Multiplicative versus additive selection in relation to genome evolution: a simulation study

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Summary
The evolution of molecular quantitative traits, such as codon usage bias or base frequencies, can be explained as the result of mutational biases alone, or as the result of mutation and selection. Whereas mutation models can be investigated easily, realistic modelling of selection-directed genome evolution is analytically intractable, and numerical calculations require substantial computer resources. We investigated the evolution of optimal codon frequency under additive and multiplicative effects of selected linked codons. We show that additive selective effects of many linked sites cannot be effective in genomes when the number of selected sites is greater than the effective population size, a realistic assumption according to current molecular data. We then discuss the implications of these results for isochore evolution in vertebrates.

1. Introduction
The efficacy of selection acting simultaneously at linked sites (a site can be a codon, a nucleotide or a gene, depending on what contributes to fitness) is reduced compared with the same selection pressure acting at independent loci (Hill & Robertson, 1966; Li, 1987). This is because linkage disequilibrium between alleles at selected loci, generated by the stochastic nature of mutation and sampling in a finite population, ‘interferes’ with the action of selection at any one locus (Felsenstein, 1974). Recent work on the effect of linkage on the efficiency of selection has stressed the fact that Hill–Robertson (HR) interference plays a major role in the effect of selection on codon usage bias and the pattern of silent polymorphism within the genome (Comeron et al., 1999; McVean & Charlesworth, 2000). The efficiency of selection can be predicted assuming independence between sites (Li, 1987; McVean & Charlesworth, 1999). Let \( v \) be the mutation rate from non-optimal codons and \( u \) the reverse mutation rate. Let \( x \) be the optimal codon frequency, \( s \) be the selection coefficient for optimal codons, and \( N \) the effective size of the diploid population. Then, at mutation–selection–drift equilibrium:

\[
E(x) = \frac{\sum_{n=0}^{\infty} (4Ns)^n \Gamma(4Nu+n+1) / \sum_{n=0}^{\infty} n! \Gamma(4Nu+n)}{n! \Gamma(4N(u+v)+n+1) / \sum_{n=0}^{\infty} n! \Gamma(4N(u+v)+n)}
\]

where \( \Gamma(.) \) denotes the Gamma function (Li, 1987). At mutation–selection–drift equilibrium, the optimal codon frequency is determined by \( Ns, N(u+v) \) (the mean mutation rate for one codon per generation), and \( u/(u+v) \) the average frequency of optimal codons without selection. There are two difficulties when several sites participate in a trait (e.g. the optimal codon frequencies of genes or of a genome). The first one is the reduced selection efficiency, e.g. codon bias, due to HR interference between the selected sites (Comeron et al., 1999; McVean & Charlesworth, 2000). The second difficulty is the way to define the contribution of each codon to the fitness of a genome.

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2. Definition of the additive and multiplicative selection schemes

We limit the present analysis to two previously published selection schemes. From Li’s (1987) definition, in the case of $L$ diallelic loci, the absolute fitness of a genome consisting of $L$ codons with $i$ optimal codons is equal to $F_i = 1 - (L - i)s'$ for negative selection, and $F_i = 1 + is$ for positive selection when additivity over codons is assumed. The positive and negative additive selection models are mathematically equivalent with $s' = s/(1 + Ls)$. This arises from the definition of the fitness is nearly equivalent when the product $Ls < 1$ under this definition. This can be deduced from the expression of the fitness of a sequence with $i + j$ optimal codons ($F_{i+j}$) over the ratio of a sequence with $i$ optimal codons ($F_i$) under both schemes (Table 1). The absolute fitness is $F_i = (1 - s')^{L-i}$ or $F_i = (1 + s)^i$. The positive and negative multiplicative selection models are mathematically equivalent if $s' = s/(1 + s).$ The additive and the multiplicative selection schemes are nearly equivalent when the product $Ls < 1$, since $(1 + s)^i \approx 1 + is$ (Li, 1987).

