Mutational Bias and Translational Selection Shaping the Codon Usage Pattern of Tissue-Specific Genes in Rice

Qingpo Liu*

College of Agriculture and Food Science, Zhejiang A & F University, Lin’an, People’s Republic of China

Abstract

The regulatory mechanisms of determining which genes specifically expressed in which tissues are still not fully elucidated, especially in plants. Using internal correspondence analysis, I first establish that tissue-specific genes exhibit significantly different synonymous codon usage in rice, although this effect is weak. The variability of synonymous codon usage between tissues accounts for 5.62% of the total codon usage variability, which has mainly arisen from the neutral evolutionary forces, such as GC content variation among tissues. Moreover, tissue-specific genes are under differential selective constraints, inferring that natural selection also contributes to the codon usage divergence between tissues. These findings may add further evidence in understanding the differentiation and regulation of tissue-specific gene products in plants.

Introduction

Within a genome, some synonymous codons are avoided readily, while some are used preferentially. Such biases are ubiquitously present in eukaryotes and prokaryotes. The processes of shaping codon usage in a given genome are very complicated, which may be primarily attributed to natural selection and neutral evolutionary forces, such as biased mutation and gene conversion. In most cases, codon usage can reflect a balance between natural selection and neutral evolution [1]. However, in some multicellular species, translational selection serves as the major determinant in shaping codon usage [2–3]. In contrast, in mammals, especially in human, the synonymous codon usage is mainly arisen from neutral processes rather than selection [4].

Interestingly, significant differences in synonymous codon usage were observed between genes selectively expressed in six adult human tissues; this tissue-specific variation of synonymous codon usage is probably not only due to translational selection but isochores structures [5]. Sémon et al. [6] confirmed the significant (but weak) difference of tissue-specific codon usage; however, they found no evidence for tissue-specific adaptation of synonymous codon usage in human [6]. In addition, it was reported that in plants and mammals tissue-specific genes are under weak selective constraints, and evolve more quickly than housekeeping genes [7–8]. Thus, different evolutionary forces must underline the evolution of synonymous codon usage of housekeeping and tissue-specific genes.

To our knowledge, whether there exists significant codon usage variation and to what extent of these differences between tissue-specifically expressed genes in plants have not been explored yet. Rice is very heterogeneous in base composition, and has an isochores structure similar to mammals [9]. It is thus intriguing to understand how about the codon usage of tissue-specific genes in rice. Particularly, although base compositional mutation bias and natural selection were found to have played crucial roles in determining the codon usage of all rice genes [10], whether the tissue-specific codon usage was affected by both factors remains unclear.

In this study, the tissue-specific codon usage in rice was evaluated using a multivariate method, internal correspondence analysis [11]. These analyses reveal a significant difference in synonymous codon usage between genes selectively expressed in different tissues. As observed in the human genome, however, this effect is weak, which represents only 5.62% of the total codon usage variability. Importantly, the GC-content variation between tissues primarily contributes to, and translational selection may play a relative weaker role in shaping the tissue-specific synonymous codon usage variability in rice.

Materials and Methods

Sequence Data

The rice (Oryza sativa) protein-coding sequences (CDSs) and expressed sequence tags (ESTs) were retrieved from the rice genomic resource [MSU pseudomolecule v7.0; ftp://ftp.plantbiology.msu.edu/pub/data/Eukaryotic_Projects/o_sativa/annotation_dbs/pseudomolecules] and GenBank database [release 189, April 2012], respectively. If there were several alternative splicing variants, the longest CDS was chosen as the representative for that gene. The total dataset contains 56,591 rice CDSs.

Gene expression patterns were estimated using EST data. A total of 1,252,989 rice ESTs were downloaded, in which ESTs belonging to cDNA libraries from suspension cell culture, callus, pooled or mix organs, or unidentified tissues were excluded. CDSs
were then used as query to search against the EST data using MEGABLAST [12] with an E-value cutoff 10−20. A sequence match is counted if MEGABLAST alignment shows at least 95% identity over 300 nt or more.

The tissue expression pattern of rice genes was also examined by analyzing the published micro-array data (GSE7951, and GSE11966; [13–14]), where the gene expression profiles in nine tissues including root, shoot, leaf, anther, embryo, endosperm, 5d-seed, and 64 51.718

were calculated using CodonW v1.4 [23] and custom PERL programs, respectively. The internal correspondence analysis [20], Spearman correlation, and ANOVA analysis were all performed using the R software v2.15 [21].

