Selection for minimization of translational frameshifting errors as a factor in the evolution of codon usage

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Received May 6, 2009; Revised July 25, 2009; Accepted August 13, 2009

ABSTRACT
In a wide range of genomes, it was observed that the usage of synonymous codons is biased toward specific codons and codon patterns. Factors that are implicated in the selection for codon usage include facilitation of fast and accurate translation. There are two types of translational errors: missense errors and processivity errors. There is considerable evidence in support of the hypothesis that codon usage is optimized to minimize missense errors. In contrast, little is known about the relationship between codon usage and frameshifting errors, an important form of processivity errors, which appear to occur at frequencies comparable to the frequencies of missense errors. Based on the recently proposed pause-and-slip model of frameshifting, we developed Frameshifting Robustness Score (FRS). We used this measure to test if the pattern of codon usage indicates optimization against frameshifting errors. We found that the FRS values of protein-coding sequences from four analyzed genomes (the bacteria Bacillus subtilis and Escherichia coli, and the yeasts Saccharomyces cerevisiae and Schizosaccharomyces pombe) were typically higher than expected by chance. Other properties of FRS patterns observed in B. subtilis, S. cerevisiae and S. pombe, such as the tendency of FRS to increase from the 5’- to 3’-end of protein-coding sequences, were also consistent with the hypothesis of optimization against frameshifting errors in translation. For E. coli, the results of different tests were less consistent, suggestive of a much weaker optimization, if any. Collectively, the results fit the concept of selection against mistranslation-induced protein misfolding being one of the factors shaping the evolution of both coding and non-coding sequences.

INTRODUCTION
It is well established that synonymous codons are used in protein-coding sequences with unequal frequencies, and in each of major model organisms, including Escherichia coli, Saccharomyces cerevisiae, Caenorhabditis elegans and Drosophila melanogaster, a subset of preferred codons was identified (1). Such codon usage bias is typically calculated using Codon Adaptation Index (CAI), which measures how frequently codons preferred by highly expressed genes are used in the given gene (2). Many factors have been reported to affect the codon usage bias, including G + C content (3), compositional strand bias (4), protein secondary structure (5), mRNA secondary structure (6), mRNA half-life (7) and transcription-induced asymmetries (8). However, two of the more important factors linked to the biased codon usage appear to be translational speed and accuracy.

There are several lines of evidence that link codon bias to translation speed. For example, it has been demonstrated that in bacterial genomes, codon bias affects translation elongation rate (9,10), i.e. gene sequences with a higher codon usage bias tend to be translated faster, probably due to the abundant supply of the cognate tRNAs for the frequent codons. These observations are consistent with the translational efficiency hypothesis.

Closely related to translational speed is translational accuracy. Factors that are responsible for slowing the translation process, such as insufficient amounts of cognate tRNAs, have been also linked to missense errors, where one amino acid substitutes for another (11). Indeed, it has been observed that codon usage bias is more pronounced in regions of greater amino-acid conservation (12–15). Again, the relationship between codon bias and minimization of the translation error rate is complex as demonstrated by the finding that, when...
factors like tRNA abundance, translation efficiency, anticodon modification, etc., are disregarded and the fitness for translation accuracy is measured by summing over the probabilities of mutations of individual codons in mRNA weighted by the similarity between the original and the mutated amino acids, then, by this simple measure, the preferred codons have not been selected optimally (16). There is also increasing evidence that the prevention of mistranslation-induced misfolding is an important force shaping codon usage (17).

Unlike missense errors, processivity errors caused by premature termination or shifted reading frames in elongation (frameshifting) cannot be tolerated even in less conserved (and thus presumably functionally less important) regions. These errors are more devastating, leading to waste of energy to generate and then degrade non-functional peptide chains. Moreover, a substantial fraction of frameshifting errors would generate misfolded proteins that could exert toxic effects on the cell (17). The frequency of frameshifting errors was estimated to be \( \sim 10^{-5} \) per codon (18), that is, similar to the frequency of missense errors, for which an upper bound estimate of \( 4 \times 10^{-4} \) per codon has been reported (19).

Several studies have tried to link codon usage gradient to optimization against processivity errors. Recently, Eyre-Walker (20) argued that, if there is selection against processivity errors, codon bias would be expected to increase along the gene sequence from 5' to 3'. Qin et al. (21) measured codon bias using effective number of codons, along the direction of translation in four prokaryotic and two eukaryotic genomes, and showed that codon bias indeed increases along the direction of translation in yeast and some prokaryotes. A similar trend was reported by Stoletzki and Eyre-Walker (13) in an analysis of codon bias in E. coli genes. However, the trend of increasing codon bias along the gene sequence was rather weak in each of these cases. For example, in the gene set selected in their study, the fraction of optimal codons only increased by 0.051 every 1000 bp (13). Stoletzki and Eyre-Walker attributed the gradient to the higher cost of a processivity error when it occurs towards the end of the translation process rather than at the beginning. A gradient of nucleotide and codon usage in E. coli genome was also observed by Hooper and Berg (22).

