Gene expressivity is the main factor in dictating the codon usage variation among the genes in *Pseudomonas aeruginosa*

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Abstract

Codon usage biases of all DNA sequences (length greater than or equal to 300 bp) from the complete genome of *Pseudomonas aeruginosa* have been analyzed. As *P. aeruginosa* is a GC-rich organism, G and/or C are expected to predominate in their codons. Overall codon usage data analysis indicates that indeed codons ending in G and/or C are predominant in this organism. But multivariate statistical analysis indicates that there is a single major trend in the codon usage variation among the genes in this organism, which has a strong negative correlation with the expressivities of the genes. The majority of the lowly expressed genes are scattered towards the positive end of the major axis whereas the highly expressed genes are clustered towards the negative end. This is the first report where the prokaryotic organism having highly skewed base composition is dictated mainly by translational selection, though some other factors such as the lengths of the genes as well as the hydrophobicity of genes also influence the codon usage variation among the genes in this organism in a minor way. © 2001 Elsevier Science B.V. All rights reserved.

Keywords: Synonymous codon usage; Correspondence analysis; Translational selection; *Pseudomonas aeruginosa*; Highly expressed genes; Lowly expressed genes

1. Introduction

It is well established that the codon usage pattern is non-random and species-specific. It has also been reported that there is significant variation of codon usage bias among the different genes within the same organism (Grantham et al., 1981; Wada et al., 1990). Biased usage of synonymous codons may result from various factors. In organisms having highly skewed base composition, it was observed that compositional constraints are the main factors in determining the codon usage variation among the genes (Ohkubo et al., 1987; Ohama et al., 1990; Andersson and Sharp, 1996b; Andersson et al., 1998; Musto et al., 1998). It was suggested that translational selection manifests the codon usage biases of highly expressed genes and subsequently it was reported that preferred codons in highly expressed genes are recognized by most abundant tRNAs (Ikemura, 1981a,b, 1982; Bennetzen and Hall, 1982). In some unicellular organisms it has been reported that both translational selection and compositional constraints operate in dictating the codon usage variation among genes (Ikemura, 1981a,b; Gouy and Gautier, 1982; Andersson and Sharp, 1996a; Romero et al., 2000b; Ghosh et al., 2000). In recent years, codon usage analyses on several complete genomes has indicated that the physical location of each gene on the chromosome determines codon usage patterns (Kerr et al., 1997). It has also been reported that in different organisms, replication-translational selection is the main source of variation of codon usage biases among the genes (McInerney, 1998; Lafay et al., 1999; Romero et al., 2000a). Recently, it has been reported that the hydrophobicity of each gene is one of the main factors in determining the codon usage variation among the genes in *Mycobacteria* (de Miranda et al., 2000).

*Pseudomonas aeruginosa* is a GC-rich Gram-negative bacteria. It has been considered as one of the major opportunistic human pathogens during the past century (Stover et al., 2000). *Pseudomonas aeruginosa* is a well-studied microorganism and is the largest prokaryote whose
complete genome has been published (Stover et al., 2000). Thus, extensive possibilities, earlier unavailable, have opened up to gain insight into the biology of this microorganism. In the present study, we have used all coding sequences (greater than or equal to 300 bp) and analyzed the codon usage data with a view to understanding the genetic organization of the P. aeruginosa genome. Our results demonstrate that translational selection is the major source of codon usage variation among the genes in this highly skewed compositional genome.

2. Materials and methods

The complete genome of P. aeruginosa has been downloaded from www.ncbi.nlm.nih.gov/genbank/genomes. To minimize sampling errors we have taken only those sequences which are greater than or equal to 300 bp and have the correct initial and termination codons. Finally, 5239 sequences were selected for data analysis. The coding sequences from the complete genome were retrieved using a program in C, developed by us.

Relative synonymous codon usage (RSCU) was used to study the overall codon usage variation among the genes. RSCU is defined as the ratio of the observed frequency of codons to the expected frequency if all the synonymous codons for those amino acids are used equally (Sharpe and Li, 1986). RSCU values greater than 1.0 indicate that the corresponding codon is more frequently used than expected, whereas the reverse is true for RSCU values less than 1.0.

GC_{3s} is the frequency of (G + C) and A_{3s}, T_{3s}, G_{3s}, and C_{3s} are the distributions of A, T, G and C at the synonymous third positions of codons. N_c is the effective number of codons used by a gene, generally used to measure the bias of synonymous codons and independent of amino acid compositions and codon numbers (Wright, 1990). The values of N_c range from 20 (when one codon is used per amino acid) to 61 (when all the codons are used with equal probability). The expected value of N_c under random codon usage can be expressed as:

\[ N_c = 2 + s + \left\lfloor \frac{29}{s + (1 - s)^2} \right\rfloor \]

where \( s = \frac{\text{GC}_{3s}}{\text{AC}_{3s}} \).

