}C_i + s_i D_{AC(i)} + s_i D_{CB(i)},
\]

(10)

where \( D_{CB(i)} \) are the LD between the \( C \) and \( B \) loci in the \( i \)th subpopulation.

According to the conditional probabilities of \( y_{0i}, y_{1i}, \) \( y_{2i}, \) and \( y_{3i}, D_{AC(i)} \) and \( D_{CB(i)} \) can be, respectively, given by

\[
D_{AC(i)} = p_{A_i} p_{a_i} (p_{B_i} (y_{0i} - y_{1i}) + p_{b_i} (y_{2i} - y_{3i}))
+ (p_{a_i} (y_{0i} - y_{2i}) + p_{A_i} (y_{1i} - y_{3i})) D_{AB(i)}),
\]

(11a)

\[
D_{CB(i)} = p_{B_i} p_{B_i} (p_{a_i} (y_{0i} - y_{1i}) + p_{b_i} (y_{1i} - y_{3i}))
+ (p_{b_i} (y_{0i} - y_{2i}) + p_{B_i} (y_{2i} - y_{3i})) D_{AB(i)}).
\]

(11b)

The first part on the right-hand side of Eqs. (11a) or (11b) is the amount without the influence of the LD between the \( A \) and \( B \) loci, and the second part is the increment due to the LD between the \( A \) and \( B \) loci generated by gene flow. The above equation analytically demonstrates that the presence of the LD among linked selected loci can enhance the effects of background selection. The proportion of \( D_{AC(i)} \) and \( D_{CB(i)} \) explained by the component of \( D_{AB(i)} \) can be assessed by \( (p_{a_i} (y_{0i} - y_{2i}) + p_{A_i} (y_{1i} - y_{3i})) D_{AB(i)} / D_{AC(i)} \) and \( (p_{b_i} (y_{0i} - y_{2i}) + p_{B_i} (y_{2i} - y_{3i})) D_{AB(i)} / D_{CB(i)} \), respectively.
Letting $\Delta y_{ij} = 0 (j = 0, 1, 2, 3)$ in Appendix C, we obtain four non-linear equations for calculating the steady-state conditional probabilities $(y_{00}, y_{11}, y_{2i}, y_{3i})$,

$$
\begin{pmatrix}
  h_{00} & \ldots & h_{05} \\
  \vdots & \ddots & \vdots \\
  h_{30} & \ldots & h_{35}
\end{pmatrix}
= \begin{pmatrix}
  y_{00} & y_{11} & y_{2i} & y_{3i} & y_{03} & y_{13} & y_{21} \\
\end{pmatrix}^T
$$

where

$$
h_{00} = s_1 p_{a_1} + s_2 p_{b_1} - u_1 - u_2 - v - \tilde{m}
- (1 + 2s_1 p_{a_1} + 2s_2 p_{b_1}) (P_{a_1 b_1} (1 - s_1)) r_1
+ P_{a_1 b_1} (1 - s_2) r_2 + P_{a_1 b_1} (1 - s_1 - s_2)
\times (r_1 + r_2 - r_1 r_2), \ldots,
$$

$$
h_{35} = \frac{1 + 2s_1 p_{a_1} + 2s_2 p_{b_1}}{P_{a_1 b_1}} (r_1 r_2 (1 - s_1 - s_2) P_{a_1 b_1} P_{a_1 b_1}),
$$

$$
g_0 = -\tilde{m} \frac{\tilde{p}_{ABC}}{P_{A_1 B_1}}, \ldots, g_3 = -\tilde{m} \frac{\tilde{p}_{abc}}{P_{a_1 b_1}}.
$$

The analytical solution is algebraically complicated, but can be numerically solved with Mathematica model solutions.

Our numerical examples demonstrate that both $D_{AB(i)}$ and $D_{AC(i)}$ increase with the migration rate (Fig. 1a). The proportion of $D_{AC(i)}$ explained by the component of $D_{AB(i)}$ can be more than 20% when $\tilde{m} = 0.1$ (Fig. 1b). Both $D_{AB(i)}$ and $D_{AC(i)}$ decrease with the recombination rate (Fig. 2a,b). The proportion of $D_{AC(i)}$ explained by the component of $D_{AB(i)}$ can be substantially increased when the three loci are tightly linked, and more than 20% of $D_{AC(i)}$ can be brought about when $r_1 = r_2 = 0.0005$ (Fig. 2c).

3.3. General case

From the preceding two- and three-locus analyses, extension can be obtained to a more general case where a neutral locus links to an arbitrary number of selected loci among which the interaction may exist. The change in the frequency of the neutral locus $C_i$ in the $i$th subpopulation, regardless of the magnitude of the linkage disequilibria among selected loci, can be generally expressed as

$$
\Delta p_{C_i} = -\tilde{m}(p_{C_i} - \tilde{p}_{C_i}) - v p_{C_i} + A_i,
$$

where $A_i = \sum_{j=1}^{n} s_j D_{M_j C(i)}$, in which $M_j$ represents the wild-type allele at the $j$th selected locus. Effects of the LD among multiple selected loci are included in $D_{M_j C(i)}$.

