Going against the flow: Maintenance of alongshore variation in allele frequency in a coastal ocean.

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ABSTRACT

Understanding the processes that develop and maintain diversity in coastal communities is an important challenge, particularly given the conservation and management needs of these ecosystems. Such diversity patterns include not only species distributions and interactions, but also variation in genetic diversity. Alongshore variations in allele frequency along coastal oceans are controlled by interactions between physical and biological forces. Here we model these interactions in terms of Lagrangian descriptors of nearshore physical oceanography, the life history dynamics of an individual species, and habitat quality. This model allows us to define population boundaries within the species range as a function of the ocean circulation, spatial habitat variability, and larval characteristics. From this, we can find quantitative criteria for the persistence of regions of alongshore genetic variation. Our results show quantitatively that the existence of phylogeographic patterns in species with high dispersal capacity along uniform coasts with typical mean currents is surprising, and requires either strong selection gradients, alongshore variation in the ocean currents and/or habitat quality, or a mixture of both. The importance of this work for marine reserve design and understanding community-wide evolutionary patterns are discussed.
A classic problem in the study of marine biodiversity is how patterns of genetic differentiation arise in an environment with few apparent barriers to dispersal, in species with high capacity for dispersal due to larval/juvenile life history, and in regions where coastal currents should promote population mixing (Palumbi 1994). Most phylogeographic studies of coastal species emphasize the historical changes that could have initiated allopatry and the process of genetic differentiation. Little attention is generally given to the processes that then must maintain spatial genetic discontinuities in these populations, except in situations where physiological or other barriers are apparent (e.g. the Florida peninsula preventing gene flow between the Gulf and Atlantic lineages; Avise 1994, Lee and O’Foighil 2004). We will quantitatively examine what processes allow spatial genetic discontinuities to persist in a coastal ocean for species with a planktonic dispersal stage. Some – possibly most – of these patterns (including gradual clines and phylogeographic ‘breaks’; see Avise 2000) may reflect the disparity between migration and gene flow caused by selection (Palumbi 1994, Barton and Hewitt 1989, Sotka et al. 2003, Sanford et al. 2003, Hare et al. 2005).

Many phylogeography studies involve an inherent assumption that the gene flow process being described is selectively neutral – that is, that the pattern is evidence of dispersal limitation alone, rather than incorporating factors that limit gene flow such as selection or ecological interactions (as well as stochastic factors associated with historical vicariance and the genealogical data themselves; Haydon et al. 1994, Irwin 2002). Here we present a model that describes the interaction between selection (defined by the relative reproductive success of individuals) and physical oceanography. We find that it is unlikely that a cline can persist in species with high dispersive capacity in an advective environment for an ecologically significant time without selection favoring the downstream allele, where downstream is defined with respect to the mean direction of larval transport.

We find that there are two interrelated mechanisms by which a cline or other phylogeographic pattern (hereafter, any population genetic discontinuity among different
geographic regions will simply be referred to as a cline) can persist in the coastal ocean in the face of non-random dispersal of larvae. First, a spatial gradient in selection can be sufficient to allow a cline to persist (Endler 1977, Hare et al. 2005). We extend prior results by quantifying the strength of the selection gradient needed to allow a cline to persist as a function of larval characteristics and local oceanographic conditions. Secondly, alongshore variability in habitat quality or larval transport can create a region that is recruit-limited, leading to increased local reproductive success and population retention. By explicitly including spatial variability in population dynamics in our model, we find that these regions can allow a cline to persist at lower selection levels than in the absence of alongshore variation.

Our model allows earlier results of Nagylaki (1978) and others who studied systems with biased directional dispersal to be applied to a coastal ocean without the \textit{a-priori} demarcation of the spatial extent of populations. Through analysis of the effect of population dynamics on the retention of alleles one can discover how to define populations (demes) in such a system, identify “source” and “sink” populations, and show how these populations are related to the oceanic circulation that drives dispersal and alongshore variation in habitat quality. Without such a quantitative description of how abiotic forces interact to maintain geographic variation in genetic markers, we are left with only a static view of how allelic diversity is regulated in natural communities. This model shows how the observed spatial patterns of genetic differentiation arise from the interaction of the circulation of the ocean, larval dispersal, and spatial variation in habitat quality.

\textbf{A Model of a Benthic Species with Planktonic Dispersal.}

The theoretical results derived below are obtained for an idealized model of a coastal benthic organism with a planktonic larval stage. The retention of a population of this idealized species in the presence of mean and variable alongshore flows has been described in Byers and
Pringle (2006; hereafter BP). In this section we describe this organism, and a simple numerical model that describes its dispersal and population dynamics. Each adult produces $N_{\text{larvae}}$ offspring per lifetime that can recruit and grow to reproductively competent adulthood, if there is sufficient empty habitat for the larvae to settle in (it is important to note that $N_{\text{larvae}}$ does not include density dependent mortality effects caused by habitat scarcity at settlement, but incorporates mortality in the larval pool, and mortality after settlement but before reproductive competency). The organism is assumed to live in a coastal habitat narrow enough that its larvae can spread across the cross-shelf dimension of the habitat in a single generation, so that the habitat can be treated as one-dimensional. The habitat has a carrying capacity of $H_{\text{dens}}$ adults per meter. These and all other parameters are defined in Table 1.

Each larva is dispersed a variable distance downstream from its parent. The mean dispersal distance is $L_{\text{adv}}$, and the standard deviation of the dispersal distance is $L_{\text{diff}}$ (see Figure 1). For simple circulation and behavioral regimes, $L_{\text{adv}}$ is the distance the larvae are advected by the mean currents and $L_{\text{diff}}$ is the distance eddies and stochastic current fluctuations would diffuse the larvae. BP quantifies these transport parameters as a function of larval behavior, inter- and intra-seasonal current variability, and the interaction of current variability and multiple spawning events. BP describes how these parameters may be estimated in regions of complex, spatial varying coastal circulation. In order to draw order-of-magnitude conclusions for typical organisms, we present from BP a simple estimate of $L_{\text{adv}}$ and $L_{\text{diff}}$ for an organism with simple larval behavior in an ocean whose circulation statistics vary little over the dispersal distance of a larva. Following Siegel et al. (2003), it can be shown that larvae that remain in the plankton for a time $T_m$ will move on average a distance

$$L_{\text{adv}} = U T_m,$$

(1)

where $U$ is the alongshore flow averaged over the time the larvae are in the plankton. Not all plankton will move the same distance, however, due to the random fluctuations in the currents associated with eddies, fluctuating wind driven currents, and other sources of temporal variability
in the ocean currents. These fluctuations lead to spread in larval settlement locations, and the standard deviation of the larval settlement position will be, from Siegel et al. (2003),
\[ L_{\text{diff}} = (\sigma^2 \tau_L T_m)^{0.5}, \] (2)
where \( \sigma \) is the standard deviation of the alongshore currents, and \( \tau_L \) is the Lagrangian timescale of the fluctuations of the mean alongshore currents, all defined at the depth or depths that the larvae reside. \( U, \sigma, \) and \( \tau_L \) are well-characterized properties of the coastal ocean, given an understanding of the vertical distribution of the larvae. Siegel et al. find that these simple estimates perform well in Central California. A comparison of dispersal distance of many species by Shanks et al. (2003) has found that (1) overestimates dispersal distance for many species. However, the linear relation between time in plankton and dispersal distance found in (1) remains valid in this comparison. All of the parameters described above are a function of the location of the parents. This perspective is analogous to a Lagrangian reference frame in fluid dynamics (see Siegel et al. 2003), and must be kept firmly in mind when interpreting the results presented below.