Gene expressivities were measured by calculating the parameter codon adaptation index (CAI) as defined by Sharpe and Li (1987).

All the above-mentioned parameters were calculated by using the program CodonW 1.3 (available at www.molbio-l.ox.ac.uk/cu). Correspondence analysis (Greenacre, 1984) available in CodonW was used to investigate the major trend in codon usage variation among genes.

GC skew, defined as the ratio of \((G - C)\) to \((G + C)\) along the DNA sequences, was calculated using a sliding window of 60 kb and a step size of 6 kb.

3. Results and discussion

3.1. Overall codon usage analysis

A total of 5239 genes were used in this study. Since P. aeruginosa is a GC-rich genome, it is expected that G and/or C containing codons will predominate in the coding regions. Table 1 shows the overall RSCU values of this organism. It is evident that codons ending in G and/or C are predominant in the entire coding region. However, the overall codon usage values may obscure some heterogeneity of codon usage bias among the genes that might be superimposed on the extreme genomic composition of this organism.

3.2. Heterogeneity of codon usage

Two indices, viz. effective number of codons used by a gene (N_c) and (G + C) percentage at the third synonymous codon positions (GC_{3s}) are generally used to study the codon usage variation among the genes in any organism. Fig. 1

Table 1 Overall codon usage data of P. aeruginosa genes^a

<table>
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<th>AA</th>
<th>Codon</th>
<th>N</th>
<th>RSCU</th>
<th>AA</th>
<th>Codon</th>
<th>N</th>
<th>RSCU</th>
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</tr>
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<td>Thr</td>
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<td>Ala</td>
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^a AA, amino acids; N, number of codons; RSCU, cumulative relative synonymous codon usage of 5239 genes.
shows the $N_c$ distribution of different genes in *P. aeruginosa*. The $N_c$ values range from 23.43 to 60.86 (with a mean of 32.72 and standard deviation of 4.77), indicating that there is a wide variation of codon usage bias among the genes. The heterogeneity of codon usage biases among the genes is further confirmed from the distributions of (G+C) at the third synonymous codon positions, shown in Fig. 2. From this figure it is obvious that (G+C) at the synonymous third position of codons varies from 23 to 97% with a mean of 86% and standard deviation of 6%. These results indicate that apart from compositional constraints, other trends might influence the overall codon usage variation among the genes in *P. aeruginosa*.

3.3. Exploring different factors in determining the codon usage variation

3.3.1. The $N_c$ plot

Wright (1990) suggested that a plot of $N_c$ against GC$_{3s}$ could be effectively used to explore the codon usage variation among the genes. It was demonstrated by Wright (1990) that the comparison of the actual distribution of genes with the expected distribution under no selection could be indicative, if the codon usage bias of the genes had some influence other than the genomic GC composition. In other words, if GC$_{3s}$ is the only determinant of the codon usage variation among the genes then the values of $N_c$ would fall on the continuous curve (the $N_c$-GC$_{3s}$ plot, see Fig. 3), representing random codon usage. From Fig. 3 it is evident that only a small number of points lie on the expected curve, mostly in the GC-rich regions. This indicates that compositional constraints play a role in defining the codon usage variation among those genes. There are also a large number of points which lie well below the expected curve (having low $N_c$ values), which indicate that these genes have an additional codon usage bias, other than genomic GC composition.

3.3.2. Multivariate statistical analysis

To explore the other possibilities in shaping the codon usage variation among the genes in *P. aeruginosa* we have subjected the data to multivariate statistical analysis, a method that has been successfully used to study the codon usage variation among genes in different organisms (Romero et al., 2000a,b; Ghosh et al., 2000; Wright and Bibb, 1992; Stenico et al., 1994; Musto et al., 1999). Correspondence analysis is one of the multivariate statistical analyses in which the data are plotted in a multidimensional space of 59 axes (excluding Met, Trp and stop codons) and then the most prominent axes contributing to the codon usage variation among the genes are determined. In the present work, correspondence analysis has been performed on RSCU values to minimize the effects of amino acid composition. Fig. 4 shows the positions of genes along the first two major axes. The first major axis accounted for 18.5% of the total variation while none of the other axes accounted for more than 5%. It must be remembered that...
although the first principal axis explains a substantial amount of variation of codon usage among the genes in this microorganism, its value is still lower than found in other organisms studied earlier (Alvarez et al., 1994; Musto et al., 1998). The low value might be due to the extreme genomic composition of this genome. It is also obvious from Fig. 4 that the majority of the points are clustered in a spherical shape around the origin of axes, having an average $N_c$ value of 32.72 and standard deviation of 4.77. This indicates that these genes have more or less similar codon usage biases. There are very few points that are widely scattered along the positive side of the first major axis with an average $N_c$ value of 45.04 and standard deviation of 6.42, indicating that codon usage biases of these genes are not homogeneous. From these results it can be stated that there is a single major trend in codon usage variation among the genes in *P. aeruginosa*.