Fig. 1. Effects of the dependence between the selected A and B loci: (a) Changes in $D_{AC(i)}$ and $D_{AB(i)}$ with the migration rate. (b) The proportion of $D_{AC(i)}$ explained by the component of $D_{AB(i)}$. $D_{AB(i)}$ is calculated according to Eqs. (7) and (8) while $D_{AC(i)}$ is calculated according to Eq. (11a). The settings of other parameters are the mutation rate for the A and B loci $u_1 = u_2 = 10^{-5}$ and for the neutral locus $v = 10^{-4}$, the selection coefficients $s_1 = s_2 = 0.02$, the recombination rates between the A and C loci or the B and C loci $r_1 = r_2 = 0.01$ and between the A and B loci $r = 2r_1$, the LD between the A and B loci in migrants $D_{AB} = 0.03$, the migrant allele frequencies $\tilde{p}_A = \tilde{p}_B = 0.8$, and the conditional probabilities in migrants for the neutral locus $C$ under different backgrounds $\tilde{y}_0 = 0.8, \tilde{y}_1 = \tilde{y}_2 = 0.7$, and $\tilde{y}_3 = 0.2$.

The cumulative effects $A_i$ from multiple selected loci are likely substantial when they closely link to the neutral locus.

4. Population differentiation

4.1. Equal spatial selection

We now examine the effects of background selection on population differentiation using the classical island model (Wright, 1969). Background selection can change both the effective population size and the neutral allele frequency. In the preceding section we have shown the systematic change of allele frequencies at a neutral locus as a result of linkage to selected loci. Previous
studies showed that background selection can reduce the effective subpopulation size \(N_e^i\) for the neutral locus (e.g. Nordborg et al., 1996) and hence affect the genetic drift process. Here we include both effects in deriving the expression for population differentiation. According to Eq. (4) of Nordborg et al. (1996), the steady-state effective size for the \(i\)th subpopulation under the impacts of background selection, denoted by \(N_e^i\), can be approximated by

\[
N_e^i = N_e e^{-\lambda_i},
\]

where

\[
\lambda_i = \sum_{j=1}^{L} \frac{1 - p_{M,j}}{(1 + \bar{f}_j/\bar{p}_j)^2}.
\]

We assume that the effective subpopulation size is not affected by gene flow although the effective size of the whole population is changed (Wright, 1943).

Denote by \(\sigma^2\) the variance of allele frequencies among subpopulations after pollen and seed flow and background selection. Suppose that the number of subpopulations \((n)\) is large. According to Eq. (13), \(\sigma^2\) can be calculated by

\[
\sigma^2 = \frac{1}{n} \sum_{i=1}^{n} (p_{C,i} - \bar{p}_C)^2
\]

\[
\approx E_{\phi}(((1 - \bar{m})(p_{C,i} - \bar{p}_C) - v p_{C,i} + A_i)^2)
\]

\[
= ((1 - \bar{m})^2 - 2v)\sigma^2 + \Pi + \bar{A}^2,
\]

where \(E_{\phi}\) represents the expectation with respect to the allele frequency distribution among subpopulations, \(\Pi = 2(1 - \bar{m})E_{\phi}((p_{C,i} - \bar{p}_C)A_i)\), and \(\bar{A}^2 = E_{\phi}(A^2)\). Note that the expectations of the terms involving coefficients of \(v^2\), \(mv\), and \(vA\) are neglected in deriving Eq. (15).

When the selection coefficients for the same allele at any selected locus are equal among all subpopulations, i.e. \(s_1 = \cdots = s_{nj}\), then \(\lambda_1 = \cdots = \lambda_n\) and the effective subpopulation size is the same, i.e. \(N_e^1 = \cdots = N_e^n = N_e^p\). The effects of background selection are equal among subpopulations, i.e. \(A_1 = \cdots = A_n = A\), and \(\bar{A}^2 = A^2\). The second term on the right-hand side of Eq. (15) varnishes, i.e. \(\Pi = 0\). According to Hu and Ennos (1999), the steady-state variance of allele frequencies after genetic drift, can be written as

\[
\sigma^2 = \left(1 - \frac{1}{2N_e^p}\right) ((1 - \bar{m})^2 - 2v)\sigma^2 + \frac{\bar{A}_C(1 - \bar{p}_C)}{2N_e^p}.
\]

Population differentiation at the neutral locus, denoted by \(F_{st} (\equiv \sigma^2/\bar{p}_C(1 - \bar{p}_C))\), can be obtained by substituting Eq. (14) into Eq. (16),

\[
F_{st} = \frac{1}{1 + 4N_e^p(\bar{m} + v)e^{-\lambda}} \left(1 + \frac{2N_e e^{-\lambda} - 1}{\bar{p}_C(1 - \bar{p}_C)} \bar{A}^2\right).
\]