The second line of supporting evidence for the reduction of processivity errors via codon usage was also proposed by Stoletzki and Eyre-Walker (13) who observed higher codon bias in longer genes. They reasoned that this trend was unlikely to be an artifact of correlations with expression because gene length is negatively correlated with expression and concluded that the most natural explanation for the observed bias is selection against processivity errors. However, the exact relationship between codon usage bias and gene length has been controversial with some studies suggesting a negative correlation (12,23). Eyre-Walker (20) found that codon usage is positively correlated with gene length in E. coli. Moriyama and Powell (24) reported a positive correlation between codon usage bias and gene length in E. coli but negative correlations in S. cerevisiae and D. melanogaster genes. The finding for S. cerevisiae has in turn been contradicted by Coghlan and Wolfe (25), who found a positive correlation between codon usage and gene length in this organism. These conflicting results serve as an illustration of how difficult it is to discover and confidently measure subtle dependencies between codon usage bias and other biological factors. Furthermore, although the increase of codon bias with the protein length might be suggestive of possible optimization against processivity errors, such increased codon bias, in and by itself, does not provide sufficient evidence for optimization against frameshifting error. For example, it has been observed that longer proteins are more highly conserved during evolution (26), leaving possibilities of alternative explanations.

In this work, we develop a more direct argument in support of the hypothesis that codon usage is optimized to prevent processivity errors, specifically, frameshifting errors. The mechanism of frameshifting errors is not completely understood but work on programmed frameshifting (27,28) suggests that programmed frameshifting and frameshifting errors might be mechanistically related (29), and several models have been proposed to explain these mechanisms (29–31). Although the models differ in details, they generally assume that the two essential elements for frameshifting are (i) translational pause and (ii) tRNA slippage. Hence we refer to this concept as the pause-and-slip model. More specifically, we based our approach on the model of Farabaugh and Bjork (29), which incorporates all essential elements of the pause-and-slip concept and remains relatively simple.

We used the pause-and-slip model to design the Frameshifting Robustness Score (FRS) that estimates how resistant a sequence is to frameshifting errors during translation. Given such direct measurement, we compared values of FRS for real sequences and a set of control sequences generated from real sequences by random permutations of synonymous codons. We studied general trends of FRS in genomes of model organisms and report evidence that codon usage is optimized to reduce frameshifting errors in B. subtilis, S. cerevisiae and S. pombe whereas the support for the hypothesis for E. coli was weak.

MATERIALS AND METHODS

Data

The gene-coding sequences of B. subtilis were obtained from the Pasteur Institute (http://genolist.pasteur.fr/SubtiList/). The gene-coding sequences of S. pombe were obtained from the Sanger Institute (http://www.sanger.ac.uk/Projects/S_pombe/). The gene-coding sequences of S. cerevisiae were obtained from Man and Pilpel’s compilation (32). The E. coli K12 genome sequence was downloaded from EcoGene (http://ecogene.org) and then the gene coding sequences were extracted based on the annotation available at EcoGene.

Codons at the start and at the end of E. coli gene sequences have been reported to be under different selection pressure as compared to other codons. We repeated all our computation on E. coli after first removing the first 50 codons and the last 20 codons of each gene sequence,
following the practice of Stoletzki and Eyre-Walker (13). Unless otherwise stated, the conclusions were the same as those reached without removing codons.

Within the gene sequence set of each organism, we first identified two subsets of equal size with low and high CAI values. We also selected four groups of gene sequences based on length and CAI, covering four combinations: short sequences with low CAI, short sequences with high CAI, long sequences with low CAI and long sequences with high CAI. We sorted gene sequences of each organism by their CAI values computed using Sharp and Li’s method (2). The codon coefficients, required for computing CAI, of \( E. coli \) and \( S. cerevisiae \) were taken from the same reference (2). The codon coefficients of \( B. subtilis \) and \( S. pombe \) were taken from Sharp et al.’s work (33). Because of the precision used in the paper (33), some codon coefficients were 0.00, which would make the work (33). Because of the precision used in the paper (33), some codon coefficients were 0.00, which would make the work (33). Because of the precision used in the paper (33), some codon coefficients were 0.00, which would make the work (33). Because of the precision used in the paper (33), some codon coefficients were 0.00, which would make the work (33). Because of the precision used in the paper (33), some codon coefficients were 0.00, which would make the work (33). Because of the precision used in the paper (33), some codon coefficients were 0.00, which would make the work (33). Because of the precision used in the paper (33), some codon coefficients were 0.00, which would make the work (33). Because of the precision used in the paper (33), some codon coefficients were 0.00, which would make the work (33). Because of the precision used in the paper (33), some codon coefficients were 0.00, which would make the work (33).

We had to consider two factors before deciding on the group size \( n \). One is that each group should contain enough gene sequences so that our computational results were statistically representative. The other factor is that the difference of CAI values in the groups should be large enough so that we could perform meaningful comparison. Based on the two factors, we took \( n \) to be 300.

In addition, we considered four gene length thresholds \( s_l < s_u < l_l < l_u \), where \( l_l \) and \( l_u \) are approximately twice of \( s_l \) and \( s_u \), respectively. Then we selected gene sequences with lengths between \( s_l \) and \( s_u \) bp and gene sequences between \( l_l \) and \( l_u \) bp. We called them short and long sequences. Among the short sequences, we again took 300 sequences with the highest/lowest CAI values as the groups \( g_{sh} \) and \( g_{sl} \). Similarly, among long sequences, we took 300 sequences with the highest/lowest CAI values as the groups \( g_{lh} \) and \( g_{ll} \). The length thresholds were selected so that the difference between average CAI of groups \( g_{sh} \) and \( g_{sl} \) is about the same as the one between average CAI of groups \( g_{lh} \) and \( g_{ll} \). For \( B. subtilis \), we set \( s_l = 600, s_u = 800, l_l = 1200 \) and \( l_u = 1800 \). For \( S. pombe \), we set \( s_l = 700, s_u = 1100, l_l = 1300 \) and \( l_u = 2100 \). For \( S. cerevisiae \), we set \( s_l = 600, s_u = 900, l_l = 1200 \) and \( l_u = 1700 \). And for \( E. coli \), we set \( s_l = 600, s_u = 900, l_l = 1200 \) and \( l_u = 1800 \).