Some of the individuals in the population carry allele(s) that give the individual a fitness advantage over other individuals. This is quantified with a selection coefficient \( s \) against all other alleles in the population, as in Hartl and Clark (2000). The models below will be for alleles in haploid organisms, or alleles at mitochondrial loci. They will also apply to alleles in the nuclear (diploid) genome as well, if the selective advantage is additive, but the derivations below would have to be altered in the case of overdominance or higher-order gene interactions. In the comparison to observed clines presented below, it must be remembered that the required selection is not necessarily imposed on the genetic markers that actually identify the cline; the individual, not the gene, could be in this case the unit of selection (for a discussion of this, and its implications for the retention of other novel diversity, see Barton and Hewitt 1989).

In order to illustrate and confirm some of the results below, it is useful to have an individual-based numerical model of this idealized organism. The computer model is a
straightforward implementation of the organism described above. The habitat in the computer is a
one-dimensional array of settlement sites each of which can hold one adult, and each adult
produces $N_{\text{larvae}}$ propagules that move away from the parent as described above. The larval
dispersal kernel used is Gaussian. However, the results below are not sensitive to this under
typical circumstances, as described in BP. If the larvae land on a site inhabited by an adult, they
die. If several larvae arrive at the same site at the same time, one is randomly chosen as a
survivor, with a weighting appropriate for selection, which results in a logistic density
dependence. Thus in the numerical model selection enters through competition, a form of soft
selection (sensu Wallace 1975), though the theory discussed below is developed for both soft and
hard selection mechanisms. The code for the model is in MATLAB and FORTRAN, and is
available from the authors.

**The Ephemeral Nature of a Cline between Neutral Alleles in a Uniform**
**Advective Environment.**

To understand how a gradient in allele frequency can be maintained against a mean
current in an organism such as described above, we will first examine the behavior of such a
gradient between two neutral alleles in a spatially uniform coastal ocean with a spatially uniform
habitat. To do so, a population of the idealized organism described above is separated into two
groups that differ only by a selectively neutral “marker” allele, with one allele (or allele class)
found in one portion of the habitat, and the other allele predominant in the remaining
“downstream” portion of habitat (Figure 2). The population density of the entire domain is at its
carrying capacity. The progeny of an offspring inherit the allele from their parent.

Individuals with either allele have the same dispersal characteristics, with their larvae
moving on average $L_{\text{adv}}$ downstream with a standard deviation of $L_{\text{diff}}$. In figure 2, the temporal
evolution of this cline is shown. While both allelic populations move downstream, the upstream
allele (maintained under the same conditions that maintain the population along the coast, see BP) rapidly displaces the downstream allele, and the downstream allele is lost from the population.

The cline moves downstream because, in this region with a saturated habitat, each adult leaves on average only one offspring to replace an adult that has just died. This propagule is on average $L_{adv}$ downstream from the adult, and so the cline moves on average a distance $L_{adv}$ downstream over a single lifetime. For organisms with relatively long planktonic larval stages ($T_m$ on the order of many days or weeks), mean dispersal distances are often tens to hundreds of kilometers per generation (Strathmann 1985, Kinlan and Gaines 2003, Shanks et al. 2003). For such organisms, a cline between two neutral alleles in a spatially uniform domain would sweep through a typical species’ geographic range in a few tens to hundreds of generations. For most species, this is a short time compared to either the time it takes to generate a cline through isolation by distance and random genetic drift, or the time since Pleistocene glaciation – a commonly proposed cause of historical allopatry.

However, the downstream alleles in figure 2 could persist in the domain if the adults carrying these alleles produced, on average, enough surviving offspring. If this occurred, the stochastic component of larval dispersal could return sufficient offspring to the location of their parents to maintain the downstream allele in the domain. In the next sections, we discuss how the individuals with the downstream allele might be able to produce more than one surviving offspring, and define explicit criteria for the persistence of the downstream alleles in the domain.

**When Can an Allele be Retained in a Region?**

To understand the persistence of a cline in gene frequency along a coastal ocean, it is first necessary to understand when an allele introduced into the center of a uniform coastal habitat with a mean alongshore current will persist. This ocean has alongshore uniform hydrodynamics and habitat, and thus $L_{adv}$, $L_{diff}$, $H_{dens}$ and $N_{larvae}$ remain constant along the coast. Individuals
carrying the allele are introduced into the habitat at a density much less than the maximum population density that could persist in the habitat, but in large enough numbers that the allele is unlikely to be lost through stochastic fluctuations in gene frequency.

To understand the persistence of the introduced allele, we use a result from BP for the persistence of a population. BP find that the descendants of an individual of a species introduced into an empty habitat – regardless of allelic composition – will persist and be retained in that area, and will grow to populate that area, if

\[ \ln(N_{\text{larvae}}) > \frac{L_{\text{adv}}^2}{2L_{\text{diff}}^2} \]  

Eq. (3) is valid as long as the domain into which the species is introduced is much larger than \( L_{\text{diff}} \) and \( L_{\text{adv}} \), and if the dispersal kernel is neither leptokurtic (which would make it easier for the introduced species to persist), nor of a finite extent not significantly greater than \( L_{\text{adv}} \) (BP).

When (3) and these criteria are met, enough larvae are returned to the location of the parents by the stochastic component of larval dispersal (\( L_{\text{diff}} \)) to replace adult mortality, despite the downstream transport of most larvae by the mean currents (\( L_{\text{adv}} \)).

The dynamics of the retention of a small number of alleles introduced into a population in which they are initially scarce are identical but for one change, for the underlying question is similar: when can the descendents of an allele introduced into a region persist in that region, despite the tendency of the mean currents to transport most copies of the allele downstream? The only change is that the relevant growth parameter is not the number of offspring an adult would leave behind in a region in which the species is scarce (\( N_{\text{larvae}} \)), but is instead \( N_{\text{allele}} \), the number of copies of the allele carried by surviving – and successfully recruiting – offspring for each copy present a lifetime earlier when the allele is scarce. Thus, the criterion for the retention of the allele introduced into a population is

\[ \ln(N_{\text{allele}}) > \frac{L_{\text{adv}}^2}{2L_{\text{diff}}^2} \]  

(this criterion will be explicitly tested in a numerical model below). When this criterion is met, enough copies of an allele successfully recruit to the location of the introduction by the stochastic
component of larval transport that the population of the allele is maintained there, and indeed the
distribution of the allele would tend to extend upstream (BP), despite the transport of the average
larvae downstream. This can be seen in the bottom panel of figure 3, in which the introduced
allele grows in frequency at and upstream of the location of introduction. If the criterion is not
met, the total number of copies of the allele in the population would still tend to increase, but this
increase would be insufficient to maintain the allele frequency at the point of introduction. Thus
the density of the allele at the point of introduction would decay to zero, the mean location of the
introduced alleles would drift downstream, and eventually all copies of the allele would be lost
off the downstream edge of the domain. This is seen in the top panel of figure 3 and in figure 2.