Clearly, \(F_{st} \) is greater than the population differentiation under the purely neutral process, denoted by \(F_{st,b} \) (\(= 1/(1 + 4\bar{m})\)) for diploid nuclear genes (Hu and Ennos, 1999).
4.2. Unequal spatial selection

The more general situation is that the selection coefficients of the same allele at any selected locus are unequal among subpopulations, and so is the effective subpopulation size among subpopulations, i.e. $N_{e1} \neq \cdots \neq N_{en}$. Eq. (15) remains effective since it is derived before the occurrence of genetic drift. The steady-state increment in the variance of allele frequencies after genetic drift, denoted by $\Delta \sigma^2_{D}$, is

$$\Delta \sigma^2_{D} = E\phi \left( \frac{p_c(1 - p_c)}{2N_e} \right) = E\phi \left( \frac{p_c(1 - p_c)}{2N_e} + (\lambda_i + \xi_i^2/2! + \cdots) \frac{p_c(1 - p_c)}{2N_e} \right)$$

$$= \frac{1}{2N_e} \left( \bar{\beta}_c(1 - \bar{\beta}_c) - \sigma^2 \right) + \rho,$$

where $\rho = E\phi((\lambda_i + \xi_i^2/2! + \cdots) \frac{p_c(1 - p_c)}{2N_e})$ is the increment part due to the background selection. Therefore, the steady-state equation for the variance of allele frequencies, equivalent to Eq. (16), is expressed as

$$\sigma^2 = \left( 1 - \frac{1}{2N_e} \right) \left( ((1 - \bar{\mu})^2 - 2\bar{\nu})\sigma^2 + \Pi + \Lambda^2 \right)$$

$$+ \frac{\bar{\beta}_c(1 - \bar{\beta}_c)}{2N_e} + \rho.$$ (18)

Denote by $F_{st2}$ the population differentiation at the neutral locus. Rearranging Eq. (19) yields

$$F_{st2} = \frac{1}{1 + 4\sqrt{N_e}(\bar{\mu} + \bar{\nu})(1 + \xi)}.$$

(20)

where

$$\xi = \frac{(2N_e - 1)(\Pi + \Lambda^2) + 2N_e\rho}{\bar{\beta}_c(1 - \bar{\beta}_c)}.$$

In the presence of inbreeding in each subpopulation, the variance effective population size reduces to $N_e(1 + F_{is})^{-1}$ where $F_{is}$ is the inbreeding coefficient (Caballero and Hill, 1992). Thus, the assumption of random mating can be relaxed by replacing $N_e$ in Eqs. (17) or (20) with $N_e(1 + F_{is})^{-1}$. Also the assumption of the large number of subpopulations can be relaxed by replacing $\bar{\mu}$ with $(n/(n - 1))^2\bar{\mu}$ in which $n$ can be an arbitrary number of subpopulations (Hu, 2000).

When the estimates of $\hat{F}_{st2}$ and $\hat{F}_{st,b}$ are available, the increment in $F_{st}$ due to the effect of background selection can be estimated according to the general formula of Eq. (20),

$$\hat{\xi} = \frac{\hat{F}_{st2}}{\hat{F}_{st,b}} - 1.$$

(21)

In order to look at the amount of increment in $F_{st}$ at the neutral locus due to its linkage to selected loci, the above analytical results are applied to the two- and three-locus cases under the model of equal spatial selection. The steady-state $F_{st}$ under the purely neutral process equals 0.2320 for $\bar{\mu} = 0.015$, 0.0847 for $\bar{\mu} = 0.05$, and 0.0444 for $\bar{\mu} = 0.1$, with the settings of other parameters $N_e = 50$, $n = 30$, and $v = 5 \times 10^{-4}$. In the two-locus case, the proportion of increment in $F_{st}$ is about 1.68% for $\bar{\mu} = 0.015$, 3.23% for $\bar{\mu} = 0.05$, and 4.10% for $\bar{\mu} = 0.1$ when the neutral locus tightly links to the selected locus $A$ ($r_1 = 0.0001$) (Fig. 3a). The proportion of increment in $F_{st}$ decreases with the recombination rate. In the three-locus case, the proportion of increment in $F_{st}$ is about 4.22% for $\bar{\mu} = 0.015$, 7.86% for $\bar{\mu} = 0.05$, and 9.16% for $\bar{\mu} = 0.1$ when $r_1 =
0.0001 (Fig. 3b). Although the proportion of increment in \( F_{st} \) decreases with the recombination rate, they are greater than those in the two-locus case (Fig. 3a,b). Note that the parameters for the neutral locus \( C \) and its linkage to the \( A \) (or \( B \)) locus in the above two cases are comparable: \( p_C = 0.8 \), \( \bar{p}_{AC} = 0.76 \), and \( \bar{p}_{AC} = 0.01 \) in the two-locus case; \( \bar{p}_{C} = 0.7957 \), \( \bar{p}_{AC} = 0.7769 \), and \( \bar{p}_{AC} = 0.0187 \) in the three-locus case. These numerical results indicate that a certain amount of increment in \( F_{st} \) can be attained when the neutral locus tightly links to selected loci or when the migration rate is high.

