

Evaluation of Codon Biology in *Citrus* and *Poncirus trifoliata* Based on Genomic Features and Frame Corrected Expressed Sequence Tags

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Abstract

Citrus, as one of the globally important fruit trees, has been an object of interest for understanding genetics and evolutionary process in fruit crops. Meta-analyses of 19 *Citrus* species, including 4 globally and economically important *Citrus sinensis*, *Citrus clementina*, *Citrus reticulata*, and 1 *Citrus* relative *Poncirus trifoliata*, were performed. We observed that codons ending with A- or T- at the wobble position were preferred in contrast to C- or G- ending codons, indicating a close association with AT richness of *Citrus* species and *P. trifoliata*. The present study postulates a large repertoire of a set of optimal codons for the *Citrus* genus and *P. trifoliata* and demonstrates that GCT and GGT are evolutionary conserved optimal codons. Our observation suggested that mutational bias is the dominating force in shaping the codon usage bias (CUB) in *Citrus* and *P. trifoliata*. Correspondence analysis (COA) revealed that the principal axis [axis 1; COA/relative synonymous codon usage (RSCU)] contributes only a minor portion (~10.96%) of the recorded variance. In all analysed species, except *P. trifoliata*, Gravy and aromaticity played minor roles in resolving CUB. Compositional constraints were found to be strongly associated with the amino acid signatures in *Citrus* species and *P. trifoliata*. Our present analysis postulates compositional constraints in *Citrus* species and *P. trifoliata* and plausible role of the stress with GC₃ and co-evolution pattern of amino acid.

Key words: *Citrus*; coevolution; mutational bias; *Poncirus trifoliata*; stress; GC₃ biology

1. Introduction

Genome composition (such as GC- and AT-content) and subsequent balance of codon usage and eukaryotic translation machinery play an important role in the evolution of the nucleotides at the wobble position.¹ Degeneracy in biased codon usage explains the concept behind the usage of same amino acids with multiple synonymous codons except methionine (Met) and tryptophan (Trp).² This genome composition bias leads to the usage of some codons at a higher frequency as compared to the synonymous

codons for encoding the particular amino acid. These differences in the usage of the synonymous codons have also been one of the factors for the evolution of proteome diversity and can help us to understand the evolution of those proteins that have structural differences in spite of being conserved at the sequence level.^{3–7}

Two major paradigms of codon usage were proposed as plausible answers to clarify the non-randomness of codon usage at intra- and inter-species levels: (i) natural selection that is expected to yield a correlation with codon bias in highly expressed genes,⁸

rapidly regulated and variably expressed genes, and (ii) neutral processes, such as mutational biases (MBs), where some mutations occur more often than others across the genome of an organism because of local variations in the base composition. Several lines of evidence support the argument that the usage of synonymous codons with unequal frequencies in both prokaryotic and eukaryotic genes is the result of a complex balance between MB and/or natural selection and genetic drift.^{9–12} It has been suggested that the codon bias could be positively selected because of a more efficient and accurate translation, and favoured codons may correspond to the most highly expressed genes.^{2,13} In addition to neutral and selection processes, GC-biased gene conversion, which depends on the local recombination rate, is an important factor in shaping codon and amino acid usage.¹⁴

To date, wide variations have been observed in codon usage patterns in many organisms and have provided clues to understand the evolution of genes and gene families. The factors that potentially affect biased usage of codons are MB that correlates the codon usage bias (CUB) with the genomic GC content, Hill–Robertson effect explaining the interference of selection of one locus with another locus, translational selection, and a cumulative effect of replication and translational selection in shaping the codon usage across the genes of several bacterial species.^{15–19} It has been shown that CUB and protein functional conservation play a major role for the decelerated evolution of whole genome duplication in *Saccharomyces cerevisiae*.²⁰ Several other factors that might affect the codon usage are amino acid conservation and hydrophobicity,²¹ gene expression,²² mRNA folding stability, codon–anticodon interaction, and gene length.²²

Citrus is a diploid genus of the Rutaceae family, whose cultivated forms are important for human diet with more than 122 MT of annual world fruit production. We have systematically analysed codon usage patterns and genomic heterogeneity using a meta-analysis approach that involves multivariate tools and codon usage indices such as relative synonymous codon usage (RSCU) and effective number of codons (Nc) in the studied gene sets.^{23,24} In present study, we inferred global pattern of CUB, mutational pressures, and association of GC₃ with potential stress events. We have collectively analysed horticulturally important *Citrus* species that includes recently sequenced double haploid *Citrus sinensis* genome²⁵ and 18 additional *Citrus* species with expressed sequence tags (ESTs) counts of more than 1000 per species. To make this analysis comparative, we have also included *Poncirus trifoliata* that is considered to be a distant relative of *Citrus* genus. These

species show wide variation in traits such as cold and drought tolerance and are of commercial importance. We have restricted our analyses to *Citrus* genera to develop resource information for *Citrus* species.