Identification of Orthologous Gene Pairs

The amino acids, CDS, and EST sequences of *Sorghum bicolor* and *Brachypodium distachyon* were downloaded from the Phytozome (version 8.0; http://www.phytozome.net/) and GenBank databases (release 189, April 2012), respectively. Totally, 208,841 and 128,092 *S. bicolor* and *B. distachyon* ESTs were downloaded, which were then rearranged as that described in rice. The orthologous gene pairs of rice, and *S. bicolor*, and *B. distachyon* were obtained from the PhylomeDB (v3.0; [24]) and MetaPhOrs [25] databases, where the orthologs were identified using the phylogeny-based approaches [24–26] instead of simple BLAST search [27]. The tissue- and non-tissue-specific genes identified in rice were used as query to search against the *S. bicolor* and *B. distachyon* sequences to identify their orthologous counterparts in each of the two species, using a custom PERL program. In order to efficiently distinguish the tissue-specific and non-tissue-specific genes in *S. bicolor* and *B. distachyon*, the ESTs originated from rice and the two plant species were mixed together to construct an EST sequences pool, considering that the three species are phylogenetically close to each other. After that, the identified orthologous genes in *S. bicolor* and *B. distachyon* were separately searched against the EST sequences pool using MEGABLAST [12]. The tissue expression pattern of a given gene was inferred based on their EST profiling. Here, if its transcript is detected in only one tissue, the corresponding gene was referred to be tissue-specific. On the contrary, a gene was regarded to be non-tissue-specific, if it was expressed in two tissues or more but not significantly abundance in any tissues. The amino acid sequences of orthologs were aligned using MAFFT v6.6 [28], and then the codon-alignments of CDS sequences were generated based on the resulting amino acid alignments using a custom PERL program. The Yang and Nielsen method [29] implemented in the yn00 program of the PAML v4.4 package [30] was employed to calculate the pair-wise synonymous (dS) and non-synonymous (dN) distance between the orthologous genes of rice and *S. bicolor* and *B. distachyon*.
Results and Discussion

Variation in Synonymous Codon Usage between Tissues

The data set consisting of 675 CDSs from seven tissues were subjected to CodonW and custom PERL programs to investigate the synonymous codon usage of rice tissues. From Table 1, it is obvious that the average ENC value in one tissue is significantly different from the ones in other tissues. The similar trend can be observed by analyzing the CAI values, another commonly used measure of codon bias. In addition, ENC values are significantly negatively correlated with CAI values (Spearman’s correlation coefficient $r = -0.964$, $p = 0.003$), a result that is consistent with those in analyzing all rice genes [10]. It is worthy to note that the average ENC and CAI values of genes selectively expressed in 5d-seed are significantly greater and smaller than that of other tissues, indicating that 5d-seed-tissue-specific genes may tend to select synonymous codons randomly. The reverse case was found in embryo, where the average ENC value is significantly smaller than other tissues, indicative of stronger codon usage bias. These observations suggest that remarkable variation in synonymous codon usage should have occurred between different tissues in rice. Compared with the genes that are tissue-specific, non-tissue-specific genes have the weakest codon usage bias, and are usually expressed at a lower level (Table 1).

However, ENC only measures the overall departure from random synonymous codon choice, and hence which codons are used more frequent than others can not be clearly presented. As a result, two genes exhibiting the same degree of overall bias (ENC value) may actually differ dramatically in their particular choice of synonymous codons [5]. To cope with this problem, Plotkin et al. [5] developed a new method to measure the distance between two genes by counting the number of amino acids that exhibit significantly different codon usage. However, this method is also defective because it is sensitive to the length and the amino acid composition of proteins [6]. Thus, for this study, the internal correspondence analysis (ICA) was adopted to further investigate the variability of synonymous codon usage among tissue-specific genes in different tissues.

In ICA, the total codon usage variability was decomposed into within-block and between-block variability, consisting of the amino acid usage (between-AA) variability ($b$, $e$, $h$), synonymous codon usage (within-AA) variability ($a$, $d$, $g$), and variability of between- ($d$, $e$, $f$) or within tissues ($a$, $b$, $c$). The decomposition of global variability yields nine elementary analyses ($a$–$i$). In each peculiar analysis, the contribution to the total variance is indicated, where only the first 10 eigenvalues are represented to allow for a direct visual comparison.

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Figure 1. Internal correspondence analysis of tissue-specific genes in rice. The global codon usage variability was decomposed into within-block and between-block variability, consisting of the amino acid usage (between-AA) variability ($b$, $e$, $h$), synonymous codon usage (within-AA) variability ($a$, $d$, $g$), and variability of between- ($d$, $e$, $f$) or within tissues ($a$, $b$, $c$). The decomposition of global variability yields nine elementary analyses ($a$–$i$). In each peculiar analysis, the contribution to the total variance is indicated, where only the first 10 eigenvalues are represented to allow for a direct visual comparison.
variability are due to variability in synonymous codon usage (within-AA; Fig. 1g) and to variability in amino acid usage (between-AA; Fig. 1h), respectively. However, the effect of variability in synonymous codon usage between tissues is much smaller as compared to other sources of variability, which only accounts for 5.62% of the total variability in codon usage (Fig. 1i).

