

Molecular Biology of Codon Usage in One-Horned Rhino (*Rhinoceros unicornis* L)



Biotechnology

KEYWORDS : Codon usage, Synonymous codons, Indian one-horned rhino

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ABSTRACT

Background: This is the first report on the codon usage pattern in Indian one-horned rhino (*Rhinoceros unicornis* L). No literature is available on codon usage bias in Indian one-horned rhino till to date. Indian one-horned rhino

is the pride of Assam, being the state animal of Assam. The phenomenon of unequal use of synonymous codons i.e. some codons are more preferred than others in genes is known as codon usage bias. Synonymous codons which vary from two to six codons encode a particular amino acid and these generally differ in the 3rd codon position. However, the usage pattern of synonymous codons differs not only among the organisms but also among the genes within an organism. The codon bias is commonly found in highly expressed genes.

Result and Discussion: In this study, we analyzed the codon usage pattern in some selected nuclear genes of one-horned rhino using bioinformatics approach. We found that the overall GC% ranged from 41.3 to 57.2 % with an average of 51.46 %. The GC3 % of rhino genes ranged from 53.3 to 73.6 % with an average of 65.2 %. ENC value varied from 50 to 60 with a mean value of 55. We found a significant negative correlation of ENC with GC ($r=-0.934^{**}$, $p<0.01$) and ENC with GC3 ($r=-0.979^{**}$, $p<0.01$). In addition, a significant positive correlation was found between GC12 and GC3 ($r=0.896^{**}$, $p<0.05$) of the genes. Further we observed significant positive correlation between A and A3 % ($r=0.964^{**}$, $p<0.01$), C and C3 % ($r=0.951^{**}$, $p<0.001$), G and GC3% ($r=0.959^{**}$, $p<0.001$).

Conclusion: We found that the codon usage bias in the analyzed coding sequences (cds) of rhino is not remarkable. Codon usage bias prefers GC bases at the 3rd codon position and most frequent codons end with G. Nucleotide constraint as well as compositional constraint under mutation pressure might have made major contribution in the codon usage bias in the coding sequences of the nuclear genes in one-horned rhino.

Introduction

Indian one-horned rhino (*Rhinoceros unicornis* L) is a highly endangered species belonging to the Schedule-I of animals of the Indian wildlife (Wildlife Protection Act, 1972) and it is widely distributed in the tall, wet grassland and riverside forests, foot hills of the Himalayas, particularly confined to the protected areas in Assam, West Bengal and Uttar Pradesh within India [1]. *Rhinoceros unicornis* is the pride of Assam and it is also the state animal of Assam. Chromosome number of rhino is $2n=82$ or 84 in which the male is heterogametic with XY allosomes but the female is homogametic with XX chromosomes [2]. For establishing the relationship between *Rhinoceros unicornis* and other mammals, the mitochondrial DNA sequences were used [3].

The sequence of three nitrogen bases which codes for a particular amino acid is the basis of genetic code. Using the genetic code biological information is transferred from DNA to protein. Sixty one codons of the genetic code encode 20 standard amino acids but only three codons act as stop codons. Due to degeneracy of the genetic code, some amino acids are encoded by more than one codon. The level of degeneracy varies from two to six and the synonymous codons generally differ in the 3rd position (wobble position). Some amino acids are encoded by only one codon (met, trp). Synonymous codons are those which code for the same amino acid. The phenomenon of non uniform usage of codons in which some codons are more preferred than others is the codon usage bias. It is mainly caused by mutation and translational selection [4][5][6][7]. Optimal codons are the more frequently used codons whereas non-optimal codons are the less frequently used codons. Non optimal codons usually correspond to less abundant tRNA molecules in the cell than optimal codons do [8][9][10][11][12] and the pause of translational machinery there [13]. For efficiency in translation process, the optimal codons may be selected for and are used frequently in highly expressed genes [14][15]. Several factors which affect the codon usage bias in an organism are gene expression level, composition bias (%GC content and GC-skew), gene length, RNA stability and recombination rate [16][17][18]. The objective of the present study is to analyze the codon usage bias in the available nuclear genes of one-horned rhino as no literature is available on this as-

pect in rhino.

Table 1. Correlation coefficient between overall nucleotide and the nucleotide at 3rd codon position in rhino

Nucleotide (%)	A3 %	T3 %	G3 %	C3 %	GC3 %
A %	0.964**	0.417	-0.832*	-0.688	-0.846*
T %	0.308	0.975**	0.651	-0.723	-0.753
G %	-0.825*	-0.603	-0.970**	0.546	0.865*
C %	-0.759	-0.706	0.689	0.951**	0.884*
GC %	-0.868*	-0.719	0.908*	0.824*	0.959*

* $p=0.05$, ** $p=0.01$, respectively

Materials and Methods

Sequence data

The complete coding sequences (CDS) of six genes (only these are available in database) encoding the proteins (viz. estrogen receptor alpha, estrogen receptor beta, CD28, HFE protein, gamma-interferon and interleukin-4) in Indian one-horned rhino (*Rhinoceros unicornis*) were retrieved from the National Centre for Biotechnology Information (<http://www.ncbi.nlm.nih.gov/Genbank/>). The cds sequences having exact multiple of three bases are considered for this analysis using the standard genetic code.

