On the meaning and existence of an effective population size

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ABSTRACT

We investigate conditions under which a model with stochastic demography or population structure converges to the coalescent with a linear change in time scale. We argue that this is a necessary condition for the existence of a meaningful effective population size. We find that such a linear time scale change is obtained when demographic fluctuations and coalescence events occur on different time scales. Simple models of population structure and randomly fluctuating population size are used to exemplify the ideas and provide an intuitive feel for the meaning of the conditions.

INTRODUCTION

In population genetics, simplifying assumptions are necessary in order to turn complex biological systems into caricatures that are, on the one hand, simple enough to analyze, and on the other hand, realistic enough to capture key features of the process under investigation. These simple models often make assumptions that are clearly violated in most populations; yet they are of great importance since their simplicity allows one to make predictions about the patterns of polymorphism that are expected under these assumptions. Data can then be compared with these predictions to detect deviations from the simplifying assumptions. In other words, these simple models serve as null models. The standard population genetic null model, the Wright–Fisher model, and its retrospective cousin, the standard coalescent, are examples of this.

These null models have proved surprisingly difficult to reject, despite what appear to be major violations of the assumptions. On the positive side, this means that they manage to capture something essential about the way natural populations behave; in other words, they are robust to changes in the assumptions and are therefore good approximations to real systems. More disturbingly, this lack of sensitivity also implies that certain seemingly important features of the population history cannot be detected using polymorphism data.

The “robustness” of the coalescent process is one of the three properties listed by KINGMAN (2000) as fundamental to this process. Even if a natural population does not fulfill all the assumptions of the Wright–Fisher model, it can sometimes behave in all important respects (i.e., those that are observable from a sample, and hence derivable from the genealogy) like a Wright–Fisher population. This happens when the ancestral process can be approximated by Kingman’s coalescent with the population size replaced by a so-called coalescent effective population size, $N_e$.

The concept of effective population size predates the coalescent era and has traditionally been used to rescale a given population model so that it behaves, with regard to certain properties, like a simple Wright–Fisher model with constant population size. The three properties of the Wright–Fisher model most commonly used in defining effective population sizes are: (i) the probability of identity by descent of two alleles chosen at random, (ii) the variance in offspring allele frequency and (iii) the leading non-unit eigenvalue of the allele frequency transition matrix. They correspond, respectively, to the “inbreeding effective population size,” the “variance effective population size” and the “eigenvalue effective population size.” In some demographic models the three effective sizes are equal but in others they can differ considerably or simply not exist (EWENS 1982; ORIVE 1993). Even when they can be defined, they may lead to complex formulae involving demographic parameters that are practically impossible to measure.

We propose that the coalescent effective population size, when it exists, provides a more general and consistent notion of effective size that is less likely to be misused. Too often, one reads of “the” effective population size without reference to the particular notion being considered. Since
the coalescent essentially embodies all of the information that can be found in sampled genetic data, one can argue that, if anything deserves the title of “the effective size,” it is the coalescent effective size.

When population genetic data is consistent with that expected under a neutral, panmictic, constant-size Wright–Fisher model, one would like to say that there exists an effective population size. In fact, our point of view is that the existence of an effective population size is equivalent to a situation in which it is not possible to reject the basic Wright–Fisher model. This condition is more informative than the classical concepts of effective size, even in a theoretical setting, since the existence of, say, an inbreeding effective size does not imply that the population behaves like a Wright–Fisher model in any other respect. By definition, the coalescent effective size only exists when the scaling of time to retrieve the standard coalescent is independent of time; but, when this is the case, the appropriately re-scaled population behaves precisely as the Wright–Fisher model in all respects. We shall therefore assess conditions under which the coalescent effective size exists, both analytically and through computer simulations. In the simulations we use two simple demographic models, one with randomly fluctuating population size, and the other with subdivided populations linked by migration.

The coalescent, is a random tree that allows one to characterize ancestral relationships between individuals (genes) in a sample when the population size is (reasonably) large (Kingman 1982a,b,c). The probabilistic structure of Kingman’s coalescent (sometimes referred to as the standard coalescent) is quite simple. If we start with a sample of \( n \) individuals, we wait a random time \( T_n \) that is exponentially distributed with mean \( 1/(\binom{n}{2}) \). At this time, two randomly chosen ancestral lineages coalesce, leaving \( n-1 \) distinct lineages. The lineages continue coalescing in this way until we reach a single common ancestor for the sample. We thus obtain a sequence \( T_n, T_{n-1}, \ldots, T_2 \) of inter-coalescence times that are independent and exponentially distributed with

\[
E[T_k] = 1/c_k, \quad k = 2, \ldots, n,
\]

where

\[
c_k = \left( \frac{k}{2} \right) = k(k-1)/2
\]

is the number of ways to choose an unordered pair from \( k \) objects. The time to reach the most recent common ancestor is thus the sum

\[
T_{\text{MRCA}} = T_n + T_{n-1} + \cdots + T_2
\]

with expected value

\[
E(T_{\text{MRCA}}) = \sum_{k=2}^{n} \frac{1}{c_k} = 2(1 - 1/n).
\]

Since the standard coalescent corresponds to a neutral Wright–Fisher model, the pairs of lineages that join at coalescence times are always chosen at random. Thus, the probabilistic structure of the coalescent tree is determined by the pure death process that keeps track of the number of ancestral lineages as time recedes into the past.