In order to test whether the observed small proportion of variability in synonymous codon usage between tissues (Fig. 1i) is caused just by chance, 1,000 independent permutations were performed by randomly associating the genes and tissues, and the ICA was repeated accordingly. It was found that the observed value (5.62%) is significantly greater than that obtained by chance ($p < 0.001$), indicating that tissue-specific genes are indeed significantly different in synonymous codon usage that depends on the tissue where they are expressed.

As reported, the identification of tissue-specific genes is always determinant with the sensitivity of the method employed to detect expression [6]. To test this hypothesis, two groups of new datasets were presented. One is composed of genes that are only detected in one of the seven tissues by EST profiling. After running the ICA on this new dataset, I found that the fraction of variability of synonymous codon usage between tissues decreases up to 3.6%. Alternatively, if the tissue specificity was defined based on the definition proposed by Wang et al. [15], only 0.8% of total variability could be attributed to the variation in synonymous codon usage between tissues. However, using a combination of the two methods, this fraction of value climbs up to 5.62%, which is nearly 2.5 times of that reported in human (2.3%, [6]), where the tissue-specific genes were defined only based on the EST abundance of transripts. As known, EST sequences are often used to estimate the expression profiling of protein-coding genes. However, the sensitivity of this method always depends on cDNA libraries’ origin, EST data quality as well as the numbers of EST sequences in each library, which can greatly influence the final annotation of tissue-specific genes [31]. In addition, if a gene has a SPM value close to 1, as proposed by Wang et al. [15], the corresponding gene can be designated as tissue-specific. According to the definition, the SPM value >0.9 is used as empirical cutoff in classifying tissue-specific genes [15]. In this regard, this method is also defective, because in some cases genes that are expressed at particularly high levels in a given tissue but also expressed at relatively lower levels in other tissues are also considered as tissue-specific. In this study, the tissue-specific genes identified by the two methods were integrated effectively, thus the obtained results should be more robust than simply adopting either EST profiling or Specificity Measure (SPM).

**Table 2. Base composition of coding sequences of tissue- and non-tissue-specific genes in rice.**

| Tissue   | No. Seqs | GC1         | GC2         | GC3         | GC all         |
|----------|----------|-------------|-------------|-------------|----------------|
| Root     | 49       | 0.59±0.010  | 0.48±0.013 a | 0.73±0.030 a | 0.60±0.014 a   |
| Shoot    | 33       | 0.59±0.015  | 0.47±0.014 a | 0.70±0.040 a | 0.59±0.016 a   |
| Leaf     | 356      | 0.55±0.005  | 0.43±0.005 cd| 0.57±0.010 b | 0.52±0.006 b   |
| Anther   | 108      | 0.58±0.007  | 0.42±0.007 cd| 0.71±0.019 a | 0.58±0.010 a   |
| Embryo   | 29       | 0.58±0.014  | 0.45±0.016 cd| 0.69±0.033 a | 0.57±0.018 a   |
| Endosperm| 36       | 0.56±0.013  | 0.41±0.012 e | 0.50±0.030 bc| 0.50±0.017 bc   |
| 3d-seed  | 64       | 0.53±0.008  | 0.46±0.008 e | 0.51±0.019 cd| 0.48±0.010 c   |
| Non-specific | 1187      | 0.54±0.001  | 0.42±0.001 de| 0.46±0.003 d | 0.47±0.002 c   |

| Tissue   | No. Seqs | A        | T        | C        | G        |
|----------|----------|----------|----------|----------|----------|
| Root     | 49       | 0.21±0.002 f | 0.18±0.001 g | 0.29±0.002 b | 0.30±0.001 a |
| Shoot    | 33       | 0.20±0.003 g | 0.20±0.002 e | 0.29±0.003 a | 0.29±0.003 b |
| Leaf     | 356      | 0.24±0.000 c | 0.23±0.000 c | 0.24±0.000 e | 0.27±0.000 c |
| Anther   | 108      | 0.25±0.002 e | 0.19±0.001 f | 0.28±0.001 c | 0.29±0.001 b |
| Embryo   | 29       | 0.23±0.003 d | 0.18±0.002 h | 0.28±0.003 d | 0.29±0.002 b |
| Endosperm| 36       | 0.27±0.004 a | 0.22±0.002 d | 0.23±0.004 f | 0.27±0.002 c |
| 3d-seed  | 64       | 0.26±0.001 b | 0.25±0.001 a | 0.21±0.001 h | 0.26±0.002 d |
| Non-specific | 1187      | 0.27±0.000 a | 0.25±0.000 b | 0.22±0.000 g | 0.25±0.000 e |