The tRNA gene copy numbers in \( B. subtilis, S. pombe \) and \( E. coli \) were extracted from GrtRNADB, a database containing tRNA genes predicted by tRNAscan-SE (34). The tRNA gene copy number in \( S. cerevisiae \) was extracted from Percudani et al.’s result (35), in which the yeast genome was scanned by Pol3scan (36).

**Methods**

**Requirements for tRNA weak binding and slippage.** We considered that there are two kinds of base pairs: standard pairs and wobble pairing at the third base. A cognate tRNA carries the standard Watson–Crick pairing or nearly standard wobble pairing matching anticodon for a given codon, while a near-cognate tRNA allows mildly and less favorable wobble pairing at the third base. For a near-cognate tRNA to slip and bind the codon at the shifted frame, we require that the anticodon form at least two base pairs with the codon. The rules of wobble pairing were adopted from several publications (37–39; see also Supplementary Tables). Furthermore, it is well known that some anticodons of tRNA undergo certain modifications in vivo. For example, base U in an anticodon can be modified to 5-methyl-2-thiouridine derivatives, 2′-O-methyluridine, 5-hydroxymethyluridine derivatives and other forms. The complementary base depends on the modification. For example, U with some modifications can pair with A and G whereas other modifications allow U to pair with A, G. Consequently, anticodon modifications have to be taken into account in the definition of sets \( V^+ \) and \( R^+ \) that are used to compute the FRS. We obtained anticodon modification data from the compilation of tRNA sequences and other references (40,41) (the full list is included in Supplementary Data). In \( S. pombe \), when modification of an anticodon was unknown, we used the known modification of the same anticodon in \( S. cerevisiae \).

**Permutation test.** Given a real gene sequence, 1000 sequences were generated by randomly permuting synonymous codons. More specifically, for each one of the 20 amino acids we record all positions of this particular amino acid in the sequence. Then we randomly permute the codons on these selected positions. This procedure is carried out separately for each amino acid. This way, both CAI and amino acid composition were preserved in the random sequences. Then, FRS of the real sequence was tested against the vector of FRS of random sequences by one sample \( t \)-test. In this way, when given a set of real gene sequences, we could check how many real gene sequences have FRS that is significantly lower or higher compared to the FRS of their random counterparts. After obtaining these two numbers, we applied a \( \chi^2 \)-test to check the significance. The null hypothesis is that these two numbers should be equal.

**Basic statistics of FRS.** For sequences of similar length, we measured if FRS of sequences with high CAI was significantly higher than FRS of sequences with low CAI. Given two sequence groups, FRS of the sequence in each group was computed and put in a separate group. Then two groups of FRS would be compared using unpaired \( t \)-test.

**Comparing FRS at the beginning and towards the end of a sequence and gradient computations.** To measure if FRS of sequence segment at the end of a gene sequence was significantly higher than the one of sequence segment at the start of that sequence, for each sequence in a group, we computed FRS of the first 100 and the last 100 codons and put two scores in two groups separately. Then, we performed a paired \( t \)-test to compare two groups of FRS. To perform the gradient test, for each sequence in a given group, we computed the pair (FRS, \( n_i \)), \( i = 1, \ldots, m \), where \( FRS \) is the FRS of the \( i \)-th \( k \) codons of the gene and \( n_i = i^k \). Then we pooled all the data together and fit a straight line with linear regression between FRS and \( n_i \). The slope of the line was taken as the gradient of FRS along genes’ length in the group.
We set $m = 40$ and $k = 5$. The first 50 codons were removed from *E. coli* gene sequences when we computed FRS gradient.

RESULTS

The pause-and-slip model

Our frameshifting model (Figure 1) is based on the pause-and-slip mechanism introduced by Farabaugh and Bjork (29). These authors proposed that translational frameshifting is a two-step process. First, there is a translational pause, caused by an empty ribosomal A-site resulting from insufficient amount of a cognate tRNA and the ensuing competition between the cognate tRNA with a near-cognate tRNA. Since near-cognate tRNA forms a suboptimal bond with the codon, it would fall off more easily than the cognate tRNA. As a result, the A-site might remain unoccupied for a longer time. We refer to this step as the ‘pause’ step. After the near-cognate tRNA successfully but weakly binds the codon at the A-site, it translocates to the P-site. In the second (slip) step, while at the P-site, the weakly bound near-cognate tRNA might slip to the left or right in case its anticodon is also capable of binding the codon in the shifted frame. Because here we focus on uncovering possible optimization against frameshifting errors rather than the impact of a missense error followed by a frameshifting error, we considered only those competing near-cognate tRNAs that are isoacceptors, i.e. carry the same amino-acid as the cognate tRNA.

If a tRNA slips to the left/right by one nucleotide, we refer to such an event as $-1$ or $+1$ frameshifting, respectively, where the sign indicates the frameshifting direction. Although, following Farabaugh and Bjork (29), we used the pause-and-slip scheme to model both $-1$ and $+1$ frameshifting errors, one has to keep in mind that these two events are not fully symmetric because the slip step in $-1$ frameshifting is likely to involve additional site-dependent factors not included in the model (42). Figure 1 illustrates $+1$ frameshifting under the pause-and-slip model.