5. Simulations

5.1. Method

To confirm the analytical results simulation study was conducted according to the sequence of events in the life cycle of hermaphrodite plants. Simulation starts from an initial adult reference population that begins subdivision and produces many subpopulations. The allele frequencies at the selected loci are initially set to be the same as in the reference population. The conditional probabilities for a diallelic neutral gene in the reference population under different backgrounds of selected gametes are also assumed, thus the allele frequencies of the neutral locus can be calculated. The frequencies of two- or three-locus gametes in migrants (seeds and pollen grains) are assumed to be equal to those in the initial reference population. Constant selection coefficients among subpopulations are examined although a more complicated pattern of selection can be modeled.

The detailed simulation procedure is as follows. Given the initial parameter settings, calculate the frequencies of two- or three-locus gametes in pollen and ovules according to the assumption of Wright–Fisher’s model. Then calculate the gamete frequencies after pollen flow. According to the assumption of random combination between pollen and ovules, calculate the genotype frequencies in seeds so formed. Seed flow is then considered and the genotype frequencies are calculated after seed flow. Assume that the mutation at the neutral locus is a deterministic process, with a probability of \( v \) from the allele \( C \) to the allele \( c \), per generation, and then calculate all genotype frequencies after mutation. The model of multiplicative viability is employed to calculate the fitness of each genotype in any subpopulation and the genotype frequencies after selection. A sampling process (genetic drift) is then conducted according to the phenotype frequencies after selection, given an effective subpopulation size (\( N'_{el} \)). Gamete frequencies in ovules and pollen grains are then calculated according to the segregation ratios of gametes from individual cross (10 crosses in the two-locus case and 36 crosses in the three-locus case). The above steps are repeated until the population differentiation reaches steady distribution from generation to generation.

Five thousand independent data sets are created per generation, and each is used to calculate population differentiation. From these replicated datasets, means and standard deviation of \( F_{st} \) are calculated. The predicted results are obtained according to Eq. (17), where the LD is calculated from Eq. (6) in the two-locus case and Eq. (11a, b) in the three-locus case. The steady-state values of \( y_j \) (\( j = 0,1,2,3 \)) in calculating the expected \( F_{st} \) values are calculated according to Eq. (12) with the Mathematica tool.

5.2. Results

Let \( n = 30 \), \( v = 5 \times 10^{-4} \), and \( N_e = 50 \). In the purely neutral process, the theoretical expectation of \( F_{st} \) at steady state equals 0.2320 when \( \bar{m} = 0.015, 0.0847 \) when \( \bar{m} = 0.05 \), and 0.0444 when \( \bar{m} = 0.1 \). In the presence of background selection, our simulations clearly show that population differentiation is enhanced, especially when the neutral locus is closely linked to the selected loci or when migration rate is high. In the two-locus case, for example, the average proportion of increment in \( F_{st} \) at the 200th generation (steady-state value) reduces from 9.48% for \( r_1 = 0.01–2.97% \) for \( r_1 = 0.1 \) (Fig. 4a,b), although these estimates are greater than the expected values of 1.12% and 0.25% (Fig. 3a), respectively. The expected values of \( F_{st} \) are within the range of one standard deviation of empirical results. The general pattern for the change of \( F_{st} \) with generation is that the average \( F_{st} \) is initially small when the reference population starts subdivision and all subpopulations are formed by sampling from the same reference population. Then population differentiation gradually increases and approaches a steady distribution after 160 generations (Fig. 4a). Compared with the results in the case of loose linkage (\( r_1 = 0.1 \)), population differentiation displays a greater fluctuation in the case of tight linkage (\( r_1 = 0.01; \) Fig. 4b).

Population differentiation quickly reaches a steady distribution with the increase of migration rate (Fig. 5a,b). The average proportion of increment in \( F_{st} \) generally increases with migration rate. For example, the average proportion of increment in \( F_{st} \) at the 200th generation increases from 9.48% for \( \bar{m} = 0.015 \), to 10.5% for \( \bar{m} = 0.05 \), and to 16.9% for \( \bar{m} = 0.10 \) (Fig. 5a), although these estimates are greater than the expected values of 1.12%, 2.28%, and 2.74% (Fig. 3a), respectively. All expected \( F_{st} \) values are within the range of one standard deviation of empirical results.