How Kingman’s coalescent relates to the ancestry in a given population genetic model is described most easily for the haploid Wright–Fisher model with fixed population size \( N \), and no selection or recombination. In this panmictic model, when \( N \) is sufficiently large and time is measured in units of \( N \) generations, the ancestry of a sample is approximated by Kingman’s coalescent.

Thus to do a calculation for the Wright–Fisher model one does the analogous calculation for the coalescent process and then interprets \( t \) units of coalescent time to be \([Nt]\) generations, where \([Nt]\) is the largest integer less than or equal to \( Nt \). For example, the mean time to go from \( k \) lineages to \( k-1 \) in the coalescent is \( E[T_k] = 1/c_k \). Thus, in the Wright–Fisher model with population size \( N \), it takes on average \( N/c_k \) generations for \( k \) lineages to coalesce down to \( k-1 \). With the appropriate scaling of time this approximation works well beyond the Wright–Fisher model and there are variations of the coalescent that incorporate the effects of selection, recombination, spatial structure, and demographic variation.

Mathematically, the above can be expressed as follows. Let \( A(t) \) denote the number of lineages in the standard coalescent \( t \) units of (coalescent) time in the past, and let \( A_N(\tau) \) be the number of ancestors \( \tau \) generations in the past for the discrete-time neutral Wright–Fisher model corresponding to fixed population size \( N \). Convergence to the coalescent then means \( A_N([Nt]) \to A(t) \), as \( N \) tends to infinity. If we are dealing with a different population model that has some quantity fluctuating over time, we will say “averaging occurs” if the corresponding discrete-time ancestral process satisfies \( A_N([Nt]) \to A'(ct) \) for some constant \( c \). This means that, after converting \( t \) units of coalescence time to \([Nt] \) generations, the ancestral process for the discrete-time model can be approximated by the standard coalescent with time speeded up by a factor \( c \). This scaling factor \( c \) then allows us to define the coalescent effective population size by

\[
N_c = N/c.
\]

Notice that speeding up the coalescent by a factor \( c \) is equivalent, as far as sample data is concerned, to multiplying the mutation rate by the factor \( 1/c \). Thus, we get the same genealogy as for a neutral, panmictic, constant size Wright–Fisher model with a different mutation rate. Indeed, if there is a scaling constant \( c \) such that the appropriately-scaled ancestral process converges to \( A'(ct) \), then the sampled data cannot be distinguished from that arising in a standard neutral Wright–Fisher model.
In this paper, we seek to determine when, in the presence of stochastic demography (e.g., fluctuating population size or spatial structure), we can define an effective population size, $N_e$, that allows us to do, approximately, all coalescent-based calculations in the same way as we would for a Wright–Fisher model with size $N_e$. Our approach involves both theoretical analysis and simulations.

Our analytical results will provide general insight into the effects of fluctuating population size and geographical structure on the genealogical process. We will see, for example, that these effects can be averaged to get a coalescent effective size if population size fluctuations and migration rates are sufficiently rapid. To get a feel for how these limiting results apply to real populations and to gauge their robustness, we use simulations. For this, we quantify the effects of deviations from the standard constant-size Wright–Fisher model with Fu and Li’s $F$ statistic (Fu and Li 1993), one of many statistics designed to detect such deviations. The $F$ statistic is defined by

$$F = F(\pi, \eta_s, S) = \frac{\pi - \eta_s \left( \frac{n-1}{n} \right)}{\sqrt{\mu_F S + \nu_F S^2}},$$

where $n$ is the sample size, $\pi$ is the average number of pairwise nucleotide differences (the average being over all possible pairs in the sample), $S$ is the number of segregating sites, $\eta_s$ the number of singletons (mutations that appear in only one individual in the sample), and $\mu_F$ and $\nu_F$ are constants given the sample size $n$. This construction yields an expected value that is nearly zero (actually, it is slightly negative), assuming the standard model and the infinitely many sites model, but is expected to deviate from zero when the assumptions are not met. For example, fluctuating size tends to produce negative values of $F$ and population subdivision leads to positive $F$.