Frameshifting robustness score

Based on the pause-and-slip model, we propose a new measure, the FRS, which estimates how resistant a given sequence is to frameshifting errors during translation. For the FRS calculation, we use tRNA gene copy number in the respective genome as a proxy for tRNA abundance. For *E. coli*, we repeated the computation using actual tRNA abundance data, with consistent results. The FRS for $+1$ frameshifting in a gene $g$ is:

$$FRS(g)^{+1} = 1 - \frac{1}{m} \sum_{i=1}^{m_c} \frac{b \sum_{i \in V_i^{+1}} n_i}{b \sum_{i \in R_i^{+1}} n_i + b \sum_{i \in R_i^{-1}} n_i + n_{tc}}$$

where $n_i$ is the number of codons in $g$ used to compute the score; the set $V_i^{+1}$ contains near-cognate tRNAs that can

Figure 1. $+1$ frameshifting in the pause-and-slip model. (A) The competition between small number of cognate tRNA and relatively large number of near-cognate tRNA keeps A-site empty for a relatively long time. (B) A near-cognate tRNA enters the A-site, forming a weak bond with the codon at the A-site. (C) The near-cognate tRNA is translocated to P-site and the binding is not stable. (D) The near-cognate tRNA slips one nucleotide to the right, forming a bond with the codon at the $+1$ shifted frame.
slip to the right at the P-site after binding the i-th codon \( c_i \) and the set \( R^i \) contains near-cognate tRNAs that are assumed to be unable to slip at the P-site after binding codon \( c_i \) (for details see ‘Materials and Methods’ section); \( n_i \) is the gene copy number of the tRNA \( t_i \); \( t_c \) is the cognate tRNA of codon \( c_i \), and \( b \) is a positive constant less than one, which we call weak binding coefficient. The multiplication by \( b \) reflects the fact that near-cognate tRNAs bind codons less stably (with a lower probability) than cognate tRNAs. Here, we report results for \( b = 0.01 \) but the results were robust for \( b \) values between 0.001 and 0.01 (data not shown). Thus the term, \( b \sum_{i \in R^i} n_i \), measures the binding potential of tRNA \( t_i \), which allows for +1 frameshifting after binding codon \( c_i \). The denominator, \( b \sum_{i \in R^i} n_i + b \sum_{i \in R^i} n_i + n_{ci} \), reflects the binding potential of all tRNAs that compete for binding codon \( c_i \). If we let

\[ p_i = b \sum_{i \in R^i} n_i \left( b \sum_{i \in R^i} n_i + b \sum_{i \in R^i} n_i + n_{ci} \right), \]

then \( p_i \) measures the probability of the occurrence of +1 frameshifting at the codon \( c_i \). So \( 1 - p_i \) measures the probability that frameshifting error does not occur at the codon. The +1 FRS of a sequence is the average of such probabilities over all codons in the sequence. A higher score indicates that the sequence is more resistant to +1 frameshifting errors. When computing the score, we did not consider stop codons, non-degenerate codons AUG and UGG, just as these codons are disregarded for computing the CAI. In addition, we disregarded those codons for which the corresponding codon in the +1 shifted frame is a stop codon. FRS for –1 frameshifting was defined in a similar way.

It would be natural to assume that the anticodon carried by a cognate tRNA is the exact Watson–Crick complement of the corresponding codon. However, this approach would not account for two important aspects of codon–anticodon interactions: anticodon modifications and wobble pairing. Anticodons often undergo certain modifications in vivo. With such modification, the base \( U \), for instance, can pair with either \( A \) and \( G \) or with \( U \), \( A \) and \( G \). Furthermore, bases in the third codon position can form a non-standard, the so called wobble pair, with the first position of the anticodon. Consequently, wobble pairing and anticodon modifications impact the FRS score. Not all wobble pairs are equally favorable. Some form nearly standard pairs whereas others are less favorable (38,39). Let us refer to a codon followed by a nucleotide \( \text{NNN} \) as a +1 frameshifting sensitive combination if a +1 frameshifting event can occur at the codon \( \text{NNN} \). Applying Crick’s original pairing rules (43), which assumes only unmodified \( G \), \( C \), \( A \), \( U \) and \( I \) in any anticodon, there are 14 +1 frameshifting sensitive combinations in \( S. cerevisiae \). With anticodon modifications, this number increases to 27. We consider a tRNA to be cognate for a given codon if its anticodon forms standard Watson–Crick pairs with the corresponding codon’s first and the second bases and forms a standard or nearly standard wobble pair at the third base. For near-cognate tRNA, the anticodon is allowed to form mildly and less favorable wobble pairs with the third base of the codon. Finally, we assume that the anticodon of a near-cognate tRNA that binds in a shifted position after slippage should form at least two base pairs in the shifted frame, where only nearly standard and mildly favorable wobble pairs are allowed. On the basis of the above binding rules, the set \( V^i \) contains all near-cognate tRNAs that can bind \( \text{NNN} \) and \( \text{NNN} \), whereas \( R^i \) contains all near-cognate tRNAs that are only capable of binding \( \text{NNN} \). Full details for base pairing in original and shifted frames are given in the ‘Materials and Methods’ section.

Statistics of the FRS

First, we selected two prokaryotes \( E. coli \) and \( B. subtilis \), and two eukaryotes \( S. cerevisiae \) and \( S. pombe \), for which (relatively) complete information on anticodon modification is available. We computed +1/−1 FRS for all available gene sequences from these organisms. In the computation of the FRS, we set the weak binding coefficient \( b \) to 0.01. The mean, the standard deviation and the minimum of the corresponding +1/−1 FRS values are listed in Table 1 (the maximum +1/−1 FRS in all four organisms was 1). In these computations, we used tRNA gene copy number as a proxy for tRNA abundance. The actual tRNA abundance has been experimentally measured in \( E. coli \) (44), so we recomputed +1/−1 FRS of \( E. coli \) with the experimental tRNA abundance data. The results were very similar to those obtained with the tRNA gene copy number (Table 1).