The change of \( F_{st} \) with generation in the three-locus case has the pattern similar to that in the two-locus case, which displays a gradual increase with time and eventually approaches a stable distribution (Fig. 6a), but has a greater fluctuation than the latter (Fig. 6b).
Compared with the results in the two-locus case, the average $F_{st}$ in the three-locus case is increased. For example, the average proportion of increment in $F_{st}$ at the 250th generation (steady-state value) is about 13.8% for $r_1 = 0.01$ and 11.6% for $r_1 = 0.1$ (Fig. 6a), although these estimates are greater than the expected values of 3.92% and 2.97% (Fig. 3b), respectively. The general pattern for the effect of migration on increasing the proportion of increment in $F_{st}$ can be observed (Fig. 7a). For example, the average proportions of increment in $F_{st}$ at the 250th generation are 13.8% for $\tilde{m} = 0.015$, 15.7% for $\tilde{m} = 0.05$, and 26.4% for $\tilde{m} = 0.10$ (Fig. 7a,b), although these estimates are greater than the expected values of 3.92%, 7.53%, and 8.88%, respectively (Fig. 3a). Again, all predicted $F_{st}$ values are in the range of one standard deviation of the empirical results. In summary, these simulation results demonstrate that the cumulative effect of background selection on $F_{st}$ at a neutral locus can be substantial if the neutral locus is closely linked to multiple selected loci.

6. Discussion

In this paper we have obtained the analytical expressions for population differentiation at a neutral locus in the island model of population structure. The increase in population differentiation as a result of background selection is analytically demonstrated under very general conditions. Although the individual effects of a single selected locus are likely to be small, the cumulative effect of multiple selected loci could be substantial. Our theoretical results can be applied to a
wide situation to map the effects of background selection in terms of population differentiation at a fine genome scale.

Although the present result is qualitatively the same as a previous study (Charlesworth et al., 1997), our approach is fundamentally different in theoretical deduction. By partitioning the total genetic diversity into the components of between and within subpopulations, Charlesworth et al. (1997) showed that the increase of $F_{st}$ due to background selection is mainly caused by the decreased diversity within populations. Our general expression of $F_{st}$ is derived using Wright’s approach and seems more rigorous.

The expression of $F_{st}$ given by Charlesworth et al. (1997) is only suitable for a pair of subpopulations. Slatkin and Wiehe (1998) have investigated the hitchhiking effects on population differentiation using the island and stepping-stone models of population structure, and presented the analytical expression of $F_{st}$ suitable for the case of two subpopulations. The present model is an extension of the classical island model (Wright, 1969) to incorporate effects of background selection and necessarily expands the study of Charlesworth et al. (1997) to an arbitrary number of subpopulations simultaneously. Further, when the haploid dispersal of pollen is set to zero, our $F_{st}$ formula can be applied to animal populations where only diploid dispersal occurs.

Until now empirical studies have been concentrated on the observations of high $F_{st}$ in the regions of low recombination as a result of hitchhiking effect, such as in Drosophila species (e.g., Stephan and Mitchell, 1992; Begun and Aquadro, 1993). Much less attention has been paid to the effects of background selection on $F_{st}$. A recent study shows that the average level of nucleotide
diversity in regions of low recombination can be used to
distinguish background selection from hitchhiking
effects (Innan and Stephan, 2003). Because population
differentiation of a neutral locus is negatively related to
the genetic diversity within subpopulations given a total
 genetic variation, an interesting question is whether the
measure of $F_{st}$ can be used to distinguish background
selection from hitchhiking effects. The properties of $F_{st}$
under hitchhiking effects (Slatkin and Wiehe, 1998)
and background selection (present study) are very similar,
displaying a negative correlation with recombination
fraction. The “spatial” pattern of $F_{st}$ along chromo-
somes is likely very similar between these two types of
processes, and this problem presents a challenge for
future study.

The present theory provides an implemental techni-
que to examine the magnitude of background selection
effects. The conventional approaches for estimating
population differentiation at neutral loci remain valid
(e.g., Weir, 1996). The problem is how to distinguish
which loci are purely neutral and not affected by
background selection or hitchhiking effects. With
molecular genome sequence data, $F_{st}$ can be calculated
for all individual SNP that are distributed on the same
chromosome, using the method introduced by Hudson
et al. (1992), and hence the pattern of genome-wide $F_{st}$
can be mapped. Those neutral loci with the smallest $F_{st}$
values can be used for approximating $F_{st,b}$, and the
effects of background selection at other individual SNP
can be estimated according to Eq. (21). The prerequi-
sites for such genomic-wide $F_{st}$ mapping are the tests of
neutrality and background selection.