Our study demonstrated that CUB in *Citrus* species and *P. trifoliata* is biased towards AT richness, and a relatively higher occurrence of A- and T-ending codons was observed. We found several interesting and diverse patterns of optimal codons, and it was observed that two optimal codons coding for Alanine and Glycine were evolutionarily conserved between the *Citrus* species and *P. trifoliata*. To the best of our knowledge, our analysis presents the first complete report on the identification of optimal codons across the entire *Citrus* genus and *P. trifoliata*, which could serve as the potential source for developing transgenic *Citrus* cultivars using codon optimization. GC₃ and evolution were correlated using Hamming distance parameter to identify suggestive evolutionary pairs of genes. Co-orthologous genes matrix identified using the alignment ratio suggested close association of *Citrus clementina*, *Citrus sinensis*, and *Citrus reticulata*, as they belong to the same phylogenetic clade. We further demonstrated that nucleotide bias has a genome-wide influence on amino acid composition of *Citrus* species and *P. trifoliata*.

2. Materials and methodology

2.1. Sequence information and processing

Our dataset consists of genome-predicted coding regions and ESTs. All the genome-predicted coding sequences of *C. sinensis* were retrieved from recently sequenced *Citrus* genome.²⁵ In case of other *Citrus* species, ESTs were downloaded from the National Center for Biotechnology Information (NCBI) EST repository (NCBI; <http://www.ncbi.nlm.nih.gov>). In addition, we downloaded putative unique transcripts for all studied species from the PlantGDB database (<http://www.plantgdb.org>). A detailed description of the data used is shown in Table 1. In case of ESTs, the ESTs were first clustered into contigs and singletons using CAP3 with default parameters.²⁶ A minimum match percentage cutoff of 95% for 40 overlapping bases was used to assign 2 sequences to a cluster.

2.2. Frame correction of ESTs

All the UniGenes (contigs + singletons) were analysed for frame correction and prediction of protein-coding region using FrameDP.^{27,28} Briefly, the following pipeline was implemented using FrameDP to identify open reading frames (ORFs): firstly, each EST was compared against the TAIR database (*Arabidopsis* Information Resource; <http://www.arabidopsis.org/>) using BLASTX²⁹ with E-value = 10^{-4} , identity

Table 1. Genomic composition of coding region of *Citrus* species and *P. trifoliata* at GC, GC₁, GC₂, GC₃, and GC_{3s}

No.	<i>Citrus</i> species	Genes/EST ^a	Unigenes count	Genes ^b	GC	GC ₁	GC ₂	GC ₃	GC _{3s}
1	<i>C. sinensis</i>	44 275	—	41362	43.92	50.4	40.33	41.03	38.8
2	<i>C. aurantifolia</i>	8219	7550	4715	47.78	52.96	43.87	46.52	44.73
3	<i>C. aurantium</i>	14 584	11 952	6787	46.93	52.47	42.59	45.72	43.94
4	<i>C. clementina</i>	118 365	51 591	33 765	47.04	51.84	42.51	46.76	44.99
5	<i>C. clementina</i> × <i>C. tangerina</i>	1843	1283	677	44.66	51.67	39.77	42.55	40.47
6	<i>C. jambhiri</i>	1017	858	701	45.86	51.79	41.13	44.66	42.67
7	<i>C. japonica</i> var. <i>margarita</i>	2924	1628	588	46.11	52.67	41.42	44.25	42.35
8	<i>C. limettoides</i>	8188	7933	3964	46.47	50.76	42.75	45.9	44.25
9	<i>C. limonia</i>	11 045	9761	4256	45.82	51.13	41.37	44.96	43.17
10	<i>C. medica</i>	1115	896	711	45.65	51.91	40.9	44.13	42.15
11	<i>C. reshni</i>	5768	3735	2644	45.2	51.51	40.57	43.52	41.5
12	<i>C. reticulata</i>	55 980	46 170	30 816	46.85	52.3	42.77	45.48	43.62
13	<i>C. reticulata</i> × <i>C. temple</i>	5823	3685	2253	46.48	52.41	41.64	45.41	43.41
14	<i>C. sinensis</i> × <i>P. trifoliata</i>	1837	1522	992	46.07	51.67	41.41	45.15	43.23
15	<i>C. sunki</i>	5216	4688	1746	47.58	51.51	43.58	47.64	46.17
16	<i>P. trifoliata</i>	62 695	35 740	19 445	46.78	52.33	42.67	45.35	43.52
17	<i>C. unshiu</i>	19 072	9289	4592	45.67	51.72	41.14	44.15	42.21
18	<i>C. paradisi</i>	8039	4621	2517	45.32	52.19	40.51	43.27	41.3
19	<i>C. paradisi</i> × <i>P. trifoliata</i>	7954	3335	2596	45.23	51.96	40.77	42.98	40.96