Notably, with the exception of \( B. subtilis \), +1 FRS was significantly higher than −1 FRS in the other three organisms (\( P < 1 \times 10^{-10} \)). The relatively lower −1 FRS might indicate a lower pressure for optimizing against −1 frameshifting.

Because it has been proposed that frameshifting robustness should be higher for longer and more highly expressed genes, to capture such differences appropriately, the genes in each organism were classified into four subsets (see ‘Materials and Methods’ section for details of the definitions): \( g_{sh} \)– short sequences with high CAI values; \( g_{sl} \)– short sequences with low CAI values; \( g_{lh} \)– long sequences with high CAI values; and \( g_{ll} \)– long sequences with low CAI values. Within our length threshold, short proteins are typically single-domain whereas long proteins are typically multi-domain ones.

First, the FRS values of gene sequences in the \( g_{sl} \) and \( g_{sh} \) sets, and in the \( g_{ll} \) and \( g_{lh} \) sets were compared using an unpaired t-test. For \( B. subtilis \), \( S. cerevisiae \) and \( S. pombe \), the FRS values of sequences in \( g_{sh} \) and \( g_{sl} \) were significantly higher than the FRS values of sequences in \( g_{sh} \) and \( g_{ll} \), respectively (Table 2). Thus, compared to the sequences with lower CAI of similar length, the sequences with higher CAI had higher FRS, consistent with the expectation that the latter, presumably highly expressed, genes should be more strongly optimized against frameshifting errors.

In contrast, in \( E. coli \), the +1 FRS values of sequences in \( g_{ll} \) were significantly higher than the FRS values of
The symbols ‘>’ indicate the mean FRS of sequences in gsh is higher/lower than the mean FRS of sequences in gll. ‘?’ is used for tests resulting in $P > 0.1$. The probability that the observation is by chance is shown in the parentheses.

Ec, E. coli, Bs, B. subtilis, Sc, S. cerevisiae and Sp, S. pombe.

| gsh versus gll | Ec/C0 | Bs/C0 | Sc/C0 | Sp/C0 |
|----------------|-------|-------|-------|-------|
| gll versus gsh | < (1e-4) | (3e-13) | > (0) | > (0) |

The probability for the correlation is shown in parentheses. ‘?’ is used for tests resulting in $P > 0.05$.

Ec, E. coli, Bs, B. subtilis, Sc, S. cerevisiae and Sp, S. pombe.

sequences in gsh ($P < 1e-4$), and the $-1$ FRS values of sequences in gll were significantly higher than the sequences in gsh ($P < 0.02$); other comparisons for E. coli had insignificant $P$-values (Table 2). It has been reported that in E. coli the codons at the beginning and at the end of a gene sequence are under different selection pressures (22,45), so we repeated the computations after removing 50 codons at the start and 20 codons at the end of genes. With this modification, $+1$ FRS of sequences in gll remained significantly higher than the ones of sequences in gsh ($P < 3e-3$) whereas the other comparisons had insignificant $P$-values.

FRS is correlated with CAI and protein abundance

To further characterize the relation between FRS and CAI, we computed, for each organism, the partial correlation coefficients between FRS and CAI for all sequences, and for sequences in two sequence groups (gsh and gll) of low and high CAI values, respectively. Additionally, we computed the same correlation in each group gsh, gll, with gsh and gll, to control for the potential effect of gene length and CAI level. The results are shown in Table 5, Figure 2 and Supplementary Data. To ensure that the value of binding coefficient did not significantly affect the results, we repeated the computations for $b = 0.001$, with consistent results (data not shown).

For B. subtilis, S. cerevisiae and S. pombe, $+1$ FRS showed significant correlation with CAI (except for B. subtilis in group gsh). This result confirms that gene sequences with high CAI, on average, appear to be more

We computed partial correlation between FRS and protein abundance/gene length taking gene length/protein abundance (46) (to the best of our knowledge, no large-scale protein abundance data set for B. subtilis is available) as the control variable (Table 4). The results are consistent with the results obtained for CAI. Again, we observed a significant positive correlation of FRS with protein abundance level in S. cerevisiae and S. pombe but not in E. coli.

To further characterize the relation between FRS and CAI, we computed, for each organism, the Pearson correlation coefficients between FRS and CAI for all sequences, and for sequences in two sequence groups (gsh and gll) of low and high CAI values, respectively. Additionally, we computed the same correlation in each group gsh, gll, with gsh and gll, to control for the potential effect of gene length and CAI level. The results are shown in Table 5, Figure 2 and Supplementary Data. To ensure that the value of binding coefficient did not significantly affect the results, we repeated the computations for $b = 0.001$, with consistent results (data not shown).

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Figure 2. The relation between +1 FRS and CAI of gene sequences in four groups of B. subtilis: (A) g_{sl}, (B) g_{sh}, (C) g_{sl}, and (D) g_{sh}.