The rest of the paper is arranged as follows. In the next section, we discuss the case of randomly fluctuating population size. We begin with a fairly thorough treatment of the analytical results that describe when one can and cannot get a coalescent effective population size. This is followed by simulations of Fu and Li’s $F$ for a special case of the model. This section will be followed by a more abbreviated one dealing with population structure. Again, we begin with an analytical discussion and follow it with simulations in a special case. A discussion section summarizes the results and in the Appendix we show how to get the nonlinear time change for “intermediate” rates of population size fluctuations.

**FLUCTUATING POPULATION SIZE**

In this section, we discuss the effects of stochastically fluctuating population size on haploid, neutral, single-locus genealogies. This differs in a fundamental way from coalescent theory in the presence of deterministically-varying population size.

If these size fluctuations are fast compared to the coalescent time scale, then they will affect the coalescent only in an average sense. In this case there will be an effective population size and the genealogy will be given by Kingman’s coalescent with a linear time change. If, on the other hand, “macroscopic” size fluctuations occur on the same time scale as coalescences, then the resulting genealogical process will be described by Kingman’s coalescent run on a nonlinear, stochastic time scale. In this case, there is no effective population size. The object that one would like to think of as an effective size in this case changes with time instead of being constant; essentially, there is only an “instantaneous” effective size.

**Fast fluctuations – averaging**

One often sees in population genetics the claim that, when population sizes fluctuate, there is an effective size given by the harmonic mean of the possible sizes. To understand when this works and, more importantly, when it does not, let us begin with a simple calculation.

Suppose the sizes of a population have some fixed discrete set of possible values, denoted by $N_1, N_2, \ldots$, and assume that these sizes are all multiples of some large value $N$; say $N_i = x_iN$ for each $i$, where the $x_i$ are fixed positive numbers. As is typical in coalescent theory, we think of $N$ as a parameter which gives the magnitude of population size. Denote by $M_N(0), M_N(1), M_N(2), \ldots$ the sequence of population sizes backwards in time; i.e., $M_N(0)$ is the size of the current generation, $M_N(1)$ is the size of the previous generation, etc. The simplest possible model of randomly fluctuating size would assume that $M_N(0), M_N(1), M_N(2), \ldots$ form an independent, identically distributed sequence with probabilities $p_i = \mathbb{P}(M_N(\tau) = N_i)$ for each time $\tau \geq 0$. Suppose for simplicity that we have a Wright–Fisher model of reproduction, so that the probability of two randomly chosen individuals in generation $\tau - 1$ having a common parent in generation $\tau$ is $1/M_N(\tau)$. Then we compute the probability that two individuals do not have a common ancestor $[Nt]$ generations in the past (i.e., their ancestral lineages have not yet coalesced):

$$\mathbb{P}(\text{no coalescence in } [Nt] \text{ generations})$$

$$= \mathbb{E} \left[ \prod_{\tau=1}^{[Nt]} \left( 1 - \frac{1}{M_N(\tau)} \right) \right]$$

$$= \left( 1 - \sum_i p_i \cdot \frac{1}{x_i N} \right)^{[Nt]}$$

$$\rightarrow \exp \{- t \sum p_i / x_i \}$$
as $N \to \infty$. In the first equality, we conditioned on the values of the population sizes and then averaged over all possibilities; in the second equality we have used the assumption that sizes were i.i.d. to bring the expectation inside the product and turn the product into a power.

This calculation suggests that the pairwise coalescence rate, when $N$ is large, should be given by $\sum p_i/x_i$. This means that the pairwise coalescence probability in one generation is of the form $\frac{1}{N} \sum p_i/x_i$. To match the Wright–Fisher dynamics, this last quantity is set to one over the effective population size; i.e., the effective size $N_e = (\sum p_i/N_i)^{-1}$ is given by the harmonic mean of the possible sizes.

The above calculations depend on population size being independent between generations, which is hardly a realistic assumption. However, using the methods in Nordborg and Krone (2002), one can extend this to the case where the sequence $(M_N(\tau))_{\tau \geq 0}$ is allowed to change every generation according to a discrete-time Markov chain with state space $\{N_1, N_2, \ldots\}$ and unique stationary distribution $(\pi_1, \pi_2, \ldots)$. Then, as in the i.i.d. example, the effects of fluctuating size “average” between coalescence events, this time giving an effective pairwise coalescence rate of $\sum \pi_i/x_i$ and hence an effective size $N_e = (\sum \pi_i/N_i)^{-1}$ which is again a harmonic mean with the averaging being done with respect to the stationary distribution. Jagers and Sagitov (2003) obtain similar results for general reproduction models and stationary Markovian population size with a finite number of states.

**Slow fluctuations**

Similarly, if the large changes in population size are sufficiently slow compared to coalescence events, then in the limiting $(N \to \infty)$ coalescent all the coalescences will occur before there are any changes in size. This means that the limiting coalescent will correspond to a model in which the population size does not change and hence, in a simplistic way, one can think of having an effective size given by the initial size. This situation does not, however, entail averaging.