Means of GC% were calculated at the first, second, and third positions. GC_{3s} represents the GC at the third synonymous position.

^aIn *C. sinensis*, genome-predicted coding regions are used, whereas for the rest of the species, the number represents the EST count in the column (genes/EST).

^bSelected genes above 300 bp threshold.

percent (%) = 40% over 100 amino acids.²⁷ Secondly, the training dataset was generated from the BLASTX results, and, subsequently, the training matrix was calculated, which represents the coding style of the species. Thirdly, a collection of putative protein-coding sequences (CDSs) was generated for each *Citrus* species and *P. trifoliata* based on its homology with known protein dataset and on coding style recognition matrix.

2.3. Sequence filtering and GC variation

From the set of corrected sequences, we discarded proteins shorter than 100 amino acids to create a reliable dataset for all the studied *Citrus* species and *P. trifoliata* for further analysis.²³ The final sequence dataset was subsequently analysed by tabulating the frequency of GC at the first, second, and third codon positions (GC₁, GC₂, GC₃, and GC_{3s}, respectively) using in-house written Perl and C++ scripts. GC₃ is defined as the fraction of cytosines (C) and guanines (G) in the third position of the codon: $GC_3 = 3(C_3 + G_3)/L$ for the ORF of length L, whereas GC_{3s} is defined as G + C base composition at the third synonymously degenerate position of codons. To define

GC₃-rich and GC₃-poor groups, we have selected 5% of the genes with the highest and the lowest GC₃ values. For the genes in the GC₃-rich and-poor groups, we have computed two measures: (i) positional gradients of GC₃ and (ii) CG₃ skew that are defined as: CG₃ skew is the difference in fraction of cytosines (C) and guanines (G) in the third position of the codon divided by the sum of C and G in the third position: $CG_3\text{-skew} = (C_3 - G_3)/(C_3 + G_3)$ and positional gradient GC₃ as a $[G_3(x) + C_3(x)]/N_{seq}$, where x is the distance measured as number of codons from the first ATG, and Nseq is the number of sequences.

2.4. Indices of codon usage and correspondence analysis

The effective number of codons (Nc) provides an independent measure of CUB, regardless of the gene length.²³ The expected Nc values were computed according to the equation proposed by Wright, which assumes equal use of G and C (A and T) in degenerate codon groups.²³

$$Nc = 2 + s + \left\{ \frac{29}{[s^2 + (1 - s)^2]} \right\}, \text{ where } s = GC_{3s}.$$

We have further analysed RSCU using multivariate analysis for all the 59 informative codons (excluding Met, Trp, and the three stop codons).^{30,31} As proposed earlier, if RSCU values are close to 1.0, it indicates that all the synonymous codons are used equally without any bias towards the usage of a particular codon in a gene of length L . In our study, axis 1 (COA/RSCU) and axis 2 (COA/RSCU) represent the first and second major axes of correspondence analysis. The Kyte–Doolittle scale was used to calculate the hydropathy score that is the arithmetic mean of the sum of hydropathic indices of each amino acid. This scale also provides information on transmembrane or surface regions.³²

2.5. Identification of optimal codons

Optimal codons are believed to achieve faster translation rates and high accuracy, therefore the effect is more pronounced in highly expressed genes.³³ For the identification of optimal codons, we used 10% of total genes from extreme ends of the principal axis of correspondence analysis (axis 1; COA/RSCU). Codon usage was then compared using χ^2 contingency (χ^2) of the two groups, and codons whose frequency of usage was significantly higher at three different levels of statistical precision (P -value < 0.5 ; P -value < 0.01 ; P -value < 0.001) in highly expressed genes when compared with lowly expressed genes were defined as the optimal codons. We also identified the genes related to the ribosomal proteins and observed the association of the ribosomal proteins with axis 1 (COA/RSCU) and axis 2 (COA/RSCU) of the correspondence analysis to optimize the identification of the optimal codons.