Table 4. Pearson partial correlation between FRS and protein abundance/gene length for E. coli, S. cerevisiae and S. pombe with length/protein abundance as the control variable

| +1 FRS versus protein abundance | −1 FRS versus protein abundance | +1 FRS versus length | −1 FRS versus length | Number of tested genes |
|---------------------------------|---------------------------------|---------------------|---------------------|------------------------|
| Ec                              | −0.091 (0.0038)                 | −0.045 (?)          | 0.11 (3.69e-4)      | −0.036 (?)             | 1005                   |
| Sc                              | 0.19 (1.63e-31)                 | 0.18 (6.64e-29)     | −0.022 (?)          | 0.051 (1.71e-3)        | 3839                   |
| Sp                              | 0.23 (5.15e-19)                 | 0.15 (1.14e-8)      | −0.15 (2.46e-9)     | −0.062 (0.017)         | 1465                   |

The probability for the correlation is shown in parentheses. “?” is used for tests resulting in $P > 0.05$.

Ec, E. coli, Bs, B. subtilis, Sc, S. cerevisiae and Sp, S. pombe.

Table 5. Correlation between +1/−1 FRS and CAI in sequence groups of E. coli, B. subtilis, S. cerevisiae and S. pombe

| +1 FRS versus CAI | +1 FRS versus CAI | +1 FRS versus CAI | +1 FRS versus CAI | −1 FRS versus CAI | −1 FRS versus CAI | −1 FRS versus CAI | −1 FRS versus CAI |
|-------------------|-------------------|-------------------|-------------------|-------------------|-------------------|-------------------|-------------------|
| Ec                | −0.06 (1e-4)      | 0.31 (1e-16)      | 0.37 (3e-12)      | 0.47 (1e-16)      | 0.06 (4e-5)       | 0.26 (1e-16)      | 0.35 (3e-12)      | 0.23 (1e-15)      |
| Bs                | 0.05 (0.4)        | 0.14 (0.02)       | 0.23 (5e-5)       | 0.23 (7e-5)       | −0.03 (0.7)       | −0.004 (0.9)      | 0.04 (0.5)        | 0.07 (0.2)        |
| Sc                | −0.07 (0.2)       | 0.32 (1e-8)       | 0.31 (4e-8)       | 0.14 (2e-2)       | 0.24 (5e-5)       | 0.30 (7e-7)       | 0.27 (3e-6)       | 0.16 (6e-3)       |
| Sp                | 0.15 (0.01)       | 0.1 (0.09)        | 0.21 (2e-4)       | 0.22 (1e-4)       | −0.23 (5e-5)      | 0.04 (0.5)        | 0.24 (3e-5)       | 0.07 (0.2)        |
| g_{sl}            | −0.16 (4e-3)      | 0.31 (6e-8)       | 0.41 (1e-13)      | 0.47 (4e-16)      | 0.21 (3e-4)       | 0.34 (1e-9)       | 0.42 (3e-14)      | 0.30 (9e-8)       |
| g_{sh}            | 0.23 (6e-5)       | 0.07 (0.2)        | 0.17 (4e-3)       | 0.24 (4e-5)       | −0.20 (5e-4)      | −0.08 (0.1)       | 0.16 (4e-3)       | 0.11 (0.05)       |
| g_{sl}            | −0.17 (3e-3)      | 0.38 (1e-11)      | 0.44 (1e-15)      | 0.50 (1e-16)      | 0.35 (4e-10)      | 0.42 (5e-14)      | 0.44 (1e-15)      | 0.37 (3e-11)      |

The probability for the correlation is shown in parentheses. Ec, E. coli, Bs, B. subtilis, Sc, S. cerevisiae and Sp, S. pombe.
resistant to +1 frameshifting errors than sequences with lower CAI. Furthermore, within the same length range (groups $g_{st}$ and $g_{bh}$, and groups $g_{h}$ and $g_{ln}$), sequences with higher CAI showed a stronger correlation between FRS and CAI. In addition, for all 3 organisms, long gene sequences with high CAI ($g_{ln}$) showed the strongest correlation between FRS and CAI. For these 3 organisms, the relation between −1 FRS and CAI was similar to the one observed for +1 frameshifting with the exception that for $B. subtilis$ −1 FRS for sequences in $g_{l}$ and $g_{y}$ showed insignificant correlation with CAI. The correlation between FRS, especially −1 FRS, and CAI for sequences in groups $g_{l}$ and $g_{h}$, was often less significant, suggesting that differences in gene length might obscure the relationship between the two factors.

In $E. coli$, there was no consistent pattern of correlation between FRS and CAI. Surprisingly, for genes with low CAI, the +1 FRS showed significant correlation with CAI values but for genes with high CAI, there was significant negative correlation with CAI. Furthermore, −1 FRS also showed either a negative or a positive correlation with CAI depending on the sequence group (Table 5).

**Frameshifting robustness of gene sequences is higher than expected by chance**

In our next test, we asked whether the FRS values of real protein-coding sequences differed from the values calculated for sequences in which synonymous codons were randomly permuted (see ‘Materials and Methods’ section). For each organism, gene sequences longer than 200 codons were selected, and for each such gene, 1000 random sequences were generated by permuting synonymous codons. Under this procedure of random sequence generation, the amino-acid sequence and the codon composition remain unchanged but the context that affects frameshifting robustness is randomized. We compared the value of FRS for the real and permuted sequences by testing how many real gene sequences had FRS significantly ($P < 0.01$) lower or higher compared to random sequences. The test results are not sensitive to $P$-value cut-off. We repeated the test for $P$-value cut-off set at 0.001, 0.0001 and 0.00001, with very similar results. The results indicate that, as one might expect, +1 FRS scores were, on average, higher than the corresponding values for randomized sequences in $E. coli$ and $B. subtilis$; however, the two yeasts did not show this difference (Figure 3A). We repeated this analysis with the FRS computed only for the last 200 codons in each real and permuted sequence. In this comparison, significantly higher +1 FRS values were observed for all four organisms (Figure 3C). The $p$-value for $E. coli$, $B. subtilis$ and $S. cerevisiae$ in the last 200 codon comparison is $< 2.2e-16$ and the $p$-value for $S. pombe$ is $< 0.004$. The comparison of −1 FRS revealed higher values in real sequences compared to randomized sequences for three organisms but $E. coli$ was once again an outlier and showed the opposite trend (Figure 3D).