For a given $s'$, negative additive selection against non-optimal codons is more effective on codon usage bias than multiplicative selection (McVean & Charlesworth, 2000). This arises from the definition of the additive and the negative selection scheme (Table 1). In the case of positive selection, the ratio $F_{i+j}/F_i$ is greater under the multiplicative selection scheme than under the additive selection scheme, so that one expects multiplicative selection to be more effective than additive selection. In the case of negative selection, the contrary is true: multiplicative effects are less effective than additive effects (Table 1). The negative additive selection scheme is restricted to the parameter space $Ls < 1$, so that we have used the positive selection scheme. The total number of synonymous sites is estimated to be greater than the effective population size, $N$, in many species (Kondrashov, 1995). For example, the total number of codons in Drosophila is about $L \approx 7 \times 10^6$ (total number of genes: 13600 (Adams et al., 2000), average protein length 500 amino acids) and the effective population size is about $N \approx 10^8$ (Sawyer & Hartl, 1992); in the worm Caenorhabditis elegans $L \approx 7 \times 10^6$, and in the plant Arabidopsis thaliana $L \approx 10^7$. Estimates of effective population sizes are lacking in the latter species. Selection on codon usage has been demonstrated in Drosophila (Shields et al., 1998), Caenorhabditis and Arabidopsis (Stenico et al., 1994; Duret & Mouchiroud, 1999). As a consequence, since selection is effective when $Ns \approx 1$ (Kimura, 1983) it is reasonable to study the case $Ls > 1$ (where $L$ is the total number of codons in a genome).

3. Results

We explored the efficiency of selection under positive directional selection on optimal codons under both selection schemes, using Li’s (1987) multisite model. The equilibrium frequency of the optimal codon $x_{eq}$ was computed 100 times every $2N$ generations to ensure independence between values (Li, 1987), allowing $10(u+v)$ generations to reach equilibrium (Tachida, 2000). The corresponding value predicted from the single-site theory was calculated from each run (for this haploid model, $4N$ is replaced by $2N$ in equation 1). We used $N(u+v) = 0.05$ and $0.005$, $N = 10^2$, $10^3$ and $10^4$, $L$ from 32 to 38400 and $Ns = 0.1, 0.4, 1, 4$ and 10. The number of recombination events per generation was assumed to follow a Poisson process with parameter $cNL$. Each simulation was run at least four times. Fig. 1 shows that, for positive selection over sites, additive effects of optimal codons are less effective than multiplicative effects. An increase in $Ls$, $L$ constant, equivalent to a decrease in $N$ keeping $Ns$ and $N(u+v)$ constant, dramatically reduces the efficacy of additive selection over sites (Fig. 1A, B). Thus, the selection–mutation–drift equilibrium optimal codon frequency depends on $N$ and $L$ with additive effects of selected sites.

A methodological consequence is that the calculation times (which increase with increasing population size) cannot be reduced by keeping $N(u+v)$ and $Ns$ constant under additive effects when $Ls > 1$. This restriction has not been noted in the previous simulation studies comparing additive and multi-

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<tr>
<th>Fitness</th>
<th>Positive selection</th>
<th>Negative selection</th>
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<tr>
<td>$F_{i+j}$</td>
<td>$(1+s)^{i+j}$</td>
<td>$1 + is + js$</td>
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<tr>
<td>$F_i$</td>
<td>$(1 + s)^i$</td>
<td>$1 + is$</td>
</tr>
<tr>
<td>$F_{i+j}/F_i$</td>
<td>$(1 + s)^{i+j}$</td>
<td>$1 + s(1 + is)$</td>
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Table 1. Comparison of multiplicative versus additive selection under positive and negative selection. $F_i$ is the absolute fitness of a sequence of $L$ sites with $i$ optimal sites.
Multiplicative versus additive selection

Fig. 1. Evolution of the optimal codon frequency $x_{eq}$ in a sequence of 384 completely linked codons for $N(u+v) = 0.05$ (A) and $N(u+v) = 0.0005$ (B), under mutational bias, $u/(u+v) = 30\%$, increasing selection and different population sizes. Bold lines, $(\pm)$ prediction from the single-site theory (equation (1), see text); dashed lines, multiplicative effects (open circles, $N = 10^3$; filled circles, $N = 10^6$); continuous lines, additive effects (filled squares, $N = 10^4$; open circles, $N = 10^6$; filled circles, $N = 10^9$).

Multiplicative selection schemes (Li, 1987; McVean & Charlesworth, 2000). When effects of optimal codons are multiplicative, the relevant parameters are $N(u+v)$, $Ns$ and $L(u+v)$ (Tachida, 2000; McVean & Charlesworth, 2000). The effect of HR interference increases with $L(u+v)$, the mean number of mutations per genome per generation (Fig. 2). HR interference becomes negligible, that is $x_{eq}$ is well predicted from the single-site theory, for small $L(u+v)$ under multiplicative or additive codon effects when $Ls < 1$ (Fig. 1B; Fig. 2 with $L = 32$).

