Expectation of the Site Frequency Spectrum

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Abstract

The site frequency spectrum describes variation among a set of n DNA sequences. Its i’th entry (i = 1, 2, ..., n−1) is the number of nucleotide sites at which the mutant allele is present in i copies. Under selective neutrality, random mating, and constant population size, the expected value of the spectrum is well known but somewhat puzzling. Each additional sequence added to a sample adds an entry to the end of the expected spectrum but does not affect existing entries. This note reviews the reasons for this behavior.

In a sample of n DNA sequences, a polymorphic nucleotide site can divide the sample into 1 mutant and n − 1 non-mutants, into 2 mutants and n − 2 non-mutants, and so on. The number of copies of the mutant must be at least 1 and no more than n − 1 if the site is polymorphic. The site frequency spectrum describes the number of sites that fall into each of these n − 1 categories. It is widely used as a summary of variation among DNA sequences.

The expectation of the spectrum has a very simple form under random mating, selective neutrality, and constant population size, provided that the mutation rate is so low that we can ignore the possibility of multiple mutations at the same site—the so-called “infinite sites” model of mutation [4]. The expected spectrum for samples of sizes 2 through 5 looks like this:

| Sample size | Expected spectrum (singletons, doubletons, ...) |
|-------------|-----------------------------------------------|
| 2           | θ                                              |
| 3           | θ, θ/2                                         |
| 4           | θ, θ/2, θ/3                                   |
| 5           | θ, θ/2, θ/3, θ/4                              |
| Etcetera    |                                               |

Here, “singletons” are sites with one copy of the mutant allele, “doubletons” are sites with two copies, and so on. θ = 4Nu, where N is population size and u the mutation rate per sequence per generation. The expected number of sites with i copies of the mutant allele is θ/i [1, Eqn. 22]. It is remarkable that as we increase sample size, the number of mutants in each category doesn’t change. We merely add a new category at the end. To explore the cause of this behavior, we begin with a graphical argument.

Figure 1 shows the gene genealogy of a sample of size 3, with branch lengths equal to their expected values [2]. There are two coalescent intervals: a recent one on the left, with three lines of descent, and an ancient one on the right, with two. Imagine first
that we could examine this sample at the end of the ancient interval, just before the upper lineage bifurcates. Our tree would have two branches, and the expected length of each is $2N$ generations. Thus, we expect $2N u = \theta/2$ mutations on each branch, and $\theta$ mutations altogether, all of which are singletons.

At the time of the coalescent event, half of these singletons become doubletons, and this is why we end up with $\theta/2$ doubletons at the left edge of the graph. There are also new singletons, which arise via mutation during the recent interval. How many? The expected length of this interval is $2N/3$ generations, and there are 3 lines of descent, so the total branch length within the interval is $2N$. Consequently, the recent interval will add $\theta/2$ to the expected number of singletons—exactly the number that we lost when the upper lineage bifurcated. At the end of the second interval, we expect $\theta$ singletons and $\theta/2$ doubletons, just as in the table above.

To carry this argument farther, we must make it algebraic, and our algebraic argument paraphrases that of Hudson [3]. As in the graphical argument, we census mutations at the recent (tipward) end of each coalescent interval. For the $k$th interval (the one with $k$ lineages), let $s_{i,k}$ denote the expected number of polymorphic sites at which the derived allele is present in $i$ copies. There can be no polymorphic sites if there is only one lineage, so $s_{i,1} = 0$ for all $i$. Singletons require special treatment. Their expected number obeys

$$s_{1,k} = \frac{\theta}{k-1} + \left(1 - \frac{1}{k-1}\right)s_{1,k-1} \quad (1)$$

The first term on the right accounts for singleton mutations that arose within the current ($k$th) interval. This is simply the number of new mutations, a standard result from coalescent theory. The second term is the contribution from the $s_{1,k-1}$ singletons that existed during the $(k-1)$th interval. Of these, a fraction $1 - 1/(k-1)$ remain singletons because they lie on a lineage that did not bifurcate.

Consider now those mutations with $i > 1$ copies at the end of interval $k$. These can arise in two ways: either the mutation was present in $i - 1$ copies at the end of interval $k-1$, and one of the copies bifurcated, or else it was present in $i$ copies, none of which bifurcated. We consider these cases separately.

**Mutations that increased in number.** During the interval with $k-1$ lineages, $s_{i-1,k-1}$ is the expected number of mutations present in $i - 1$ copies. At the end of this interval, a single random lineage bifurcates. Each lineage has the same chance, $1/(k-1)$, of being the one that bifurcates, so a mutation with $i - 1$ copies has probability $(i - 1)/(k-1)$ of including the lineage that bifurcates. In this

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1To derive the expression, note that the expected duration of the interval is $4N/[k(k-1)]$. Multiply by $k$ to get the total branch length and then by $u$ to get the expected number of mutations.
case, its number will increase from \( i - 1 \) to \( i \).

Thus, \( \frac{i-1}{k-1}s_{i-1,k-1} \) is the contribution to \( s_{i,k} \) from mutations that increased in number.

**Mutations that did not increase.** During interval \( k - 1 \), \( s_{i,k-1} \) is the expected number of mutations present in \( i \) copies. With probability \( 1 - i/(k - 1) \), none of these were on the lineage that bifurcated. Thus, \( (1 - i/(k - 1))s_{i,k-1} \) is the contribution to \( s_{i,k} \) from mutations that did not increase.

Summing these contributions,

\[
s_{i,k} = \frac{i-1}{k-1}s_{i-1,k-1} + \left(1 - \frac{i}{k-1}\right)s_{i,k-1} \tag{2}
\]

for the case in which \( i > 1 \).

Consider the sequence \( s_{1,2}, s_{2,3}, s_{3,4}, \) and so on. This is the case of a “lucky” mutation whose count grows as fast as possible, because one of its copies happens to bifurcate at the end of each coalescent interval. It is of interest because it simplifies our equations. With \( k = i + 1 \), the second term in Eqns. 1 and 2 each disappear, leaving

\[
\begin{align*}
s_{1,2} &= \frac{i-1}{i}s_{i-1,i} \\
s_{i,i+1} &= \frac{i-1}{i}s_{i-1,i} \quad \text{if } i > 1
\end{align*}
\]

If you work your way through this sequence, you will discover that \( s_{i,i+1} = \theta/i \) for all \( i \).

Now we are in a position to discover why the entries of the expected spectrum do not change as we add DNA sequences to the sample—or in other words, why \( s_{i,k} \) does not depend on \( k \). Its value is given by the formula derived above for \( k = i + 1 \):

\[
s_{i,k} = \theta/i \tag{3}
\]

for all \( i \) and for all \( k > i \), as shown by Fu [1, Eqn. 22].

**References**

[1] Yunxin Fu. Statistical properties of segregating sites. *Theoretical Population Biology*, 48(2):172–197, October 1995.

[2] Richard R. Hudson. Gene genealogies and the coalescent process. In Douglas Futuyma and Janis Antonovics, editors, *Oxford Surveys in Evolutionary Biology*, volume 7, pages 1–44. Oxford University Press, Oxford, 1990.

[3] Richard R. Hudson. A new proof of the expected frequency spectrum under the standard neutral model. *PLoS ONE*, 10(1):e0118087, 2015.

[4] Motoo Kimura. The number of heterozygous nucleotide sites maintained in a finite population due to steady flux of mutation. *Genetics*, 61:893–903, 1969.