*Note: Data are reported as means ± SD. Within a column, mean values followed by different letters (a, b, c, d, e, f, and g) mean significant difference at the 0.05 level ($p<0.05$).*
Figure 2. Illustration of GC3 content variation for genes selectively expressed in different tissues. For each tissue, the distribution of GC3 content is represented by a boxplot, where the lower, middle, and top horizontal lines of the boxes represent the 25%, 50%, and 75% quantiles, respectively. Initials of tissue names are indicated: se, seed; le, leaf; en, endosperm; sh, shoot; an, anther; ro, root; em, embryo.

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Figure 3. Comparison of synonymous $d_s$ (a) and nonsynonymous substitution rates $d_N$ (b) between rice and B. distachyon, and sorghum orthologous gene pairs for tissue-specific and non-tissue-specific classes of genes. The symbol ** indicates significant difference at the 0.01 level ($p<0.01$).

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To test whether the observed significant difference in synonymous codon usage between tissues can be solely explained by the differences in base compositions, the ICA analysis of within-AA and between-tissues variability in codon usage (Fig. 1d) was analyzed, where a single major trend in codon usage was identified: the first axis may account for 83.1% of the total variation in synonymous codon usage between tissues. Thus, the first axis can be used as the major factor explaining the variation in codon usage between tissues. It was observed that Axis 1 coordinates are strongly negatively correlated with GC, and GC3 content (Spearman’s correlation coefficients $r = -0.964$, $p = 0.0028$; and $r = -0.929$, $p = 0.0067$, respectively). Particularly, the GC3 content of tissue-specific genes varies extensively, and the variation of GC3 content between genes within a given tissue is even much larger than the variability between tissues (Fig. 2). Thus, the weak differences of GC content between genes specifically expressed in different tissues mainly contribute to the observed differences of synonymous codon usage between tissues.

In addition, a significantly positive Spearman correlation between Axis 1 coordinates and intronic GC content was found ($r = 0.955$, $p = 0.0008$). Under mutational bias alone, the correlations between Axis 1 and exonic and intronic GC contents should be similar [3]. Thus the observed inverse correlations may indicate that natural selection should have played a role in shaping the distinct synonymous codon usage in different tissues. This was confirmed by the observations that CAI values are significantly positively correlated with GC3, but not with intronic GC content in all tested tissues (data not shown). If tissue-specific genes were under selective pressure to optimize translation, it expects that introns would be not influenced by this effect [6]. Notwithstanding, no significant correlation between Axis 1 coordinates and CAI values was found (Spearman’s correlation coefficient $r = -0.75$, $p = 0.066$). Overall, the results indicate that the tissue-specific variability of synonymous codon usage is mainly due to base compositional mutation bias, while translational selection contributes to, with a relative smaller fraction, in shaping this phenomenon. Thus, although rice and human share a similar isochore genomic structure [9], the features of codons that are responsible for the tissue specificity of synonymous codon usage in rice are not totally the same as that in the human genome [6].

**Selective Constraints on Tissue- and Non-tissue-specific Genes**

The analyses of tissue-specific genes reveal that the evolution of synonymous codon usage of a given gene may essentially depend on the genomic environment where it is located [6,32]. Because weak tissue-specific variation of GC3 content exists, I would expect that tissue-specifically expressed genes may be strongly biased to cluster into GC-rich or GC-poor regions along the rice chromosomes. However, like in the human genome, this study did not find any significant clustering of tissue-specific genes along rice chromosomes. Conversely, these genes are distributed throughout the genome (see Fig. S1). Actually, housekeeping but not tissue-specific genes are prone to cluster in chromosomal regions [33]. Moreover, the genes that are selectively expressed in different tissues have near similar distributions of CDS lengths (Table 1).

**Supporting Information**

**Figure S1** Distribution of tissue-specific genes onto rice chromosomes. The tissue names were used to represent the genes specifically expressed in that tissue. (PDF)

**Table S1** EST abundance, CAI values, and functional annotation of 71 highly expressed genes in rice. (DOC)

**Author Contributions**

Conceived and designed the experiments: QL. Performed the experiments: QL. Analyzed the data: QL. Contributed reagents/materials/analysis tools: QL. Wrote the paper: QL. 

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