The criterion above explains only how an allele introduced into the middle of a coastal
ocean could persist. It neither explains how the growth in the number of copies of allele, \( N_{\text{allele}} \),
could come about, nor does it explain how, if (4) is met, the downstream allele in a cline does not
spread upstream and homogenize gene frequency in the upstream direction. In the following
section, we focus on the dynamics of an allele in a population that is at the carrying capacity of
the environment everywhere. In this case, selection alone can increase \( N_{\text{allele}} \) and a spatial
gradient in selection is needed to allow a cline to persist but not spread upstream. In the
subsequent section, we examine the population dynamics of a region of alongshore variability in
habitat or circulation and find a different mechanism for the persistence of a cline.

**What Selection is Needed to Allow a Cline to Persist in a Uniform Advective Environment?**

Gradients in selection have often been invoked to explain the persistence of clines in systems
with extensive gene flow (Endler 1977, Hare et al. 2005, and many others). However, we will
show below that in the presence of non-random components of larval dispersal and uniform
habitat, there is a minimum level of selection needed to allow a cline to persist. To do so, we
examine the model of a benthic organism given above in the limit that the habitat’s carrying
capacity is alongshore uniform, and the species has been present for long enough that its
population has reached the carry capacity. In this limit, we quantify the minimum level of
selection needed to allow an allele to persist, and find that it depends sensitively on the mean
\( L_{adv} \) and stochastic \( L_{diff} \) components of larval dispersal distance. In order to do so, the increase
in the number of copies of an allele that is rare in the population per existing copy of the allele per
lifetime – the \( N_{allele} \) given above – will be calculated as a function of the selection coefficient \( s \)
(see Hartl and Clark 2000) against all other alleles in a population in which the population is
habitat limited everywhere. It will then be shown with a numerical model that when there is
selection for an allele sufficient to satisfy the criterion (4) in and only in a downstream portion of
a coastal ocean, a cline in allele frequency can persist in the coastal ocean.

In order to calculate \( N_{allele} \) from the selection \( s \), we assume that the population in the
habitat is limited by the scarcity of habitat, but the model could be easily extended to some other
density dependent process. We also assume the allele of interest is scarce in the population, for
as discussed above and in BP, this is the limit of interest for retention. \( N_{allele} \) can be written

\[
N_{allele} = (\text{Larvae per adult per lifetime})*(\text{success in competition with other larvae})*(\text{fraction of habitat free})
\]

(5)

“Larvae per adult per lifetime” is defined above as \( N_{larvae} \), and is net of mortality in the larval pool
and after settlement and before reproductive competency. \( N_{larvae} \) is defined for an adult that
carries the allele with less selective advantage. The “fraction of habitat free” accounts for the
fact that in this simple model, larvae that try to settle where there is an adult will die. The
fraction of habitat free is represented by the parameter \( \gamma \). If the larvae were to have sufficient
behavior to avoid habitat occupied by adults and to seek out nearby available habitat, \( \gamma \) would be
1.

The remaining term of (5) captures the relative reproductive success of adults with the
allele of interest, and is where selection enters the equation. It can represent either soft or hard
selection mechanisms. The coefficient \( s \) is the selection coefficient against other alleles, and is defined so that if an adult that does not carry the allele of interest produces \( X \) larvae that would successfully recruit, an adult that does carry the allele of interest produces \( X/(1-s) \) larvae that successfully recruit. This term can represent an ability to out compete other larvae for habitat, in which case it represents the likelihood that a single larva carrying the allele of interest arriving at an open piece of habitat can succeed in competition against \( N_{\text{other}} \) larvae without the allele. Or the term can represent a fractional increase in fecundity or survival in the larval or juvenile pool. Assuming that there are on average \( N_{\text{other}} \) other larvae competing for the same piece of open habitat, this term is

\[
\frac{1}{1+(1-s)N_{\text{other}}} \tag{6}
\]

The increase in the number of the selected allele can now be written

\[
N_{\text{allele}} = \frac{\gamma N_{\text{larvae}}}{\{1+(1-s)N_{\text{other}}\}} \tag{7}
\]

Because we assume the favored allele is scarce in the population, \( N_{\text{other}} \) in the above equation can be eliminated by noting that if there is no selection for the allele \( (s=0) \), it will not increase in the population and \( N_{\text{allele}} \) is one. \( N_{\text{other}} \) can then be found by setting \( s=0 \) and \( N_{\text{allele}}=1 \) in (7). This solution for \( N_{\text{other}} \) can then be substituted into (7) to give

\[
N_{\text{allele}} = \frac{\gamma N_{\text{larvae}}}{\{(1-s)\gamma N_{\text{larvae}}+s\}} \tag{8}
\]

In the limit of \( s<<1 \) and \( \gamma N_{\text{larvae}}>>1 \), this can be written as

\[
N_{\text{allele}} = 1+s \tag{9}
\]

If (8) is substituted into the criterion for the persistence of an allele into the middle of a population, (4), it reduces to

\[
s > L_{\text{adv}}^2/(2L_{\text{diff}}^2) \quad \text{if } s<1 \text{ and } \gamma N_{\text{larvae}}>>1 \tag{10a}
\]

\[
\ln(\gamma N_{\text{larvae}}) > L_{\text{adv}}^2/(2L_{\text{diff}}^2) \quad \text{if } s \to 1 \tag{10b}
\]

When (10) is true, an allele introduced into the middle of an alongshore-uniform domain will grow in number and fill the domain, because the increase in the number of copies of the allele due
Note that if \( L_{adv}^2/(2L_{diff}^2) \) is large enough and \( N_{larvae} \) small enough, an allele can be lost downstream even if it is very strongly selectively favored \((s=1)\).

The derivations of these results in BP were only tested in a domain in which the ecological and physical parameters did not vary in the alongshore direction. However, for a cline in allele frequency to persist, \((10)\) cannot be true everywhere, for if it were the downstream allele would take over the entire domain. We hypothesize that if the selection for the allele only satisfies \((10)\) in a downstream portion of the domain, the cline will not move upstream, and will occur immediately downstream of the location where selection becomes strong enough to satisfy \((10)\). When we test this conjecture in the numerical model of the population, it is found to be correct (figures 4 and 5), as long as the downstream region is large compared to \(L_{adv}\) and \(L_{diff}\).

This is the same criterion found for the validity of \((3)\) in BP and references therein.

Thus for a cline to persist in a uniform environment in which the population is limited by habitat scarcity, there must be a gradient in selection so that \((10)\) is satisfied only in the downstream portion of the domain. In this limit, the selection needed for a cline to persist depends on the larval dispersal parameters \(L_{adv}\) and \(L_{diff}\). This will be discussed further below.

However, it is first necessary to examine another limit, in which a cline is maintained by an alongshore variation in either the ocean circulation or the habitat, and the subsequent reduction in habitat scarcity.

Persistence of a Cline Allowed by Alongshore Variation in Habitat and Transport.