2.6. GO and stress-associated annotation, Hamming parameter, and coevolution of amino acid composition

Stress-related genes were identified using reciprocal best bidirectional hits (RBH) using NCBI BLASTP program with E-value 10^{-30} , and gene ontology (GO) annotation of the model dicot plant *Arabidopsis thaliana* was then used by ‘guilt-by-association’ approach. To optimize our annotation pipeline, we selected GO annotations ‘involved in’ the category containing word ‘stress’ and/or ‘response’ and grouped together as ‘stress related’. The normalized Hamming distance was then evaluated according to the method described in Sablok et al.³⁴ All the frame-corrected coding regions and the genome-predicted coding regions were parsed. Subsequent proteins were used to identify co-orthologous genes, and a suggestive phylogeny was drawn using the co-orthologous matrix as described in Lechner et al.³⁵ To identify the effect of nucleotide bias on amino

acid composition, we partitioned the codons according to GC-rich (the so-called GARP amino acids: Glycine, Alanine, Arginine, and Proline) and AT-rich, FYMINK amino acids (Phenylalanine, Tyrosine, Methionine, Isoleucine, Asparagine, and Lysine) amino acids.^{36,37} We have maintained the exclusion of Leucine and Arginine from the dataset as per Singer and Hickey.³⁷

2.7. Statistical analysis

All the indices of codon usage were calculated using CodonW (<http://codonw.sourceforge.net>), and several in-house developed Perl, R, and C++ scripts were written to streamline the downstream analyses. Statistical analyses were performed using R in R studio (<http://rstudio.org/>). All the results were interpreted based on the non-parametric Spearman’s rank correlation (ρ).

3. Results and discussion

3.1. Patterns of genomic content and codon usage variation across *Citrus* species and *P. trifoliata*

The recently sequenced nuclear genome of *C. sinensis*²⁵ presents an opportunity to analyse the nucleotide compositional pressure and heterogenetic variation, which could possibly help us to understand the molecular adaptation of these horticulturally important fruit species. We have used meta-analyses approach using the predicted genomic coding regions of recently sequenced *C. sinensis* and frame-corrected ESTs of related species and *P. trifoliata*. Nucleotide composition analysis indicated that *Citrus* species are AT rich ($\sim 46\%$ GC, typical for many sequenced genomes of dicot species), and this bias towards the AT richness is dominant and is observed across all the studied species (Table 1).

Based on the above observation, we could hypothesize that these species are biased towards the A- and/or T-ending codons in the coding region across these two genera. Generally, most of the analysed dicot species prefer to use A- or T-ending codons at the third position; this observation is in accordance with a previous study in *Citrus* using a small dataset of 177 CDS regions.³⁸ We found that average GC₃ is significantly lower than GC₁ and the observed GC (P -value < 0.05) which potentially explains that the genome composition is biased towards the dominating AT usage encoding genes at the third, ‘wobble’ position.

In eukaryotes, the intra-genomic heterogeneity is high, and interspecific variation of the average GC content is low.^{39,40} In *Citrus* species, GC usage varies by position, but with much greater variance, with higher usage in position 1 ($\sim 51.8\%$ GC) and lower

GC usage in positions 2 and 3 (~41.6 versus ~44.7% and also at the third synonymous positions GC_{3s} ~42.8%). The observed results are consistent with the previous reports demonstrating preference for G in position 1 and T/A in positions 2 and 3⁴¹ and demonstrate the role of mutational pressure in the evolution of CUB across *Citrus* species and *P. trifoliata*. Furthermore, there is a significant wide variation in GC usage at the third synonymous position (GC_{3s} ~42.8%) (Table 1; Fig. 1). In our analysis, we estimated codon usage patterns of frame-corrected coding regions obtained from ESTs versus gene predictions derived from a fully assembled and annotated genome.

To look for biased association between the nucleotide content and codon usage, we plotted Nc against GC_{3s} , also described as the Nc plot (Fig. 2) that has been widely demonstrated as an important parameter to evaluate codon usage variation among genes, as this codon usage index has a definite relationship with the GC_3 (more compositionally biased DNA is expected to encode a smaller subset of codons).^{23,42} Wright²³ argued that the comparison of actual distribution of genes, with expected distribution under no selection could be indicative, if CUB of genes has some other influences other than compositional constraints. A significant correlation was found between Nc and GC_{3s} (0.466** *C. sinensis*; -0.025** *Citrus aurantifolia*; 0.070** *Citrus aurantium*; 0.125** *C. clementina*; 0.183** *C. clementina* × *Citrus tangerina*; 0.184** *Citrus jambhiri*; 0.179** *Citrus japonica*

var. *margarita*; -0.048** *Citrus limettioides*; -0.031** *Citrus limonia*; 0.093** *Citrus medica*; 0.216** *Citrus reshni*; 0.061** *C. reticulata*; 0.101** *C. reticulata* × *Citrus temple*; 0.119** *C. sinensis* × *P. trifoliata*; -0.213** *Citrus sunki*; 0.061** *P. trifoliata*; 0.139** *Citrus unshiu*; 0.160** *Citrus paradisi*; 0.212** *C. paradisi* × *P. trifoliata*; **P-value < 0.01).