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Appendix A. Changes in the conditional probability in
the two-locus case

Let $P_{ACi}^*$ and $P_{ACi}$ be the frequencies of gametes $ACi$
and $ACj$ after selection in the $i$th population, respect-
ively. The mean fitness in the $ith$ subpopulation, denoted
by $\bar{w}_i$, is approximated by $\bar{w}_i = 1 - 2p_{ACi}s_{ii}$. The
frequency of each two-locus gamete after selection
($P_{ACj}$ and $P_{AC}$) can be calculated using the conven-
tional method,

$$P_{ACj} = \frac{(1 - p_{ACi}s_{i1})P_{ACi} + \delta_0 r_1)/\bar{w}_i}{(1 - p_{ACi}s_{i1})P_{ACi} + \delta_0 r_1)/\bar{w}_i + (u_1 - v)P_{ACi} - \tilde{m}(P_{ACi} - \tilde{P}_{AC})}, \quad (A.1)$$

where $\delta_0 = P_{ACi}P_{ACi}(x_{i1} - x_{i0})(1 - s_{i1})$. The changes in
gamete frequency ($\Delta P_{ACj} = P_{ACj} - P_{ACi}$) and
$\Delta P_{AC} = P_{AC} - P_{ACi}$ can be calculated from Eqs.
(A.1) and (A.2).

From the relation of $P_{ACi} = p_{ACi}x_{0i}$, we obtained
$\Delta P_{ACj} = \Delta p_{ACi}x_{0i} + p_{ACi}\Delta x_{0i}$. Since $\Delta p_{ACi} = 0$
at steady state, the change in the conditional probability of $x_{0i}$
by

$$\Delta x_{0i} = \frac{\Delta P_{ACi}/P_{ACi} = r_1p_{ACi}(1 - s_{i1})(1 + 2p_{ACi}s_{i1})(x_{i1} - x_{0i})}{(1 + 2p_{ACi}s_{i1})(x_{i1} - x_{0i})} + (p_{ACi}s_{i1} - u_1 - v - \tilde{m})x_{0i} + \tilde{m}\tilde{P}_{ACj}/P_{ACi}, \quad (A.3)$$

where $\tilde{P}_{ACj}$ is the frequency of the gamete $ACj$
in migrants. Similarly, the change in the conditional probability
$x_{1i}$ is given by

$$\Delta x_{1i} = \frac{\Delta P_{ACi}/P_{ACi} = r_1p_{ACi}(1 - s_{i1})(1 + 2p_{ACi}s_{i1})(x_{0i} - x_{1i})}{(1 + 2p_{ACi}s_{i1})(x_{0i} - x_{1i})} + (p_{ACi}s_{i1} - s_{i1} - v - \tilde{m})x_{1i} + u_1P_{ACi}x_{0i}/P_{ACi} + \tilde{m}\tilde{P}_{ACj}/P_{ACi}, \quad (A.4)$$

where $p_{ACj}$ is the frequency of the gamete $ACj$
in migrants. Since the genetic drift process does not
change the average gamete frequency, Eqs. (A.3) and
(A.4) actually represent the per-generation changes in
the conditional probabilities of $x_{0i}$ and $x_{1i}$, respect-
ively.

Appendix B. Recurrent equation for the LD between two
selected loci

Let $D_{AB(i)}$ be the LD between the $A$ and $B$ loci in the
$ith$ subpopulation in the current generation (adults), and
$r$ be the recombination rate between them. The
gamete frequencies at the current generation, denoted
by $P_{ji}$ ($j = A, a; l = B, b$), can be expressed as $P_{ji} = P_{ji} +
\delta_{jl}D_{AB(i)}$, where $\delta_{AB} = \delta_{ab} = 1$ and $\delta_{Ab} = \delta_{aB} = -1$.
The gamete frequencies in pollen and ovules in the next
generation, denoted by $P_{ji}'$, can be expressed by $P_{ji}' = P_{ji} +
(1 - r)\delta_{jl}D_{AB(i)}$. After pollen flow the gamete
frequencies in pollen, denoted by $P_{ji}'$, can be expressed by $P_{ji}' = mp\tilde{P}_{ji}' + (1 - mp)P_{ji}'$, where $\tilde{P}_{ji}'$ is the gamete
frequency in migrants. The gamete frequencies in ovules
remain the same as those in the preceding adults.

After random combination between pollen and
ovules, the nine genotypic frequencies in seeds so
formed can be readily calculated. For example, the
frequency for the genotype $AABB$, $B_{AB}$ in seeds, denoted
by $P_{AABB}$, equals $P_{AABB}$. The frequency for the
genotype $AABBB$, $B_{AAB}$ after seed flow can be expressed as

$$P_{AABBB} = mS\tilde{P}_{AABB} + (1 - mS)P_{AABBB}, \quad (A.4)$$

where $\tilde{P}_{AABB}$ is the frequency in migrating seeds. The frequencies for
other genotypes after seed flow can be expressed