In the last section, the number of successful offspring an individual could leave was limited by the scarcity of habitat, and thus \(N_{allele}\) was driven by selection. The number of copies of an allele left by an adult can also be increased if each adult is able to leave multiple offspring.
because the environment is not, at least locally, habitat limited. These regions of enhanced reproductive success can be areas in which $N_{allele}$ satisfies (4), and thus can support a persistent cline. In order to understand these dynamics, we examine the model in the limit where alongshore variation in carrying capacity or larval transport produces regions of enhanced reproductive success.

Of course, if a species’ total population is in steady state, the average adult will only leave a single offspring behind after its death. However, this need not be true everywhere in the species range, for some regions can be source populations in which each adult has above average reproductive success, while other regions can be sink regions (figure 6). This pattern of sources and sinks can be seen most clearly in considering the entire range of a species in an environment with unidirectional mean alongshore current. Adults in the center of the domain leave, on average, one successful child per lifetime, for where the population growth is limited by density dependence children can only survive to replace adults that have died. Adults near the downstream edge of the species range largely fail to reproduce successfully as their progeny are, on average, swept downstream and beyond suitable habitat. On the other hand, adults on the upstream edge of the species range have a greater reproductive success rate because the larvae produced at the upstream edge of the species range serve to repopulate both this region and the area immediately downstream of it.

Similar regions of enhanced reproductive success can also form any place in the domain where the supply of larvae from upstream is insufficient to fully supply the downstream region with larvae, and so the population is below the local carrying capacity of the habitat. This can occur either when larval transport from upstream is interrupted, or when available habitat increases downstream. For example, the carrying capacity of the habitat triples downstream of the 1300km point of figure 6. Reproductive success is high here and the population is recruit limited, for the larvae produced in this region feel little competition from the small number of larvae advected in from the poor habitat upstream. Another region of high reproductive success
and reduced population occurs downstream of a region of reduced larval transport between kilometers 2500 and 3000, because few larvae are exported from there to settle downstream.

In these regions of enhanced reproductive success, eqs. (4) and (3) can be satisfied. Thus, upon their deaths, adults in the regions will usually be replaced with recruits that originated in the same region. These areas are thus retention areas (BP), and alleles in these regions, including any novel alleles that may arise in these regions, will tend to persist in these regions. Thus we can treat these retention regions as populations, in the classical population genetic sense. This will allow us to apply classical population genetic results to these areas, and to analyze the retention of a cline at these locations in terms of the migration between populations upstream and downstream of the change in habitat or circulation.

At the upstream edge of the species domain, there is no immigration from further upstream and the usual balance between mutation and drift will govern allelic diversity. In those regions of high reproductive success in the interior of the species range, allelic diversity will be governed not only by these processes, but also by the immigration of small numbers of individuals from upstream. To understand the relative importance of these processes, it is necessary to quantify the processes that create regions of high reproductive success and limit immigration from upstream. The parameter $f_{migrate}$ is the ratio of the larval supply from upstream to the supply that would occur if habitat and circulation in the retention zone extended infinitely upstream, so that a smaller number indicates fewer larvae from upstream settle in the region (at the upstream edge of the species range, $f_{migrate} = 0$). Where the reduced larval supply is formed by an interruption in the alongshore transport of larvae, $f_{migrate}$ is the fraction of larvae which can pass through the interruption. Where the reduced larval supply is caused by poor upstream habitat, $f_{migrate}$ is the ratio between the carrying capacity upstream of and in the retention region. When the population in the region of enhanced reproductive success is close to the carrying capacity, then $f_{migrate}$ is also approximately the ratio of migrants to local recruits that settle in the region of enhanced reproductive success (see appendix A).
If $f_{\text{migrate}}$ is much less than the mutation rate of a gene, classical population genetic results suggest that the migration rate would be insufficient to homogenize the populations upstream and downstream of the region of high reproductive success, and a cline could exist there solely due to stochastic fluctuations in the frequency of neutral alleles (Nagylaki 1978, Hartl and Clark 2000).

However, if $f_{\text{migrate}}$ is greater than the mutation rate, some selection for a downstream allele would be necessary to maintain it against immigration from upstream. The relative importance of immigration and selection in a deme has been extensively studied in cases of asymmetric dispersal (e.g. Kawecki & Holt 2002). In these studies, the frequency of the favored allele in the population receiving migrants is $(1-s/m)$ for $s<<1$, where $m$ is the ratio of migrants to local recruits in the retention region. Since $m \approx f_{\text{migrate}}$, the frequency of the favored allele in the retention region downstream of the variation in habitat or circulation is approximately

$$\text{Allele Frequency} = (1-f_{\text{migrate}}/s) \quad (11)$$

for $f_{\text{migrate}}<<1$, $s<<1$, where $s$ is the selection coefficient against all other alleles in the retention zone. This is illustrated in figure 7, and derived more formally, along with a discussion of its limitations, in appendix A. When $f_{\text{migrate}} > s$, the favored allele will be lost from the system and the cline will disappear. When $f_{\text{migrate}} < s$, the allele and the cline will persist where there is the change in alongshore larval transport or habitat.

At this point, it may seem that there is little difference between the classical problem of two populations exchanging migrants and an advective coastal environment with alongshore variation in larval transport or habitat. In both, reduction in larval transport between two regions increases the ability of a cline to persist by reducing the migration between the two populations [e.g. Slatkin 1985, Hare and Avise 1996, Rocha-Olivares and Vetter 1999, Lenormand 2002]. However, directional dispersal of the species changes the system in two fundamental ways. First, in a system with directional dispersal, if the reproductive success in the downstream region is reduced to the point that (4) is not satisfied, no non-zero $f_{\text{migrate}}$ will allow the favored allele to persist only in the downstream region. Population dynamics, in particular the reproductive
success of adults in the downstream region, fundamentally govern the ability of an allele to be
retained in an advective environment, and the enhanced reproductive success is enabled by
alongshore variation in circulation or habitat. Secondly, directional dispersal prevents an allele
from flourishing everywhere it is selectively favored, as can be seen in figure 7. In this figure, an
allele is favored in the downstream nine-tenths of the domain, and there is a reduction of larval
transport in the midst of the region where the downstream allele is favored. Since the selection
for the allele is less than that required to satisfy (10), the cline is not able to spread upstream of
the location of the interruption of larval transport. The cline is instead pinned to the region of
alongshore variability in larval transport, even though the downstream allele is favored upstream
of that point. Thus clines will persist not where an allele begins to be favored, but where there is
a retention zone driven by alongshore changes in circulation or habitat, or where the selection
exceeds the minimum set by (10).

Discussion

The maintenance of phylogeographic boundaries in a coastal ocean in a species with
a long planktonic duration: To maintain population genetic boundaries in the coastal ocean
with alongshore uniform circulation and habitat, there must be a gradient in selection for the
downstream allele. There must be sufficient selection downstream of the cline to retain the
downstream allele against advection, and insufficient selection upstream of the cline to prevent
the cline from moving upstream. In such a relatively simple coastal ocean, and for species with
uncomplicated spawning and larval behavior, we can estimate the critical level of selection at the
cline. Substituting (2) and (1) into (10) produces an expression for this critical selection threshold
as a function of the circulation parameters and larval duration:

\[
\frac{U^2 T_m}{2 \sigma^2 \tau_L} \quad \text{if } \frac{(U^2 T_m)(2 \sigma^2 \tau_L)}{} << 1
\]

\[
\quad \Rightarrow 1 \quad \text{if } \frac{(U^2 T_m)(2 \sigma^2 \tau_L)}{} >> 1
\]
Increased mean currents ($U$) and reduced variability in the currents ($\sigma$) will tend to increase the selection needed to retain an allele, while a decreased time in plankton ($T_m$) will reduce the selection needed to retain an allele.