Compositional properties

Overall nucleotide composition (A, C, T and G%) and nucleotide composition at the wobble position of each codon (A3, C3, T3 and G3%) were analyzed for one-horned rhino coding sequences using an in-house perl script developed by S. Chakraborty. The GC and GC3 indices referred to the overall GC content and that at the wobble position of synonymous codons in the cds sequences (excluding the codons encoding single amino acids met and trp and the three codons responsible for polypeptide termination).

Measures of synonymous codon usage bias

A couple of indices for measuring the codon usage bias in genes have been proposed by several workers. Some of the most widely

used measures for measuring the synonymous codon usage bias are discussed here.

Relative Synonymous Codon Usage (RSCU)

Relative synonymous codon usage (RSCU) is analysed as the ratio of the observed frequency of a codon to the expected frequency if all the synonymous codons of a particular amino acid are used equally. If the RSCU value of a codon is greater than 1.0, it indicates that the particular codon is used more frequently than the expected frequency whereas the reverse is true for RSCU values less than 1.0 [14].

$$RSCU_{ij} = \frac{X_{ij}}{\frac{1}{n_i} \sum_{j=1}^{n_i} X_{ij}}$$

where X_{ij} is the frequency of occurrence of the j^{th} codon for i^{th} amino acid (any X_{ij} with a value of zero is arbitrarily assigned a value of 0.5) and n_i is the number of codons for the i^{th} amino acid (i^{th} codon family).

Effective Number of Codons (ENC)

The effective number of codons used by a gene (ENC) is generally used to measure the codon usage bias of synonymous codons [19]. The value of ENC ranges from 20 (for each amino acid, when only one codon is used) to 61 (when all synonymous codons for an amino are randomly used). If the calculated ENC value is greater than 61 (because codon usage is more evenly distributed than expected), it is adjusted to 61.

$$ENC = 2 + \frac{9}{F_2} + \frac{1}{F_3} + \frac{5}{F_4} + \frac{3}{F_6}$$

Codon Adaptation Index (CAI)

The codon adaptation index (CAI) [28] is the most extensively used measure of codon bias in prokaryotes [20][21][22] and in eukaryotes [23][24][25]. CAI is widely used as a measure of gene expression. CAI is based on relative adaptiveness of codons. The relative adaptiveness (ω) of a codon is the ratio of the usage of each codon divided by the most abundant codon within the same synonymous family. CAI value ranges from 0 to 1, with higher values indicating a higher proportion of the most abundant codons [26]. The analysis for codon usage bias was done by using an in-house perl script developed by S. Chakraborty.

Statistical analysis

Correlation analysis was performed to identify the relationship between overall nucleotide composition and each base at 3rd codon position and correlation of ENC with GC, GC₃ and CAI. All the statistical analyses were done using the SPSS software.

Results

Compositional properties

In the coding sequences of genes in one-horned rhino, it was found that the nucleotides, A and C occurred more frequently than G and T in the overall composition. The base C occurred most frequently (average C₃ % = 33.77) and A occurred least frequently at the 3rd codon position (average A₃ % = 16.20) as shown in Fig 1. The nucleotide composition in general and that at 3rd codon position for nuclear genes in one-horned rhino suggests that compositional constraints might be influencing the pattern of codon usage bias in the coding sequences of rhino. The overall GC % of rhino nuclear genes ranged from 41.3 to 57.2 % with an average of 51.46 % but the GC₃ % of nuclear genes ranged from 53.3 to 73.6 % with an average value of 65.2%. These differences could be attributed to the increased number of C at 3rd codon position in all the cds.

Table 2. Correlation coefficients between parameters of codon usage bias in rhino

No.	Correlation between parameters	Correlation coefficient value
1	ENC and GC	-0.934**
2	ENC and GC ₃	-0.979**
3	GC ₁ and GC ₃	0.817*
4	GC ₂ and GC ₃	0.799
5	GC ₁₂ and GC ₃	0.896*

* p=0.05, ** p=0.01, respectively

ENC values of rhino cds varied from 50 to 60 with a mean of 55 and were usually of higher magnitude. This indicates that the codon usage bias is possibly not remarkable in rhino and is maintained apparently at a stable level.