$P_{aACj} = \frac{(1 - p_{ACi}s_{i1})P_{ACi} + \delta_0 r_1)/\bar{w}_i}{(1 - p_{ACi}s_{i1})P_{ACi} + \delta_0 r_1)/\bar{w}_i + (u_1 - v)P_{ACi} - \tilde{m}(P_{ACi} - \tilde{P}_{AC})}, \quad (A.1)$
in a similar way. Following the procedure of mutation and selection, the genotypic frequencies in adults in the next generation can be derived using the conventional method. The frequencies of four gametes after selection in adults, denoted by $P^s_{j}(j = A_i,a_i; l = B_i, b_i)$, are

$$P^s_{A_iB_i} = \hat{m}\hat{p}_{AB} + (1 - \hat{m} + p_{a_i}s_{1i}) + u_2 P'_{A_iB_i}, \quad (B.1a)$$

$$P^s_{A_i b_i} = \hat{m}\hat{p}_{A} + (1 - \hat{m} + p_{a_i}s_{1i} + p_{b_i}s_{2i} - u_1)P'_{A_i b_i} + u_2 P'_{A_i b_i}, \quad (B.1b)$$

$$P^s_{a_iB_i} = \hat{m}\hat{p}_B + (1 - \hat{m} - p_{a_i}s_{1i} + p_{b_i}s_{2i} - u_2)P'_{a_iB_i} + u_1 P'_{a_iB_i}, \quad (B.1c)$$

$$P^s_{a_i b_i} = \hat{m}\hat{p}_B + (1 - \hat{m} - p_{a_i}s_{1i} - p_{b_i}s_{2i} + u_2)P'_{a_i b_i} + u_1 P'_{a_i b_i}, \quad (B.1d)$$

Let $D^s_{AB(i)}$ be the LD in the $i$th subpopulation in the next adult generation. According to Eqs. (B.1a)–(B.1d), the recurrent equation for LD between the $A$ and $B$ loci is derived as

$$D^s_{AB(i)} = P^s_{A_iB_i}P^s_{a_iB_i} - P^s_{A_i b_i}P^s_{a_i b_i}$$

$$= \hat{m}(D_{AB} + (p_{a_i} - \hat{p}_A)(p_{b_i} - \hat{p}_B))$$

$$+ (1 - \hat{m} - (p_{a_i} - \hat{p}_A)s_{1i} - (p_{b_i} - \hat{p}_B)s_{2i} - u_1 - u_2)(1 - r)D_{AB(i)}, \quad (B.2)$$

where $D_{AB}$ is the LD in migrants.

## Appendix C. Changes in the conditional probability in the three-locus case

Let $P^s_{A_iC_iB_i}$, $P^s_{A_iC_iB_i}$, $P^s_{a_iC_iB_i}$, and $P^s_{a_iC_iB_i}$ be the frequencies of gametes $A_iC_iB_i$, $A_iC_iB_i$, $a_iC_iB_i$, and $a_iC_iB_i$ after selection in the $i$th population, respectively. Using the assumption of multiplicative viability model, the mean fitness in the $i$th subpopulation equals $\bar{W}_i = 1 - 2p_{a_i}s_{1i} - 2p_{b_i}s_{2i}$. Using the same approach as in the two-locus case, the frequencies of the four three-locus gametes are given by

$$P^s_{A_iC_iB_i} = \left( (1 - s_{1i}p_{a_i} - s_{2i}p_{b_i})P_{A_iC_iB_i} - \delta_1 r_1 \right) \left( 1 + vP_{A_iC_iB_i} - P_{ACB} \right)$$

$$+ \frac{\delta_2 r_2 + \delta_3}{\bar{W}_i} + u_1 P_{A_iC_iB_i} - u_2 P_{a_iC_iB_i} - \hat{m}(P_{A_iC_iB_i} - P_{ACB}), \quad (C.1a)$$

$$P^s_{a_iC_iB_i} = \left( (1 - s_{1i}(1 + p_{a_i}) - s_{2i}p_{b_i})P_{a_iC_iB_i} \right)$$

$$+ \frac{\delta_1 r_1 + \delta_2 r_2 + \delta_3}{\bar{W}_i} + u_1 P_{a_iC_iB_i} - u_2 P_{a_iC_iB_i} - vP_{a_iC_iB_i} - \hat{m}(P_{a_iC_iB_i} - P_{ACB}), \quad (C.1b)$$

$$P^s_{A_iC_iB_i} = \left( (1 - s_{1i}(1 + p_{a_i}) - s_{2i}p_{b_i})P_{A_iC_iB_i} \right)$$

$$+ \frac{\delta_1 r_1 + \delta_2 r_2 + \delta_3}{\bar{W}_i} + u_1 P_{A_iC_iB_i} + u_2 P_{A_iC_iB_i} - u_2 P_{a_iC_iB_i} - \hat{m}(P_{A_iC_iB_i} - P_{ACB}) - \hat{m}(P_{a_iC_iB_i} - P_{ACB}) - P_{ACB}, \quad (C.1c)$$

$$P^s_{a_iC_iB_i} = \left( (1 - s_{1i}(1 + p_{a_i}) - s_{2i}(1 + p_{b_i}))P_{a_iC_iB_i} \right)$$