3.3.3. Effect of gene expressivities on codon usage

For a long time it has been noted that in organisms with a highly skewed base composition, mutational bias is the main factor in shaping the codon usage variation among the genes whereas translational selection plays a minor role. Overall RSCU values (shown in Table 1) and $N_c$ plot (shown in Fig. 3) provide definite indications that mutational bias is acting in this organism in dictating the codon usage variation among the genes. However, correspondence analysis indicates that there is a single major

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Fig. 3. $N_c$ plot of *P. aeruginosa* genes. The continuous curve represents the expected curve between GC$_3$s and $N_c$ under random codon usage.
trend in the codon usage among the genes in this bacterium. To assess the effect of expressivities of genes on codon usage biases, we have calculated the expressivity for each gene of this organism. The CAI has been widely used to estimate the expressivities of genes by different workers (Gutierrez et al., 1996; Nakamura and Tabata, 1997; Pan et al., 1998; Tiller and Collins, 2000) and is now considered a well-accepted measure of gene expressivities. In this study we have used the CAI to measure the expression level of a gene. A scatter diagram of the positions of genes on the first major axis was plotted against their corresponding CAI values, as shown in Fig. 5. The correlation coefficient between the positions of genes along the first major axis against their corresponding CAI values as estimated from Fig. 5 is \(-0.920 (P < 0.01)\). This is a clear indication that gene expression is the main cause for the codon usage variation among the genes in this organism. The positions of the genes along the first major axis are also significantly negatively correlated with GC\(_{3s}\) \((r = -0.896, P < 0.01)\) and with the length of the gene \((r = -0.119, P < 0.01)\). We also obtained a significant negative correlation between the positions of the genes in the first major axis and the hydropathy of each protein \((r = -0.060, P < 0.01)\) and a positive correlation with the third major axis \((r = 0.060, P < 0.01)\). The positions of the genes along the second major axis are strongly positively correlated with GC\(_{3s}\) \((r = 0.207, P < 0.01)\). These results suggest that highly expressed genes have lower (G+C) content at their synonymous third codon positions than the lowly expressed genes. It also indicates that highly expressed genes are longer. Earlier it was observed that in case of \textit{Escherichia coli} there was a strong positive correlation between the synonymous codon usage and gene length and it was proposed that longer genes should impose stronger constraints on the codon usage bias (Moriyama and Powell, 1998). The positive correlation between the synonymous codon usage and gene length in this organism is not surprising since it was observed (by comparing gene sequences of \textit{P. aeruginosa}) that \textit{E. coli} is the closest relative of \textit{P. aeruginosa} (Stover et al., 2000).
3.3.4. Influence of replication-transcriptional selection on codon usage

In recent years it was reported that in several bacteria, codon usage bias is mainly dictated by transcriptional-translational selection (McInerney, 1998; Lafay et al., 1999; Romero et al., 2000a). We explored this possibility by studying codon usage bias in the leading and lagging strands of P. aeruginosa. For locating the leading and lagging strands we have calculated GC skew along the DNA of P. aeruginosa by taking a 60 kb window and a step size of 6 kb. GC skew is generally used to locate the leading and lagging strands of a prokaryotic organism and it has been reported that in most prokaryotes there is a change in sign of GC skew near the origin of replication (Lobry, 1996; McLean et al., 1998). Fig. 6 shows the GC skew of this organism. There is a change in sign around 240 kb and the major part of the genome lies on the negative side of the GC skew curve. It has also been reported that in the leading strand there is an excess of G over C and most of the genes lie on the leading strand (McInerney, 1998; McLean et al., 1998). But in the GC skew positive side, we have obtained only 2038 genes and the rest (3201 genes) are found to lie in the lagging strand. Earlier it was reported that in most prokaryotic genomes a larger number of genes are located in the leading strand than in the lagging strand (McLean et al., 1998). In this respect, the sequence of P. aeruginosa is exceptional in having more genes in the lagging strand. Interestingly, we have also observed that of the genes clustered near the
origin of the first two major axes generated by correspondence analysis, 91% are located in the positive side of the GC skew curve and 94% are located in the negative. From this it is clear that replication-transcriptional bias has no effect in shaping the codon usage variation among the genes in this organism. In conclusion it can be asserted that codon usage bias among the genes in *P. aeruginosa* is mainly dictated by translation selection, though some other factors also have an influence in a minor way. This is obviously a new paradigm on the codon usage biases in this highly skewed base compositional genome.

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**References**


Fig. 6. GC skew (the ratio of (G−C) and (G+C)) of the *P. aeruginosa* genome. The GC skew was calculated by taking a window of size 60 kb with a step of 6 kb.