**Intermediate fluctuations – no averaging**

The remaining case, in which “large” changes in population size occur on the same time scale as coalescence events, has been treated mathematically by Kaj and Krone (2003) (see also Donnelly and Kurtz 1999). In this case, there will be no averaging of the effects of size fluctuations, and hence there is no effective population size. Rather, the size fluctuations on this scale directly affect the time scale of the coalescent in a nonlinear, stochastic manner. In other words, the coalescent in this setting is given by a time change of Kingman’s coalescent, but now the time change is a random process and not a linear function of $t$, which is what would happen in the case of averaging.

To make this precise, consider a single haploid population with size $M_N(\tau)$, $\tau$ generations in the past, and write

$$X_N(\tau) = \frac{M_N(\tau)}{N}$$

for the relative size process, where $N$ is a parameter that we will take to be large. We assume that this process, when run on the coalescent time scale, converges to a process $\{X(t) : t \in [0, \infty]\}$ with state space $I \subseteq (0, \infty)$:

$$X_N([Nt]) = \frac{M_N([Nt])}{N} \to X(t),$$

as $N \to \infty$.

We have in mind, primarily, three kinds of limit processes:

- **Case (i)** (one-dimensional diffusion). $X$ is a diffusion process with state space given by some interval $[a, b]$, where $a > 0$.
- **Case (ii)** (jump process). $X$ is a continuous-time Markov jump process with bounded jump intensities and state space $I$ given by a discrete subset of $(0, \infty)$.
- **Case (iii)** (mixture). We can also consider combinations of the above; i.e., diffusion plus occasional large jumps. This includes as a special case deterministic continuous size change (e.g., exponential decay, reflecting exponential growth forward in time) with occasional random jumps.

Note that Case (i) contains as a trivial special case the usual models of deterministic size fluctuations as discussed, for example, in Griffiths and Tavaré (1994). In other words, the diffusion coefficient is zero in such models.

Intuitively, for the diffusion limit to occur, the scaled size process $X_N(\cdot)$ should make frequent (say every generation) small jumps (of order $1/N$). For example, if $I = [a, b]$ is the state space for the limiting diffusion, the state space for $X_N(\cdot)$ might be of the form $I_N \equiv I \cap Z_N$, where $Z_N = \{0\} \cup \mathbb{Z}$ is the set of all integer multiples of $1/N$.

A typical example of Case (ii) occurs when the process $X_N(\cdot)$ jumps within a fixed discrete set (possibly finite), and the probability of jumping out of a given state in one generation is of order $1/N$. Then, for a given $N$, the holding time in a given state is geometric with parameter $p_N \sim O(1/N)$. These geometric holding times converge to exponential holding times as $N \to \infty$. As we mentioned above, the thing to keep in mind in all these cases is that “macroscopic” changes in population size occur on the same time scale as coalescence events.
We will show in the Appendix that, in the limiting coalescent, the pairwise coalescence probability during \([0,t]\) is determined by the *cumulative coalescence intensity* over the time interval \([0,t]\):

\[
Y_t \equiv \int_0^t \frac{1}{X(s)} \, ds,
\]

where \(X(t)\) is the scaled backward size process. Thus, when there are \(k\) lineages, the coalescence intensity grows like \((k^2)Y_t\). In other words, if \(A(t)\) is Kingman’s coalescent process, the limiting coalescent in the above setting is given by \(A(Y_t)\). This is Kingman’s coalescent run according to the nonlinear stochastic clock \(Y_t\). This can be envisioned as moving up the standard coalescent tree at a rate that varies according to what the current size is. Notice that the initial population size matters, unlike what happens in the case of averaging. This dependence will also be seen in the simulations when size fluctuations are sufficiently slow.

**Simulation results**

We consider a simple model in which the population size has two possible values and falls within the realm of Case (ii) above. This model has been studied by Izuka and co-workers in the context of inbreeding (Iizuka 2001) and heterozygosity (Iizuka *et al*. 2002) effective population sizes. There are four parameters — two population sizes \(N_1\) and \(N_2\) (with \(N_1 < N_2\)), and two transition probabilities, \(q_1\) and \(q_2\), giving the one-step probabilities of size changes from \(N_1\) to \(N_2\) and from \(N_2\) to \(N_1\), respectively. Thus, the size process describing the demographic process is a discrete-time Markov chain with state space \(\{N_1, N_2\}\) and unique stationary distribution \(\pi\) with \(\pi(N_1) = q_2/(q_1 + q_2)\) and \(\pi(N_2) = q_1/(q_1 + q_2)\).