$$+ \frac{\delta_1 r_1 + \delta_2 r_2 + \delta_3}{\bar{W}_i} + u_1 P_{a_iC_iB_i} + u_2 P_{a_iC_iB_i} - P_{ACB}, \quad (C.1d)$$

where

$$\delta_1 = P_{A_iB_i}P_{a_iB_i}(1 - s_{1i} - s_{2i})(y_{0i} - y_{1i})$$

$$+ (P_{A_iB_i}P_{a_iB_i} - y_{1i})y_{0i}(1 - s_{1i} - s_{2i}),$$

$$\delta_2 = P_{A_iB_i}P_{a_iB_i}(1 - s_{1i} - s_{2i})(y_{0i} - y_{1i})$$

$$+ (P_{A_iB_i}P_{a_iB_i} - y_{1i})y_{0i}(1 - s_{1i} - s_{2i}),$$

$$\delta_3 = P_{a_iB_i}(1 - s_{1i} - s_{2i})(y_{3i} - y_{1i})$$

$$+ (P_{A_iB_i}P_{a_iB_i} - y_{1i})y_{0i}(1 - s_{1i} - s_{2i}),$$

$$\delta_4 = P_{A_iB_i}(1 - s_{1i} - s_{2i})(y_{3i} - y_{1i})$$

$$+ (P_{A_iB_i}P_{a_iB_i} - y_{1i})y_{0i}(1 - s_{1i} - s_{2i}),$$

$$\delta_0 = (1 - s_{1i} - s_{2i})(P_{A_iB_i}P_{a_iB_i}(y_{0i} + y_{3i} - y_{0i}y_{3i})$$

$$- P_{A_iB_i}y_{0i}(y_{1i} + y_{2i} - y_{1i}y_{2i})).$$

$\delta_0$ is associated with the effect of double crossover among the three loci.

From the expression of $P_{A_iC_iB_i} = P_{A_iB_i}y_{0i}$, we obtain

$$\Delta P_{A_iC_iB_i} = \Delta P_{A_iB_i}y_{0i} + P_{A_iB_i}\Delta y_{0i}. \quad (C.2)$$

Since $\Delta P_{A_iB_i} = 0$ at steady state, the change in the conditional probability of $y_{0i}$ is

$$\Delta y_{0i} = \Delta P_{A_iC_iB_i}/P_{A_iB_i}$$

$$= (s_{1i}p_{a_i} + s_{2i}p_{b_i} - u_1 - u_2 - v - \hat{m})y_{0i}$$

$$+ (\delta_1 r_1 - \delta_2 r_2 + \delta_3)(1 + 2s_{1i}p_{a_i} + 2s_{2i}p_{b_i})/P_{A_iB_i}$$

$$+ \hat{m}P_{ACB}/P_{A_iB_i}, \quad (C.3a)$$

where $\Delta P_{A_iC_iB_i} = P_{A_iC_iB_i} - P_{A_iC_iB_i}$ and $P_{ACB}$ is the frequency of gamete $A_iC_iB_i$ in migrants.

Similarly, the changes in the conditional probabilities of $y_{1i}$, $y_{2i}$, and $y_{3i}$ are derived as

$$\Delta y_{1i} = (s_{1i}p_{a_i} + s_{2i}p_{b_i} + u_1 - u_2 - v - \hat{m})y_{1i}$$

$$+ (\delta_1 r_1 + \delta_2 r_2 - \delta_3)(1 + 2s_{1i}p_{a_i} + 2s_{2i}p_{b_i})/P_{a_iB_i}$$

$$+ \hat{m}P_{ACB}/P_{a_iB_i}, \quad (C.3b)$$

$$\Delta y_{2i} = (s_{1i}p_{a_i} - s_{2i}p_{b_i} - u_1 + u_2 - v - \hat{m})y_{2i}$$

$$+ (\delta_1 r_1 + \delta_2 r_2 - \delta_3)(1 + 2s_{1i}p_{a_i} + 2s_{2i}p_{b_i})/P_{a_iB_i}$$

$$+ \hat{m}P_{ACB}/P_{a_iB_i}, \quad (C.3c)$$
\[ \Delta y_{3i} = (-s_1 p_{A_i} - s_2 p_{B_i} + u_1 + u_2 - \bar{v} - \bar{m}) y_{3i} + (-\delta_1 r_1 - \delta_2 r_2 + \delta_0)(1 + 2s_1 p_{A_i} + 2s_2 p_{B_i}) / p_A p_B + \bar{m}\bar{p}_{A_i C_i B_i} / p_{A_i B_i}, \]

where \( \bar{p}_{A_i C_i B_i} \), \( \bar{p}_{A_i C_i B_i} \), and \( \bar{p}_{A_i C_i B_i} \) are the frequencies of gametes \( A_i C_i B_i \), \( a_i C_i B_i \), and \( a_i C_i B_i \) in migrants, respectively.

References