This observation suggested the possibility of a positive gradient of FRS along gene sequences.

**The FRS increases along the gene-sequence length**

The results of the above test indicated that FRS might not be distributed uniformly along the protein-coding sequence but instead might be higher in the 3′-terminal, distal part of the sequence, in accord with the selection for minimization of frameshifting error rate. To further assess this possibility, we computed the FRS of the first 100 and the last 100 codons in each of the analyzed sequences and compared the values using paired t-test (Table 6).

For $B. subtilis$, $S. cerevisiae$ and $S. pombe$, whenever the comparison revealed a significant difference, the FRS of the 100-codon segment toward the end of a gene was higher than the FRS of the segment at the start of the gene. When sequences of similar lengths were compared, the difference of the +1 FRS values for proximal and distal sequences segments were more significant for sequences with high CAI than for sequences with low CAI, except for $B. subtilis$ gene sequences in $g_{l}$ and $g_{y}$. $E. coli$ demonstrated a different behavior again. In the case of +1 FRS, the results were not significant, and in the case of −1 frameshifting, the −1 FRS values of N-terminus segments were even significantly higher than −1 FRS values of the C-terminal segments (Table 6). When the computation was repeated after removing 50 codons at the start and 20 codons at the end of the genes, this reversed trend was also observed for +1 FRS in the $g_{st}$ and $g_{ln}$ groups of $E. coli$ genes.

Finally, we computed the gradient of FRS of gene sequences in four sequence groups for $E. coli$, $B. subtilis$, $S. cerevisiae$ and $S. pombe$ (see ‘Materials and Methods’ section). We observed that, except for $B. subtilis$ sequences in $g_{ln}$, all significant +1 FRS gradients were positive, that is, the resistance to +1 frameshifting errors tends to increase along the length of genes. All −1 FRS gradients were non-significant (Table 7).

**DISCUSSION AND CONCLUSIONS**

It has long been accepted that translation speed and accuracy are two major factors that shape an organism’s codon usage bias. The hypothesis that codon usage bias is optimized to minimize missense errors during translation is fairly well supported (12,13,47). However, processivity errors seem to occur with frequencies comparable to those of misense errors and are potentially more costly. A number of recent studies provided indirect arguments in support of the hypothesis that codon usage is optimized to avoid frameshifting errors. Based on the pause-and-slip molecular model of frameshifting, we developed the FRS to assess resistance of genes to frameshifting errors on the genome scale. Using FRS, we tested whether the codon patterns in 4 well-studied model organisms, $E. coli$, $B. subtilis$, $S. cerevisiae$ and $S. pombe$, were compatible with the hypothesis that codon usage is optimized to avoid frameshifting errors.

The results indicate that the patterns of codon usage in the two yeasts, $S. cerevisiae$ and $S. pombe$ and in the
bacterium *B. subtilis* are consistent with the hypothesis of optimization against frameshifting errors (under the pause-and-slip model). In these three organisms, we observed significant positive correlation between +1/−1 FRS and the codon usage bias measured by CAI and protein abundance. Furthermore, when length-matched sets of sequences were compared, the genes with high CAI have higher FRS than the genes with lower CAI. We also reasoned that, if there is optimization against frameshifting errors, one should expect that the sequences are more highly optimized against such errors towards the end of the sequence, and for sequences with similar CAI, longer sequences would be more strongly optimized. In agreement with these predictions, we observed that FRS

![Figure 3](https://academic.oup.com/nar/article-abstract/37/20/6799/1108377)

**Figure 3.** The number of real gene sequences whose +1/−1 FRS is significantly higher (blue bar) or lower (red bar) than FRS of random sequences generated by permuting its synonymous codons in four organisms. (A) +1 FRS was computed using the whole real and random sequence. The *P*-value for *E. coli* and *B. subtilis* in the whole sequence comparison for +1 FRS score is 2.1e-9 and less than 2.2e-16, respectively. (B) −1 FRS was computed using the whole real and random sequence. (C) +1 FRS was computed using the last 200 codons of the real and random sequence. (D) −1 FRS was computed using the last 200 codons of the real and random sequence.

|                  | Ec | Bs | Sc | Sp |
|------------------|----|----|----|----|
| +1 FRS start     |    |    |    |    |
| Ec versus end    |    |    |    |    |
| +1 FRS start     |    |    |    |    |
| Bs versus end    |    |    |    |    |
| +1 FRS start     |    |    |    |    |
| Sc versus end    |    |    |    |    |
| +1 FRS start     |    |    |    |    |
| Sp versus end    |    |    |    |    |
| +1 FRS start     |    |    |    |    |
| Ec versus end    |    |    |    |    |
| −1 FRS start     |    |    |    |    |
| Ec versus end    |    |    |    |    |
| −1 FRS start     |    |    |    |    |
| Bs versus end    |    |    |    |    |
| −1 FRS start     |    |    |    |    |
| Sc versus end    |    |    |    |    |
| −1 FRS start     |    |    |    |    |
| Sp versus end    |    |    |    |    |
| −1 FRS start     |    |    |    |    |

The symbols ‘>’/‘<’ indicate the mean FRS of sequence segment at the start of the sequences in the group is higher/lower than the mean FRS of sequence segment at the end of the sequences. ‘?’ is used for tests resulting in *P* > 0.1.