Introduction of recombination between sites reduces the effects of HR interference under the multiplicative fitness scheme (Comeron et al., 1999; McVean & Charlesworth, 2000). The effect of recombination at the rate of one crossover per sequence per generation is sufficient to remove all the effects of interference (Fig. 2B, D), except for high values of $L(u+v)$, when $(u+v) \approx c$ (Fig. 2B; $L = 38400$, $(u+v) = 5 \times 10^{-3}$, $c = 3 \times 10^{-3}$). HR interference on genomes depends on the $(u+v)/c$ ratio: when the recombination rate per genome $Lc$ is not sufficient to make the $L(u+v)$ segregating sites independent $(u+v \approx c$ was not sufficient, from our simulations), there is interference and reduction of selection efficiency. When $c$ is greater than $(u+v)$, the segregating sites are independent and there is no interference: the frequency of optimal codon is well predicted from the single-site theory (Fig. 2B, D). For the additive selection scheme, the effect of recombination is not sufficient to predict $x_{eq}$ from the single-site model (Fig. 2A, C), and recombination does not increase the efficiency of selection at all for $L = 38400$. More exactly, recombination cannot increase the efficacy of selection for $Ls > 1$, as a simple consequence of the definition of the additive fitness: the relative fitness of a sequence with $i+1$ optimal sites appearing from mutation in a haploid population of size $N$ compared with the relative fitness of a sequence containing $i$ optimal sites is equal to the ratio of the absolute fitness $F_{i+1}/F_i$:

$$\frac{F_{i+1}}{F_i} = 1 + \frac{s}{1+is}.$$ 

It follows that for a given selection coefficient over site and $L$ very large, so that the number of optimal sites $i$ (at least equal to $uL/(u+v)$) is also very large, the difference between the fitness of a sequence with $i+1$ optimal sites and the fitness of a sequence with $i$
optimal sites is negligible, as $F_{i+1}/F_{i} \approx 1$. More generally, selection can not distinguish between genomes containing $i$ optimal sites and genomes containing $i+j$ optimal sites when the number of selected sites is very large.

4. Discussion

In vertebrate genomes, codon usage bias is related to the local G+C content of the genome, that is the isochore localization of the gene (Bernardi et al., 1985). The debate simmers on whether the isochore structure (for review see Bernardi, 2000) of the vertebrate genome is due to selection (Bernardi et al., 1985; Eyre Walker, 1999) or to neutral mechanisms (Wolfe et al., 1989) Gu & Li, 1994). If selection for GC nucleotides is involved, $Ls$ is far beyond 1 (the number of nucleotides in the GC-rich isochores is roughly $10^9$ and the long-term effective population size $10^1$ to $10^2$; Nachman & Crowell, 2000), so that additive effects of selection on GC nucleotides cannot be considered to be responsible for the emergence of the isochore structure in vertebrates. The molecular significance of additive or multiplicative effects of selected GC nucleotides remains an open question. Increased thermostability of GC-rich isochores has been proposed to be advantageous for warm-blooded vertebrates (Bernardi et al., 1985; but see Hughes et al., 1999; Rodrigues-Trelles et al., 2000). Biochemical studies suggest that DNA thermostability increases linearly with GC content (Marmur & Doty, 1959), whereas conformational properties of DNA would rather imply non-linear effects (the effect of replacing one AT by one GC depends on the amount of GC present in the sequence) (Foloppe & MacKerell, 1999). The estimated mutation rate in the human genome, of about $10^{-8}$ mutations per nucleotide per generation (Nachman & Crowell, 2000), is of the same order of magnitude as the estimated overall recombination rate in the human genome, $10^{-8}$ recombinations per nucleotide per generation (Weissenbach et al., 1992), although recombination rates vary greatly along the genome (Lynn et al., 2000). This suggests that HR interference may be an important process if selection plays a role in the evolution of the isochore structure.

Our analysis suggests that additive selection on codon usage over the whole genome of most eukaryotic genomes (for which the number of codons is greater than the effective population size) cannot account for codon usage bias. But, as codon usage is highly biased only in the more strongly expressed proteins (e.g. Duret & Mouchiroud, 2000), the number of selected sites in a genome must be smaller than the total number of codons, so that additive directional selection may be effective in species with large effective
population sizes, such as Drosophila. However, we are aware that the definition of the fitness of the genome as an additive or multiplicative increasing and symmetrical function of the codon content is not satisfactory, although it is the simplest way to study HR interference. Many biochemical models of the effect of codon usage on protein translation rate or translation efficiency have been developed but do not take into account the effect of linkage between genes and the resulting expected interference (Bulmer, 1991; Zhang et al., 1994; Xia, 1996). We can therefore expect more realistic models of codon usage to be developed, taking into account both optimal codon usage and linkage between the genome sequences.

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References