For most coastal oceans, the strength of the mean current $U$ is comparable to strength of the fluctuations of the current $\sigma$, so the term $U^2/(2\sigma^2)$ in (12) is of $O(1)$ (Winant et al. 1987 for Central California, Lentz 2001 for the South Atlantic Bight, Beardsley et al. 1985 for the Mid-Atlantic Bight, Hickey & Banas 2003 for the North West, Gangopadhyay et al. 2003 for the Gulf of Maine, Boicourt et al. 1998 for the Gulf Coast). Thus the magnitude of the selection needed to retain an allele will depend on the ratio $T_m/\tau_L$. Where the larval planktonic duration $T_m$ is much greater than time scale of current fluctuations $\tau_L$, it will take strong selection to allow a cline to persist in place. If the larval planktonic duration is much shorter than $\tau_L$, it will take moderate levels of selection to allow an allele to spread upstream, and only moderate selection gradients to preserve a cline in place. The time scale of current fluctuations in most coastal oceans is on the order of a few days (Brink et al. 2000, Seigel et al. 2003, Davis 1985), and so moderate values of $s$ will suffice in species whose larvae disperse in the plankton for less than a few days. However, if larvae with simple behavior are in the plankton for much more than several days, the selection coefficient needed to allow a cline to persist increases dramatically. Kingsolver et al. (2001) find that the median of observed levels of selection on quantitative traits in natural populations is about $s=0.16$, though with a long tail of much higher selection levels. For single-locus patterns of selection, a number of examples in marine populations are available for which $s$ is apparently much higher (reviewed in Eanes 1999, Rand et al. 2002). For most species with a larval planktonic duration $T_m$ longer than a few days, clines can only be retained along a coast with relatively uniform current and habitat by high levels of selection.

A cline can also persist in a region where the mean current goes locally to zero or becomes small, allowing $L_{adv}$ to locally become much less than $L_{diff}$, and thus allowing weak
levels of selection to retain an allele (10). $L_{adv}$ may also become small due to larval or spawning strategies that enhance retention. BP describe several such strategies, such as the spawning of larvae in multiple seasons with oppositely directed mean currents to reduce $L_{adv}$. Hare et al. (2005) illustrate the importance of other mechanisms that may reduce transport across a region, all of which can lead to significant genetically controlled clines.

Additionally, even for species with long planktonic durations, a cline can persist where there is a reduction in larval pressure from upstream due to alongshore variation in circulation or habitat. These regions of persistence will not be fully captured in analyses that neglect alongshore variation in population dynamics. In these regions, the selection needed to retain the downstream allele in a cline is a function only of the strength of the disruption of larvae supply from upstream (11), and is not a function of larval duration. This allows clines to persist with modest gradients of selection even for species with long planktonic durations. If $f_{migrate}$ is less than the mutation rate for an allele, the cline can persist with no selection. Given the low median levels of selection seen by Kingsolver et al. (2001), this model suggests that most clines observed in species with long planktonic durations are associated with reduced larval supply from upstream.

Of course, the magnitude of the limitation of upstream larval supply $f_{migrate}$ might be a function of larval duration and dispersal distance. For example, $f_{migrate}$ could be small for larvae in the surface waters where there is a region of mean offshore transport of the surface waters, such as in an upwelling zone. In this case, a longer larval duration would increase the likelihood of a larvae being lost offshore. This would decrease $f_{migrate}$ and allow the cline to persist with less selection. The interruption of alongshore larval supply can also be driven by a region of poor habitat, which supplies few larvae to a downstream region. For larval dispersal distances less than the size of the region of poor habitat, no larvae could pass through this region from upstream, and $f_{migrate}$ would not be a function of the dispersal distance and larval duration. However, once the larval duration increased and dispersal distances $L_{adv}$ or $L_{diff}$ exceeded the size
of the region of poor habitat, \( f_{\text{migrate}} \) would increase abruptly and the selection needed to preserve a cline would increase. Thus our model suggests that, for species with long planktonic durations and large ratios of \( L_{\text{adv}} \) to \( L_{\text{diff}} \), clines will occur either in regions of elevated reproductive success driven by reduced larval immigration from upstream, or the cline will be maintained by levels of selection much larger than the median values found by Kingsolver et al. (2001), or both.

The Central California cline in \( \textit{B. glandula} \); an Example of a cline in a species with a long planktonic duration. \( \textit{Balanus glandula} \) is a well-studied barnacle present along much of the Pacific coast of North America (Newman and Abbott 1980). Sotka et al. (2004) describe a cline in both nuclear and mitochondrial allele families for \( \textit{B. glandula} \) across a narrow region of the Central Californian coast. This phylogeographic transition spans approximately 500km of coastline between Pacific Grove, CA and Cape Mendocino, CA (Sotka et al. 2004). For this analysis, we chose parameters appropriate for this species and for the oceanographic conditions in this region. Using current meter data, we calculate in appendix B that \( L_{\text{adv}} \) is approximately 290 km and \( L_{\text{diff}} \) 240km.

In the absence of any alongshore variation in habitat or circulation the selection needed to retain an allele here would be about \( s=0.7 \), if each adult barnacle could produce many surviving offspring in a sparsely populated area. This selection is much larger than the median seen by Kingsolver et al. (2001), though is within the range of selection coefficients observed. If the selection for the downstream allele was greater than this in the cline, and less to the north, a selection gradient alone could explain the cline.

However, it is likely that the circulation in this region would tend to greatly reduce the selection needed to allow the cline to persist, and thus reducing the required selection to a more typically observed value. Physical theory suggests that the change in the continental shelf width between Pt. Reyes and Monterey Bay will cause a reduction of alongshore transport between these two points, with the excess transport on the shelf moving offshore in the surface Ekman
layer (Pringle 2002). The reduction in the alongshore transport is expected to scale as the ratio of
the shelf width at the two locations, and thus there should be a five or tenfold reduction in
alongshore transport. Evidence from floats drogued to follow the surface waters (Sotka et al.
2004) supports this theory. Floats released to the south of the cline in late winter tend to move
along the coast and beach on the coast near Monterey Bay or to the south [e.g. the winter float
tracks in Winant et al. 2003], while floats released to the north of the cline tend to move offshore
between Pt. Reyes and Pt. Arena, and then stay offshore, never to return to the coast where a larva
could settle (Sotka et al. 2004, Brink et al. 2000, Davis 1985). Nearshore observations in
Monterey Bay support the idea that alongshore transport there is opposite to the prevailing mean,
and drives offshore transport (Storlazzi et al. 2003, Rosenfeld et al. 1994, and Graham and
Largier 1997). Thus the physical environment in the area of the cline provides the interruption of
the larval transport needed to allow a cline to persist. Studies of barnacle population dynamics in
the region of the cline support the idea that the population is limited by larval supply, and not by
habitat (Roughgarden et al. 1988 and references therein), suggesting that the interruption in larval
supply has set the stage for reproductive success, and thus the retention of alleles in this region.