Codon Usage in Rhino

The overall RSCU values for 59 codons in rhino cds indicate that G and C occurred most frequently at the 3rd codon position as shown in Fig. 3. Fourteen codons end with C and these codons are TCC, AGC, TTC, CTC, TAC, TGC, CCC, CAC, ATC, ACC, AAC, GAC, GCC, GGC encoding the amino acid serine, serine, phenylalanine, leucine, tyrosine, cysteine, proline, histidine, isoleucine, threonine, asparagine, aspartate, alanine and glycine, respectively. Seven codons end with G and these are CTG, CAG, CGG, AGG, GTG, AAG, GAG encoding leucine, glutamine, arginine, arginine, valine, lysine and glutamate, respectively. Four codons ending with A nucleotide are CCA, AGA, GCA, GGA which encode the amino acid proline, arginine, Alanine and glycine, respectively. Codon ending in T is ATT which encodes the amino acid isoleucine.

For the rhino cds under study, we plotted the ENC value against overall GC, GC₃ content and found significant negative correlation of ENC with GC ($r = -0.934^{**}$, $p < 0.01$), ENC with GC₃ ($r = -0.979^{**}$, $p < 0.01$). In addition, we observed a significant positive correlation between GC₁ and GC₃ ($r = 0.817^{*}$, $p < 0.05$) and between GC₁₂ and GC₃ ($r = 0.896^{*}$, $p < 0.05$). These results suggest that nucleotide constraint under mutation pressure is the major factor for codon usage bias in these coding sequences of one-horned rhino. No significant correlation between ENC and CAI was observed in the present study.

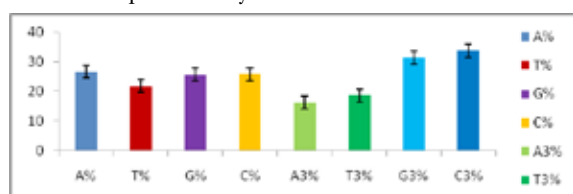


Fig. 1 Distribution of overall nucleotides (A, T, G, C%) and nucleotides at 3rd codon position (A₃, T₃, G₃, C₃) in coding sequences of rhino

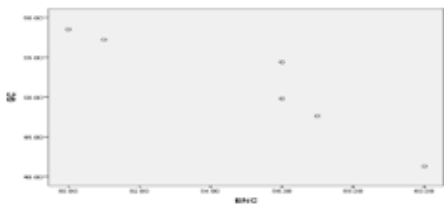
Effect of Mutation pressure on codon usage bias

To test the hypothesis that mutation pressure has no role in the evolution of codon usage bias in rhino, we compared the correlation coefficient between overall nucleotide composition A, T, G, and C% and its composition at 3rd codon position A₃, T₃, G₃ and C₃ % using Karl Pearson's correlation coefficient. Significant positive correlation was found between A and A₃ ($r = 0.964^{**}$, $p < 0.01$), C and C₃ ($r = 0.951^{**}$, $p < 0.001$), GC and GC₃ ($r = 0.959^{**}$, $p < 0.001$) whereas significant negative correlation was observed for other nucleotides. This suggests that compositional constraint under mutation pressure might affect the codon usage pattern in rhino. However, significant positive correlation was found between T and T₃ ($r = 0.975^{**}$, $p < 0.001$) but significant negative correlation between G and G₃ ($r = -0.970^{**}$, $p < 0.001$). Moreover, no significant correlation was found between G and C₃ ($r = 0.546$,

[illegible]

Discussion

Genetic analysis of endangered species such as Indian one-horned rhino is useful not only for acquiring knowledge on the biology of animals but also for enhancing the conservation strategies [27][28]. Different types of DNA repeats are found in higher eukaryotic genomes [29]. For the development of desired genetic marker, information about the organization of satellite DNA and VNTR loci of species and subspecies of rhino would be relevant [30].



We analyzed the pattern of codon usage bias in the nuclear genes of *Rhinoceros unicornis* L. Most frequent codons had C or G at its 3rd codon position. Codon usage bias in the analyzed cds is not remarkable as the ENC values are high. From this study, the overall nucleotide composition and the composition at 3rd codon position are shown in Table 1. Our analysis suggests that nucleotide composition in rhino could be under mutation pressure. Highly significant correlation between GC₁ and GC₂ and between GC₁₂ and GC₃ suggest that mutation pressure is an important factor in shaping the codon usage in this species. Further, highly significant correlation between ENC with GC and ENC with GC₃ support the hypothesis that compositional constraint under mutation pressure has influenced the codon usage bias. From our earlier work on mitochondrial DNA of mammals, we have found the same result but the only difference is that mitochondrial DNA in mammals prefers AT richness [31].

The present study is the first endeavor on the codon usage analysis in Indian one-horned rhino (*Rhinoceros unicornis* L). We have found that codon usage bias in rhino is not very remarkable. Codon usage bias prefers G or C nucleotide at the 3rd position of codons. Most frequent codons in the cds end with C at 3rd codon position. Our results support the fact that both nucleotide constraint and compositional constraint might have played a major role in the codon usage bias in the nuclear genes of Indian one-horned rhino.

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Coding sequence (CDS), Effective number of codon (ENC), Mitochondrial DNA (mt DNA)

We do not have any conflict of interest.

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