The parameter values used in our simulations were \(N_2 = 10^4\) and \(10^5\), while \(N_1\) was fixed at \(10^3\). For simplicity, we set \(q_1 = q_2\) and values used were 1, 0.75, 0.5, \(10^{-0.5}\), \(10^{-1}\), \(10^{-1.5}\), \ldots, \(10^{-6}\). The mutation probability per individual per generation was fixed at \(\mu = 0.001\). For details about the simulations, see Appendix. Dependence on initial size is one of the hallmarks of the non-averaging case. It is tempting to think that one might obtain averaging, and hence a linear time change in the coalescent, by starting the demographic process at its stationary distribution. A heuristic argument for why this will not work can be made along the following lines. Because \(q_1 = q_2\), we expect the demographic process to spend as much time in state \(N_1\) as in \(N_2\). However, because \(N_1 < N_2\), the coalescence rate is higher while the population size is \(N_1\). The combined effect is that, conditional on a coalescence event happening in generation \(\tau\), the population size of generation \(\tau\) is more likely to be \(N_1\), implying that the distribution of the demographic variable at \(\tau\) is no longer given by the stationary distribution with which we started. When the demographic process is much faster than coalescence events, however, we would expect this effect to be negligible, and the genealogy to behave as in a constant-size null model.

As predicted, departure from the null model are detected only when the fluctuations in population size are intermediate. The range of \(q\) which corresponded to the demographic process being sufficiently “intermediate” to make rejecting the constant-size null model likely appears to be given by the interval \([1/N_2, 1/N_1]\) extended by one order of magnitude on either side, i.e. \([10^{-5}, 10^{-2}]\) for \(N_2 = 10^4\), \(N_1 = 10^3\) and \([10^{-6}, 10^{-2}]\) for \(N_2 = 10^5\), \(N_1 = 10^3\) (figure 1). For \(q\) larger than \(10/N_1\), fluctuations are fast enough to give \(F\) values that are consistent with a null model with an appropriately averaged constant effective size. For \(q\) smaller than \(1/(10N_2)\), the size fluctuations are slow enough to give \(F\) values consistent with population size fixed at the initial value.

Not surprisingly, the extent to which \(F\) deviates from zero increases as the difference between \(N_1\) and \(N_2\) increases. As long as population size fluctuations are small, they have little effect.

When size fluctuations do have an effect on \(F\), the effect increases with sample size. This is expected because, as the sample size increases, so does the time to the most recent common ancestor. However, the phenomenon becomes less marked as the sample size gets very large. This is a consequence of the fact that the expected time to the most recent common ancestor reaches a limit as the sample size goes to infinity.

Figure 2 shows that \(F\) tended to be more negative when the initial population size was the larger one (\(N_2\)). The reason for this is that the coalescence rate is smaller when the population size is \(N_2\) (because \(N_2 > N_1\)). Thus, since \(q_1 = q_2\), a population size change before a coalescence event is more likely when the population size is \(N_2\) than when it is \(N_1\).

**STRUCTURED POPULATIONS**

In this section, we consider the effects of population subdivision on genealogical processes and discuss under which conditions averaging occurs. In other words, we seek conditions under which one can think of the population as being equivalent to a single panmictic unit with some constant effective size. Here, population subdivision might refer to geographical structure with, for example, fixed-sized demes connected by migration. More generally, it refers to any partitioning of the population into different “types” of individuals, with a corresponding “migration” of types. When appropriately scaled, the resulting ancestral process often converges to a “structured coalescent” (Notohara 1990; Herbots 1997) in which lineages within a deme
can coalesce, as in the coalescent, and lineages occasionally migrate between demes.

Our interest here is in finding conditions under which the limiting genealogy can be thought of as a standard coalescent relative to a single effective population size. This was discussed at length by Nordborg and Krone (2002), and we refer to that paper and the references therein for details. We contend ourselves with a brief summary of the main ideas, followed by simulations for a special case to get a feel for when these approximations work in finite populations.

If migration between subpopulations is sufficiently fast compared to the coalescent time scale, the effects of subdivision will be felt in the coalescent only in an average sense. Essentially, the migration process has time to reach equilibrium between coalescence events.

In this case there will be a coalescent effective population size and the genealogy will be given by Kingman’s coalescent with a linear time change. If, on the other hand, migration events are “intermediate” in the sense that they occur on the same time scale as coalescences, then the resulting genealogical process will be described by a structured coalescent. In this case, the genealogy cannot be thought of a standard coalescent and there is no coalescent effective population size.