Similar interactions between local adaptive processes and variation in nearshore physical
oceanography have been hypothesized to drive phenotypic and other macroevolutionary patterns

The present location and historical origin of a cline: Evidence suggests that the age of a cline
at a particular location can be much less than the age of the genetic differences that form the cline
(Felsenstein 1975, Endler 1977). The differentiation of the allelic classes which form clines in
the B. glandula population appears to have formed > 100 kya (Wares and Cunningham, 2005).
However, the change in shelf width that Pringle (2002) posits as the cause of the interruption of
larval transport at the current cline location between Monterey and San Francisco Bay only exists
at current sea levels. When the sea level was ~ 100m lower in the last ice age (Graham et al.
2003), the shelf would have vanished, and the proximate physical cause of the cline would
disappear. Thus the current mechanism for fixing the cline in place may have only existed for the
last 10,000 years.

However, this model suggests a cline could persist even without refugia. If the current
interruption of larval transport in central California were to be removed, the cline would be
swept downstream until it re-established at the next location along the coast in which larval
transport were interrupted, or one of the other physical or biological mechanisms for cline
formation discussed above existed. If the mean currents experienced by larvae were always in
the same direction, and the retention zones are sufficiently separated as to prevent larval dispersal
from the downstream to upstream retention zone, then the cline would ratchet downstream from
one extent retention zone to the next as the retention zones came and went. For these reasons,
tracking phylogeographic boundaries in marine populations necessarily involves understanding
contemporary forces that limit the movement of individuals or their larvae. Many of these
transitions, along with biogeographic transitions, may have originated due to historical events; but
their current location may not be the site at which the transition originated.

The relation of the above theory to existing phylogeographic theory: It is well established that
the location and strength, or shape, of a cline can be modified by stochastic allele frequency drift
(Felsenstein 1975, Slatkin and Maruyama 1975), and a large literature looks at the effects of
selection and genetic drift in demes that undergo asymmetric migration (e.g. Nagylaki 1978,
Holt 2002). Some of the results derived above are foreshadowed in these papers, and the links
can now be made explicit. Nagylaki (1975) showed that an allele that was favored in a region
could be lost if there was a large amount of migration into that region from areas in which the
allele was not selected for. Nagylaki (1978) pointed out that it was easier to maintain an allele in
a deme which generated a net outflow of migrants than in one which was a sink for migrants, and
Kawecki & Holt (2002) argued that selection in sink regions would be relatively ineffective (c.f. Doebeli 1995 and Lebreton et al. 2000). Lundy & Possingham (1998) argued that a deme would only contribute to preventing the fixation of an allele if it were a source of migrants.

However, to place the results of these papers in the context of a coastal ocean, it is necessary to identify the mechanisms that form populations or demes there. The tendency has been to label regions of relatively uniform allele frequency as a population. This is not appropriate, for in most of these regions any novel allelic diversity or stochastic fluctuation in allele frequency that arises will be swept downstream by the mean currents, and will not contribute to the evolution of allelic diversity at that point. We can only discover the boundaries between populations by considering the spatial structure of habitat and circulation.

Regions downstream of a reduction in larval supply can retain stochastic fluctuations in allele frequency and novel alleles that arise, and the region of enhanced reproductive success can be considered a population. These regions also export larvae downstream, so they are source regions for the downstream population. As Kawecki & Holt (2002) argue, selected alleles are relatively easily retained in these regions, even if there is some migration into the population from upstream. If immigration of larvae into this region is less than the mutation rate in the population, a cline can exist even in the absence of selection. At the upstream edge of the species range, where there can be no immigration of larvae from upstream, allele frequency will change as one would expect for a population with symmetric dispersal (e.g. Kimura 1968).

The sink populations are those downstream of the regions of enhanced reproductive success, in which density dependence limits the realized growth rate of the population. At each of these regions, the population is maintained by immigrants from upstream, and the region maintains the population downstream of it by exporting its larvae downstream. As predicted by Nagylaki (1978), Kawecki & Holt (2002) and others, it takes a very large selection coefficient to maintain an allele in the population in these sink regions (equation 10). Absent strong selection,
allele frequency at downstream regions is set by the gene frequency $L_{adv}$ upstream, until in the end it is set by the allele frequency at an upstream region of enhanced reproductive success.

In these downstream regions, the substitution rate will not equal the mutation rate. Mutants will arise at a rate proportional to the population size, but will on average be distributed further downstream each generation; given enough mutants over large distances with large population sizes, a stochastic equilibrium defined by mutation and upstream diversity will hold. This would predict increased levels of allelic diversity in downstream ‘sink’ habitats (Wares et al. 2001, Al-Rabab’ah and Williams 2004). However, as seen in figure 2 and in the associated text, the distribution of any mutant allele would shift downstream by $L_{adv}$ each lifetime. For species with mean planktonic dispersal distances of tens to hundreds of kilometers, and species ranges of thousands of kilometers in size, any mutant allele which arises outside of the downstream edge will be lost off the downstream edge of the domain in a few tens to hundreds of generations, a time perhaps insufficient to allow significant allelic diversity to develop (e.g. Hoskin 2000).

**Marine protected areas, fisheries, and the transport of clines:** The genetic homogeneity of a population along the coast has been used to argue that the population should be treated as a single population, a “stock,” for the purposes of fisheries management. The genetic homogeneity has been assumed to indicate that the population is panmictic, and thus individuals in any part of the population can replenish other parts of the population. But, as described above, this is not true in downstream regions which are sink regions, and whose population is maintained by immigration from upstream.

In such a population, harvesting could even encourage the formation of sub-populations, if the harvesting in the interior of the domain created a region of small population in the midst of a range, thus creating the interruption in larval transport needed to create locally high realized reproductive success and thus the local retention of allelic diversity. Conversely, sufficient harvesting could destroy a cline, and cause the upstream genome to invade downstream. Clines
can be maintained by weak selection in local regions where the realized growth rate is high enough to satisfy (4), as discussed above. If a fisheries reduces the growth rate by harvesting reproducing adults, the growth rate could well fall below (4), preventing retention, and allowing the cline to be swept downstream. The local population would then be taken over by the upstream population, which, while less adapted to local conditions, is anchored by some more robust retention zone farther upstream.

**Directional dispersal in other environments:** Similar genetic dynamics to those described above will exist in any system in which the dispersal of propagules is anisotropic with a preferred direction of dispersal, for example in populations with wind-driven dispersal (e.g. Antonovics 1968, Kavecki and Holt 2002, Al-Rabab’ah and Williams 2004) or populations in the open ocean. Qualitatively, the above results hold. However, the quantitative results for retention and the maintenance of clines described above will only hold if the dimension of the population range perpendicular to the direction of preferred dispersal is small enough that the population can disperse across it in $O(1)$ generations. If it is not, the quantitative results described above must be modified to include tendency of isotropic dispersal normal to the direction of preferred dispersal to reduce the growth rate at a fixed point in space, a calculation similar to the classic calculations of the relative importance of dispersal and growth made in Kierstead and Slobodkin (1953) and Wroblewski et al. (1975).