The probability for the correlation is shown in parentheses.

Ec, *E. coli*, Bs, *B. subtilis*, Sc, *S. cerevisiae* and Sp, *S. pombe*.
Ec, E. coli, Bs, B. subtilis

The probability for the correlation is shown in parentheses. 'NS' indicates gradients that were not significant ($P > 0.05$).

The results for *E. coli* were notably less consistent than those for other three organisms but nevertheless seem to be compatible with some degree of codon usage optimization for reducing frameshifting error rate. Surprisingly, in *E. coli*, we observed correlation between +1 FRS and CAI only for groups of genes with low CAI. Nevertheless, we observed the expected +1 FRS gradient for long genes with high CAI values. At present, we do not have a definitive interpretation of the anomalous results obtained for *E. coli* genes. However, combined with the observations that (i) FRS of *B. subtilis* genes was, on average, much lower than the FRS of *S. cerevisiae* and *S. pombe* genes, (ii) +1 FRS of genes of *B. subtilis* showed weaker correlation with CAI compared to genes of *S. cerevisiae* and *S. pombe* and (iii) the only significant +1 FRS gradient in *B. subtilis* is negative, it is tempting to hypothesize that prokaryotic genes, in general, are subject to weaker selection for minimization of frameshifting errors than eukaryotic genes.

Similarly, as it was the case for codon bias, the relation of FRS with gene length (when corrected for the correlation with CAI and protein abundance) is not transparent. There is clear and consistent correlation for *E. coli*, whereas the remaining three organisms show no significant correlation or even an anti-correlation. Nevertheless, for these three organisms, the gene group consisting of long genes with high CAI showed the highest positive correlation between CAI and FRS.

Comparison of +1 and −1 FRS reveals substantial differences. Compared to +1 FRS, −1 FRS was usually smaller, showed lower correlation with CAI, smaller differences between FRS values at the beginning and at the end of a gene sequence, and no significant FRS gradient. One possible explanation is that +1 frameshifting errors are more likely than −1 frameshifting errors, so gene sequences are subject to greater pressure for optimization against +1 frameshifting errors. Indeed, examination of the data in the translational recoding event database RECODE (49) shows that most of programmed framenches in *B. Subtilis, S. cerevisiae* and *S. pombe* involve +1 frameshifting.

The approach employed here to estimate optimization of protein-coding sequences for minimization of frameshifting errors has several limitations. First, we applied the simplified pause-and-slip model that was developed for explaining programmed frameshifting. Although it is believed that programmed frameshifting and frameshifting errors share the same mechanism, other frameshifting models cannot be ruled out. For instance, an often considered model invokes out-of-frame binding (without slippage) to explain programmed frameshifting (50). Thus, more experimental studies are necessary to test the validity of the pause-and-slip model. The second limitation of the approach relates to the use of the tRNA gene copy number as an approximation for tRNA abundance. Although the strong positive correlation between these two quantities has been demonstrated in many independent studies for diverse organisms, using tRNA abundance data directly would be desirable. However, this information is unavailable for most organisms and cannot be expected to become available in the near future. In contrast, tRNA gene copy number is easy to obtain computationally for any sequenced genome. In the only model organism, *E. coli*, for which the actual tRNA abundance was measured, the FRS estimates using experimental data were very close to those obtained with the tRNA gene copy number. Finally, whenever available, we used information about anticodon modifications to properly model codon–anticodon binding. Such information is essential to determine frameshifting sensitive sites and to obtain realistic values of the FRS of gene sequences. However, the list of anticodon modifications

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**Table 7. The gradient of +1/−1 FRS of the first 200 codons of gene sequence for *E. coli, B. subtilis, S. cerevisiae* and *S. pombe***

|        | Ec | Bs | Sc | Sp |
|--------|----|----|----|----|
| $g_{+1}$ | NS | NS | NS | NS |
| $g_{-1}$ | NS | NS | NS | NS |
| $g_{+2}$ | 1.4e-5 (1.2e-3) | -3.0e-5 (0.043) | 2.2e-5 (1.9e-7) | 5.2e-7 (6.4e-6) |
| $g_{-2}$ | NS | NS | NS | NS |

‘NS’ indicates gradients that were not significant ($P > 0.05$).
for each organism is not necessarily complete, so this incompleteness could also affect the results. Notwithstanding these possible drawbacks, the trends uncovered in this study were highly consistent and thus unlikely to be coincidental.

Codon usage is affected by multiple factors, such as translation efficiency, translation errors, GC content, mRNA secondary structure, and mRNA half life, and thus can be expected to be (partially) optimized for multiple biological purposes. We show here that codon usage is, at least to a certain extent, optimized to reduce frameshifting errors, a finding that fits the general concept of mistranslation-induced protein misfolding being a major factor affecting gene evolution. It remains to be shown how various factors interact with each other during evolution to shape the landscape of codon usage bias.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

**ACKNOWLEDGEMENTS**

The authors thank Dr Philip J. Farabaugh (UMBC) for insightful discussion, Dr Alejandro Schaffer and Dr Jie Zhang (NCBI/NIH) for helpful comments on the article.

**FUNDING**

Intramural Research Program of the National Institutes of Health, National Library of Medicine. Funding for open access charge: Intramural research program for the National Institutes of Health, National Library of Medicine.

Conflict of interest statement. None declared.

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Nucleic Acids Research, 2009, Vol. 37, No. 20 6809

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