To make this more precise, let us consider a scenario in which a population is broken up into a finite number, $L$, of demes and that the size of deme $i$ is the constant $N_i = a_iN$ where $a_1 + \cdots + a_L = 1$; hence $N$ is the total population size. Since coalescence probabilities in discrete-time structured models depend on the locations of the lineages in addition to the total number of lineages, the genealogy is a function of the backwards configuration process, $X(\tau) = (X_1(\tau), \ldots, X_L(\tau))$, where $X_i(\tau)$ denotes the number of ancestors in deme $i$, $\tau$ generations into the past. This is a discrete-time Markov chain whose state space consists of vectors $\mathbf{x} = (x_1, \ldots, x_L)$ which specify, at any time in the past, the number of ancestral lineages in each deme. The configuration process evolves by (backwards) migration and coalescing of ancestors with the appropriate probabilities as we move back in time one generation at a time. In general, coalescence probabilities change when the configuration process changes. Let $b_{ij}$ denote the probability that a given lineage “migrates” from deme $i$ to deme $j$ one generation back in time. For example, if the forward migration probabilities are denoted by...
Suppose that lineages migrate (backwards) independently of one another and that the backward migration process determined by the \( b_{ij} \)'s is irreducible and aperiodic with stationary distribution \( \gamma = (\gamma_1, \gamma_2, \ldots, \gamma_L) \). Finally, assume that the above backward migration probabilities scale like \( b_{ij} = \beta_{ij}/N^\alpha, i \neq j \), for some \( 0 \leq \alpha \leq 1 \). Of course, \( b_{ii} = 1 - \sum_{j \neq i} b_{ij} = 1 - N^{-\alpha} \sum_{j \neq i} b_{ij} \). In this case, we will say that \( b_{ij} \) has scaling exponent \( \alpha \).

Migration probabilities with scaling exponent \( \alpha \) correspond to migration events which take, on average, \( O(N^\alpha) \) generations to occur. In particular, when \( 0 \leq \alpha < 1 \), migration events occur much faster than coalescence events (when \( N \) is large). In this case, Nordborg and Krone (2002) show, under mild conditions, that the ancestral process for the discrete-time model can be approximated by a linear time change of the standard coalescent. The coalescence rate when there are \( r \) lineages is given by

\[
\binom{r}{2} \sum_{k=1}^{L} \frac{\gamma_k^2}{a_k}. \]

Thus, there is a scaling constant

\[
c \equiv \sum_{k=1}^{L} \frac{\gamma_k^2}{a_k} \tag{4}
\]

that gives the pairwise coalescence rate, and hence the coalescent effective population size is

\[
N_e = \left( \sum_{k=1}^{L} \frac{\gamma_k^2}{a_k N_k} \right)^{-1} = \left( \sum_{k=1}^{L} \frac{\gamma_k^2}{N_k} \right)^{-1}. \tag{5}
\]

Note that this can also be thought of as a kind of harmonic mean size in which the weighting factor \( \gamma_k^2 \) represents the stationary probability of finding two ancestral lineages together in deme \( k \). Thus, when \( 0 \leq \alpha < 1 \), the structured model can be thought of as a panmictic Wright–Fisher model with population size \( N_e \).

When the scaling exponent is \( \alpha = 1 \), migration events occur on the same time scale as coalescence events and the stochastic nature of migration does not average out in the limit. In this case, the discrete-time ancestral process converges to a structured coalescent. This maintenance of structure in the limiting genealogy results in, for example, higher variance for sample data than that expected under the standard coalescent; thus we do not expect the same pattern of variation as under the null model.

The above discussion holds in much more generality and we refer the reader to Nordborg and Krone (2002) for a full discussion. For example, if the scaling exponents are not all the same, all demes connected by fast migration (i.e., scaling exponents in the interval \([0, 1]\)) collapse down to an effectively panmictic group; all migration corresponding to scaling exponent 1 remains in the suitably-reduced structured coalescent.

### Simulations for structured populations

To emphasize the effects of scaling in subdivided populations, we employed the simplest possible model of geographical structure, with two subpopulations of equal size connected by symmetric migration. Two cases of total population size (twice the common subpopulation size), \( 10^3 \) and \( 10^4 \), were investigated and the scaled migration rate \( (\beta = bN, \text{where } b \text{ is the common migration probability}) \) varied between \( 10^{-1.0} \) and \( 10^{2.5} \), the exponent changing in increments of 0.5. The sample was divided equally between the two demes; i.e., half of the lineages started in one subpopulation and the remaining half in the other. As we have seen, an effective size is expected to exist when the migration rate is sufficiently fast.

As shown in figure 3, \( F \) did not differ much from the value that would be expected under a panmictic null model with scaled migration rate \( 10^1 \) or larger. This was true for both subpopulation sizes. In other words, \( F \) showed the effects of subdivision only for \( b < 10^{-2} \) (resp., \( b < 10^{-3} \)) when \( N = 10^3 \) (resp., \( N = 10^4 \)). Notice that the dependence on sample size is prominent only when the migration rate is very small.

We emphasize that the flat part of the graph corresponding to fast migration is predicted by the theory. The interesting thing about the simulations is that they point out how fast the migration has to be and show the effects of subdivision on \( F \) when migration is not fast enough.

### DISCUSSION

We have shown that when demographic processes and coalescent events operate on similar time scales the coalescent effective size does not exist. In other words, the genealogy cannot be expressed by a linear time scaling of the standard coalescent. As was already pointed out by (Nordborg and Krone 2002), the coalescent effective size is conceptually different from classical notions of effective size in that its existence implies that the properly scaled ancestral process converges to Kingman’s coalescent with a linear time change. This is a strong condition. Phenomena that can be reduced to an effective population size in our sense are not detectable through polymorphism data alone.