**Summary:** The population genetics of the coastal ocean has long been understood as resulting from the interplay of selection, random genetic drift, and migration (e.g. Palumbi 2003 and references therein). The interplay of these mechanisms in the coastal ocean for organisms with planktonic larval stages is simultaneously governed by the ocean circulation and local population dynamics. Thus it is only by taking the currents and population dynamics into account that the population dynamics analysis of demes and sub-populations can be mapped onto
a continuous coastal ocean, with little in the way of natural boundaries. In doing so, it has become clear that the maintenance of polymorphism in these systems is modified by advection and local population dynamics. A cline will not usually be located where an allele becomes favored. Instead, the cline will persist either where the selection for an allele begins to exceeds (10) or where an alongshore variation in habitat or circulation produces a fractional reduction in the upstream larval supply greater than the selection for an allele or the mutation rate. For species with long planktonic durations, our model suggests the latter is more likely to set the location of a cline.

Acknowledgements: This is GLOBEC submission [WILL OBTAIN WHEN ACCEPTED], and was funded by NSF OCE-0219709 to J.Pringle. Thanks go to R. K. Grosberg, S. T. Small, and R. J. Toonen for comments on the manuscript.
Appendix A: Maintenance of a cline against migration from upstream.

To quantify the relative importance of immigration and selection in the retention region downstream of alongshore variation in the circulation or habitat, we examine the allele frequency of an allele which is favored downstream of the change in circulation or habitat. The relative frequency of this allele in the retention region downstream of the interruption is $R_d$, and the total number of alleles in that region $P_{reten}$. We wish to find $R_d$ as a function of the selection $s$ against all alleles but the favored downstream allele and the fractional reduction in the ratio of larvae entering the retention region from upstream $f_{migrate}$. As part of the derivation, it is useful to define $N_m$, the number of copies of the other alleles that enter the retention region to compete for habitat.

Parameters defined only in this appendix are not listed in table 1.

The number of copies of alleles which return to the retention region each lifetime will scale as $N_{larvae}P_{reten}$. We assume that the immigration $N_m$ is much smaller than $N_{larvae}P_{reten}$, and so the number of copies of the favored downstream allele that compete for available habitat each lifetime scales as $N_{larvae}P_{reten}R_d$, and the number of copies of the other alleles are $N_m + N_{larvae}P_{reten}(1-R_d)$. From (6), we can then write the fraction of the larvae carrying downstream allele which compete successfully for the available habitat as

$$\frac{(N_{larvae}P_{reten}R_d)}{[N_{larvae}P_{reten}R_d + (1-s)(N_m + N_{larvae}P_{reten}(1-R_d))]} \quad (A1)$$

If the allele frequency is in steady state, the fraction of the larvae carrying the downstream allele that recruit successfully must equal the fraction of the allele in the population, and so we can solve for $R_d$ by setting it equal to (A1) and solving for it. This results in

$$R_d = 1 - \frac{N_m}{(N_{larvae}P_{reten})}(1-s)/s = 1 - \frac{N_m}{(P_{reten}N_{larvae})}/s \text{ if } s<<1 \quad (A2)$$

This is the classic result that the balance of immigration and selection is governed by the ratio of selection $s$ to the fractional migration rate $m = N_m/(N_{larvae}P_{reten})$ when $s<<1$.

In order to get an estimate of how $N_m/(N_{larvae}P_{reten})$ varies with the strength of $f_{migrate}$, we assume that $P_{reten}$ is of the same order as the caring capacity of the retention zone. This
assumption is valid when $N_{\text{larvae}}$ is large, as in figure 6. Then, since without the interruption in larval transport, $N_{\text{larvae}}P_{\text{reten}}$ larvae from upstream would attempt to settle in the retention zone per lifetime, with the interruption there would be $N_m = f_{\text{migrate}}N_{\text{larvae}}P_{\text{reten}}$ larvae from upstream. The fraction of the downstream allele in the retention zone is then

$$R_d = 1 - \frac{f_{\text{migrate}}}{s} \quad \text{if } s << 1$$

Equation (A3) becomes less than zero when the selection rate is not greater than $f_{\text{migrate}}$, which implies that the cline would not be maintained. Where it is greater, a cline can be maintained. The abundance of the downstream allele increases as one moves farther downstream from the retention zone, for selection will serve to increase the abundance of the allele as it is swept downstream from the retention zone.

The derivation for (A3) above makes three approximations that limit its accuracy. First, it assumes that there are none of the favored allele upstream of the variation in habitat or circulation, so that all migrants from upstream contain the un-favored alleles. Second, it assumes that $P_{\text{reten}}$ is equal to the carrying capacity of the habitat in the retention zone, while it must actually be somewhat less to allow the enhanced reproductive success needed to preserve the cline. And third, it assumes that larvae spawned in the retention zone are as likely to return to the retention zone as those from immediately upstream. The first assumption would increase $R_d$, while the second two decreases it. The exact impact of these assumptions depends on the exact nature of the alongshore variation of circulation and habitat, however in cases we have modeled it is relatively minor, changing the selection needed to allow retention by less than 30%. Equation (A3) is compared to model runs in which $f_{\text{migrate}}$ is set to 0.05 and 0.10 by removing $(1 - f_{\text{migrate}})$ of the larvae crossing a point 3/4 of the model domain from the upstream edge. The downstream allele is favored in the downstream 9/10 of the model domain. The models are then run for a large number of differing values of $s$, and the concentration of the downstream allele is shown in figure 7. Eq. (A3) is seen to capture the variability of $R_d$ with $s$ well.
Appendix B: *Balanus glandula* dispersal and life history.

*B. glandula*, like many other decapod crustaceans, has a planktonic larval form that feeds and develops in the water column for 3-4 weeks before becoming competent to settle (Brown & Roughgarden 1985). *B. glandula* broods larvae from Feb-Oct with the majority (about 90%) being released early, and about 10% of the release from Aug-Sep (settlement is then Sep-Oct) (Hines 1978). The brooding cycles for *B. glandula* are similar for populations in British Columbia, central California, and southern California (Barnes and Barnes 1956), and late larval stage individuals are found within the top 0.5m of seawater (Grosberg 1982).

$L_{\text{adv}}$ and $L_{\text{diff}}$ for *B. glandula* in the vicinity of the cline can be estimated from the mid-shelf alongshore currents measured as part of the Coastal Ocean Dynamics Experiment in the northern part of the cline, between Pt. Reyes and Pt. Arena. Lentz & Chapman (1989) estimate a mean alongshore current $U$ in the spring and early summer mid-shelf surface waters of $14 \text{ cm/s}$ southward with a standard deviation $\sigma$ of $24 \text{ cm/s}$, and a fall mean current of $7 \text{ cm/s}$ northward with a standard deviation of $18 \text{ cm/s}$. Davis (1985) estimates that the Lagrangian timescale $\tau_L$ was about 3 days. The net larval dispersal combining the early- and late-season larval releases was calculated as in BP, in which the values of $L_{\text{adv}}$ and $L_{\text{diff}}$ for each spawning event is weighted appropriately by the fraction of larvae released in each event, resulting in an estimate of $L_{\text{adv}}=287$ km and an $L_{\text{diff}}=240\text{km}$. No attempt was made to compensate for different mortality rates in the late spring versus late summer, for there is insufficient information to do so, though it is likely that more larvae are lost offshore in the spring upwelling season as surface waters are transported offshore by Ekman transport (Roughgarden *et al.* 1988). A relatively higher mortality in the spring than in the fall due to offshore advection would increase $L_{\text{diff}}$ and decrease $L_{\text{adv}}$.

Shanks *et al.* (2003) find for many species that behavior reduces the mean dispersal distance, though it remains linearly related to the time in plankton. How this would affect the results for *B. glandula* is unclear, for two reasons. First, there are not sufficient studies of the
behaviour of *B. glandula* larvae to determine the magnitude of any behavior related reduction in larval transport. Secondly, the effects on retention and allele persistence depend on the ratio of $L_{adv}$ to $L_{diff}$, and it is even less clear how this ratio might be affected by behavior. See (BP) for more details on the interaction of behavior with retention.
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Table 1: Parameters defined in the main section of the paper.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Definition</th>
</tr>
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<tbody>
<tr>
<td>$f_{migrate}$</td>
<td>Fractional change in larval transport from upstream. An $f_{migrate}$ of 0.2 indicates that the transport of larvae from upstream of a point is one-fifth of what it is elsewhere.</td>
</tr>
<tr>
<td>$\gamma$</td>
<td>Fraction of the habitat that is not occupied by adults. If larvae have sufficient behavior to avoid settling on occupied habitat, $\gamma=1$.</td>
</tr>
<tr>
<td>$H_{dens}$</td>
<td>Maximum population per unit length of habitat.</td>
</tr>
<tr>
<td>$L_{adv}$</td>
<td>Mean distance larvae recruits downstream of its parents.</td>
</tr>
<tr>
<td>$L_{diff}$</td>
<td>Standard deviation of distance larvae recruits downstream of their parents.</td>
</tr>
<tr>
<td>$m$</td>
<td>Fraction of larvae in a retention zone entering from upstream of the retention zone.</td>
</tr>
<tr>
<td>$N_{allele}$</td>
<td>Number of additional copies of an allele in the population for each existing copy of the allele in the population a lifetime earlier.</td>
</tr>
<tr>
<td>$N_{larvae}$</td>
<td>Number of larvae an adult produces per lifetime that are capable of recruiting successfully and reaching reproductive age, if habitat is available. This number is net of all mortality in the larval pool and after settlement.</td>
</tr>
<tr>
<td>$s$</td>
<td>Selection against all alleles but the allele of interest, as defined in Hartl and Clark (2000).</td>
</tr>
<tr>
<td>$\sigma$</td>
<td>Standard deviation of Lagrangian alongshore currents experienced by planktonic larvae.</td>
</tr>
<tr>
<td>$\tau_L$</td>
<td>Lagrangian timescale of alongshore currents experienced by planktonic larvae.</td>
</tr>
<tr>
<td>$T_m$</td>
<td>Time larvae spend in plankton.</td>
</tr>
<tr>
<td>$U$</td>
<td>Mean alongshore current speed experienced by larvae in plankton.</td>
</tr>
</tbody>
</table>
Figure 1. A cartoon of the larval parameters $L_{adv}$, $L_{diff}$, and $N_{larvae}$ showing the position of a single parent and the offspring left by that parent after a lifetime.
Figure 2: The evolution of a cline between two selectively neutral alleles, here represented as grey and black, in the numerical model. The larvae move on average $L_{adv}=100$ km downstream to the right each generation, with a dispersion around that mean of $L_{diff}=150$ km. The habitat is saturated, so each adult leaves one surviving offspring at its death, on average.
Figure 3. The evolution of allele density in two populations in which an allele is introduced near $y=0$. Allele density is the number of copies of a given allele per unit length. In both, $L_{adv}=300\text{km}$ and $L_{diff}=200\text{km}$. In the top panel, $N_{allele}$ is 2, which was insufficient to allow the introduced allele to persist at $y=0$. In the bottom panel, $N_{allele}$ is 4, which is sufficient to allow retention. In both panels the number of copies of the allele are increasing with time. However, note that at $y=0$, the allele density is decreasing in the top panel, while allele density is increasing at $y=0$ in the bottom panel, and the allele is also spreading upstream.
Figure 4: The allele frequency of the favored allele in the downstream portion of the model domain, as seen in figure 5. The allele is only favored in the downstream portion of the species range. The allele frequency is shown as a function of the selection $s$ normalized by the minimum selection needed to satisfy (10), $s_c$. Where the frequency of the allele is zero, selection was insufficient to retain the cline, and the downstream allele was flushed downstream and out of the habitat. Where the density is finite, selection was sufficient to maintain the cline against the mean current. The model was initialized with the favored allele in the downstream half of the domain, and the model was run until the allele frequency had stabilized. The species modeled had an $L_{\text{diff}}$ of 200km, and $L_{\text{adv}}$ is varied to change the critical selection needed to allow retention. $\gamma$ in these runs was $\approx 1.0$, for the modeled species is semelparous and spawns at the end of its lifetime.
Figure 5: The allele frequency of an allele which is selectively favored downstream (to the right) of the 1000km point of the domain, but not upstream of that point, in a numerical model. The selection is 40%, 80% and 150% greater than that needed to satisfy (10). The initial and boundary conditions are as in figure 5. $L_{adv}=100\text{km}$, $L_{diff}=200\text{km}$, and the critical value of selection to allow the cline to persist is $s=0.1958$ for a semelparaous species.
Figure 6: A) The distribution of population density as calculated by the numerical model for a finite habitat with a region of poor habitat capable of carrying only 1/3 of the normal population between 0 and 1300km, and a region of weak alongshore flow and dispersion (1/10th normal) between 2500 and 3000km. The mean current is flowing to the right, and $L_{adv}=L_{diff}=400$km outside the region of reduced alongshore flow and dispersion. B) The actual reproductive success in surviving offspring per adult, with the critical rate to satisfy (4), 1.65, marked with a horizontal line.
Figure 7: (Top) The steady-state allele frequency of the favored allele in the numerical model. The favored allele is favored downstream of 500km, and the barrier to larval transport is at 6100km. Organisms with the favored allele are released just downstream of the barrier at the start of the numerical model run, and the model is run until the populations reach steady state.

The favored allele is selected by \( s = 0.16 \), and 95% of all larvae are prevented from crossing the indicated barrier to larval transport in either direction. (Bottom) to the allele frequency of the favored allele in the population in the retention zone downstream of the barrier of larval transport as a function of the selection for the downstream allele for both \( f_{\text{migrate}} = 0.05 \) and 0.1, and the prediction of this ratio from (11). The selection is scaled by \( f_{\text{migrate}} \), so that all curves will lie upon each other. For both panels, \( L_{\text{adv}} = 200 \text{km}, L_{\text{diff}} = 300 \text{km}, \) and \( N_{\text{larvae}} = 6 \). A discussion of the validity of and errors in (11) is given in appendix A.