We have shown that convergence to the standard coalescent (with a linear time change) is not always obtained when the population size fluctuates randomly or when the
population is subdivided. Whether or not this happens, and hence whether or not there is a coalescent effective size, depends on the relative time scales at which coalescences and demographic processes are operating. This is yet another illustration of the importance of time scales first stressed in (Nordborg 1997) and then encapsulated in Möhle’s theorem (Möhle 1998). In practice, our simulations suggest that the order of magnitude of the demographic processes should be different from that of the inverse of the population size for the standard coalescent approximation to be sufficiently accurate. By “sufficiently accurate” we mean that deviations from the null model are not expected to be detected in genetic data. To monitor this we used Fu and Li’s $F$ and found that, for a simple 2-state population size model, the population size and coalescent processes operate on different time scales when the probability of a state change is not in the range from one order of magnitude less than the inverse of the smaller population size to one order of magnitude more than the inverse of the larger population size. For a simple model of population structure the same is true when the probability of migrating is not higher than the order of magnitude of the inverse of the population size. Thus the results of the two cases are similar: when there is one order of magnitude or more difference between the probability of a demographic change and the probability of a coalescence event, an effective size can be assumed.

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LITERATURE CITED


Consider the fluctuating size model discussed in the main text. Let $A_N(\cdot)$ be the ancestral process defined by $A_N(0) = n$ and $A_N(\tau) =$ number of distinct ancestors $\tau$ generations in the past, $\tau \geq 1$, where $n$ is the original sample size. To find the coalescence rate for a pair of lineages, we compute the probability (conditional on population size) that no coalescence has occurred over $[Nt]$ generations and take a limit:

$$
\mathbb{P} (\text{no coalescence in } [Nt] \text{ generations}| \{M_N(\cdot)\})
= \prod_{\tau=1}^{\lfloor Nt \rfloor} \left(1 - \frac{1}{M_N(\tau)}\right) = \prod_{\tau=1}^{\lfloor Nt \rfloor} \left(1 - \frac{1}{N X_N(\tau)}\right) \quad (A1)
$$

$$
\sim \exp\left(-\frac{1}{N} \sum_{\tau=1}^{\lfloor Nt \rfloor} \frac{1}{X_N(\tau)}\right)
\rightarrow \exp\left(-\int_0^t \frac{1}{X(s)} \, ds\right), \quad (A2)
$$

as $N \to \infty$. This suggests that, in the limiting coalescent, the coalescence probability for a pair of lineages during $[0, t]$ is governed by an exponential random variable with rate given by $\int_0^t \frac{1}{X(s)} \, ds$. Thus, as in Griffiths and Tavaré (1994) for deterministic size fluctuations and Kaj and Krone (2003) in the case of stochastic fluctuations, we define the cumulative coalescence intensity over the time interval $[0, t]$ by

$$
Y_t = \int_0^t \frac{1}{X(s)} \, ds.
$$

This applies to any pair of ancestral lineages, so when there are $k$ lineages, the coalescence intensity grows like $\left(\frac{k}{2}\right) Y_t$. In other words, the limiting coalescent process should be given by

$$
\lim_{N \to \infty} A_N([Nt]) = A(Y_t), \quad (A3)
$$

where $A(t)$ is Kingman’s coalescent, and $Y_t$ is the increasing, nonlinear stochastic time change. In a more general context, this result was proven in Kaj and Krone (2003) in terms of weak convergence for the bivariate process $\{(N^{-1}X_N([Nt]), A_N([Nt]))\}_{t \geq 0}$ towards $\{(X(t), A(Y_t))\}_{t \geq 0}$, as $N \to \infty$. In agreement with the result quoted above. Here we recognize the regime of slow demographic change by taking $a$ very small. Indeed, as $a \to 0$,

$$
\frac{1}{a} Y_{at} = \frac{1}{a} \int_0^{at} \frac{1}{X(s)} \, ds \approx t \cdot \frac{1}{X(0)}.
$$

### Relative speed of fluctuations

Following Sano et al. (2004), we give another perspective on the three time-scaling regimes for the varying size model, now starting from the limit result for the intermediate case and considering two extremes. To this end, let $a$ be a dummy variable signifying a change in speed of the pre-limit process from $X_N([Nt])$ to $X_N([aNt])$. Then $\{(N^{-1}X_N([aNt]), A_N([Nt]))\}_{t \geq 0}$ tends towards $\{(X(at), A(a^{-1}Y_{at}))\}_{t \geq 0}$ as $N \to \infty$, in agreement with the result quoted above. Here we recognize the regime of slow demographic change by taking $a$ very small. Indeed, as $a \to 0$,

$$
\frac{1}{a} Y_{at} = \frac{1}{a} \int_0^{at} \frac{1}{X(s)} \, ds \approx t \cdot \frac{1}{X(0)}.
$$

### APPENDIX

#### Nonlinear time change for intermediate size fluctuations

Consider the fluctuating size model discussed in the main text. Let $A_N(\cdot)$ be the ancestral process defined by $A_N(0) = n$ and $A_N(\tau) =$ number of distinct ancestors $\tau$ generations in the past, $\tau \geq 1$, where $n$ is the original sample size.

To find the coalescence rate for a pair of lineages, we compute the probability (conditional on population size) that no coalescence has occurred over $[Nt]$ generations and take a limit:

$$
\mathbb{P} (\text{no coalescence in } [Nt] \text{ generations}| \{M_N(\cdot)\})
= \prod_{\tau=1}^{\lfloor Nt \rfloor} \left(1 - \frac{1}{M_N(\tau)}\right) = \prod_{\tau=1}^{\lfloor Nt \rfloor} \left(1 - \frac{1}{N X_N(\tau)}\right) \quad (A1)
$$

$$
\sim \exp\left(-\frac{1}{N} \sum_{\tau=1}^{\lfloor Nt \rfloor} \frac{1}{X_N(\tau)}\right)
\rightarrow \exp\left(-\int_0^t \frac{1}{X(s)} \, ds\right), \quad (A2)
$$

as $N \to \infty$. This suggests that, in the limiting coalescent, the coalescence probability for a pair of lineages during $[0, t]$ is governed by an exponential random variable with rate given by $\int_0^t \frac{1}{X(s)} \, ds$. Thus, as in Griffiths and Tavaré (1994) for deterministic size fluctuations and Kaj and Krone (2003) in the case of stochastic fluctuations, we define the cumulative coalescence intensity over the time interval $[0, t]$ by

$$
Y_t = \int_0^t \frac{1}{X(s)} \, ds.
$$

This applies to any pair of ancestral lineages, so when there are $k$ lineages, the coalescence intensity grows like $\left(\frac{k}{2}\right) Y_t$. In other words, the limiting coalescent process should be given by

$$
\lim_{N \to \infty} A_N([Nt]) = A(Y_t), \quad (A3)
$$

where $A(t)$ is Kingman’s coalescent, and $Y_t$ is the increasing, nonlinear stochastic time change. In a more general context, this result was proven in Kaj and Krone (2003) in terms of weak convergence for the bivariate process $\{(N^{-1}X_N([Nt]), A_N([Nt]))\}_{t \geq 0}$ towards $\{(X(t), A(Y_t))\}_{t \geq 0}$, as $N \to \infty$. In agreement with the result quoted above. Here we recognize the regime of slow demographic change by taking $a$ very small. Indeed, as $a \to 0$,

$$
\frac{1}{a} Y_{at} = \frac{1}{a} \int_0^{at} \frac{1}{X(s)} \, ds \approx t \cdot \frac{1}{X(0)}.
so we have linear scaling with effective population size $X(0)$; i.e., on the coalescent time scale, the population size never changes. Similarly, $a$ very large corresponds to the fast scaling regime. In this case, assuming that $X(t)$ is ergodic with a steady state $X_\infty$, the limit $a \to \infty$ gives
\[
\frac{1}{a} Y_{at} = t \frac{1}{at} \int_0^{at} \frac{1}{X(s)} \, ds \approx t \cdot E(1/X_\infty),
\]
which is linear scaling with effective population size given by the harmonic mean $1/E(1/X_\infty)$.

Recall that $X(s)$ represents the (limiting) scaled size process. As is to be expected, the coalescence intensity $Y_t$ increases at a faster rate when $X(t)$ is smaller. Also, during such periods in which $Y_t$ increases faster, there will tend to be fewer mutations. This was brought out in earlier calculations, as well as in the simulations.

The program

The simulation program, written in C++, can be obtained from the authors upon request. The program simulates the combined effect on $F$ of the ancestral and demographic processes. The models used for the demographic processes were the simplest possible. Fluctuating population size was modeled by a two-state Markov chain and population structure by two equally sized subpopulations with symmetric migration.

Generations were discrete and population sizes finite. For each set of parameters, even sample sizes from 4 to 60 were simulated and the average value of $F$ from ten thousand runs was calculated. The program does not allow for more than two lineages to coalesce at a time. This deviation from the Wright–Fisher model will cause a negative bias in $F$, but the effect is negligible for the parameter values we used (results not shown: the sample size has to be large relative to the population size for multiple coalescences to the same individual to matter).

Every generation, a gene can mutate with a fixed probability. If no mutations occur in a given realization, $F$ is not defined. One can either define $F$ as zero or disregard these cases. We chose a mutation rate high enough so that our results were unaffected by which method was used.