We noted that most genes tend to lie below the standard trajectory path and are poised towards GC_{3s} , which clearly demonstrates that MB is acting as a major factor for the wide variation in codon usage across the *Citrus* species and *P. trifoliata* (Fig. 2). However, there might be additional factors that might influence codon usage across these species. This dependence of codon usage on genome base composition (AT- or GC-richness) has been previously reported in several unicellular genomes.^{43,44} In a genome-wide analysis of eubacterial and archaeal genomes, it has been suggested that genome-wide variance in codon usage is primarily due to MB as the GC content shows wide variation along the isochors.¹⁷ Simultaneously, alternative views associate GC variability with various factors such as transcriptional optimization, methylation, recombination, and horizontal gene transfer.⁴⁵ It can be inferred that the cloning of orthologs and homologues conserved across the *Citrus* species and *P. trifoliata* that are AT rich and have low Nc values will require few degenerate primers. On the contrary, genes that are GC rich and having high GC values will require more degenerate primers for enhancement of cloning efficiency.

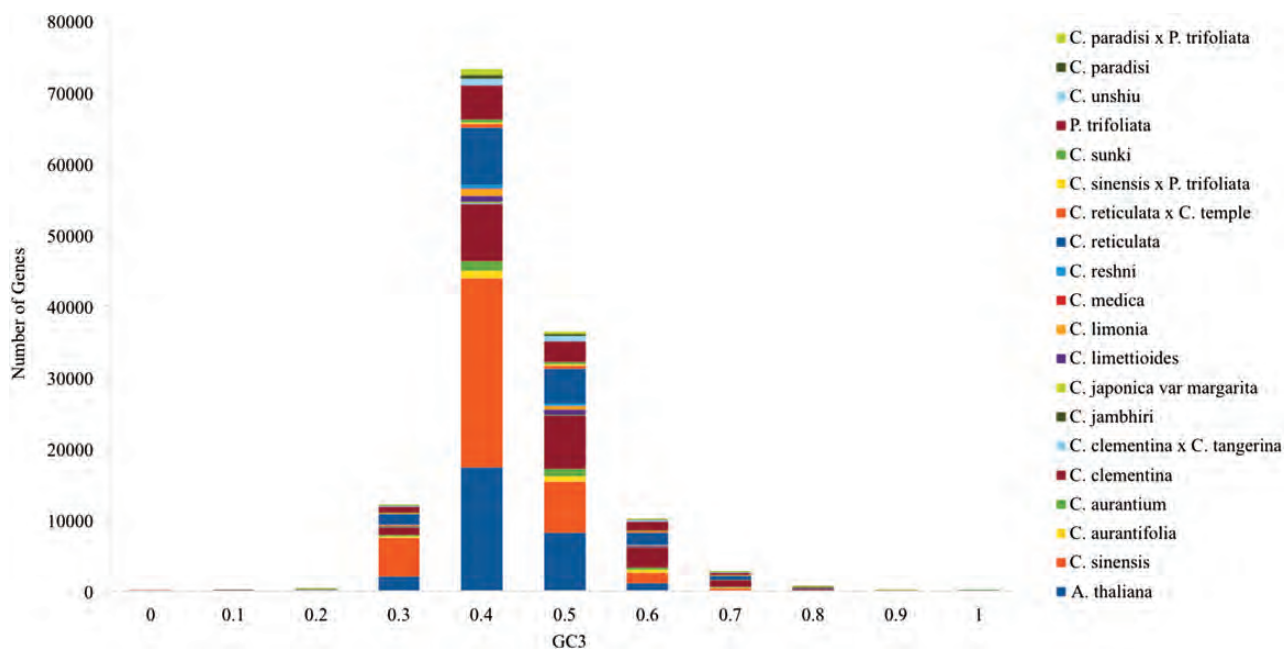


Figure 1. The distribution of GC_3 content in *Citrus* species, *P. trifoliata* and *A. thaliana* genes. The GC_3 content showed unimodal distribution.

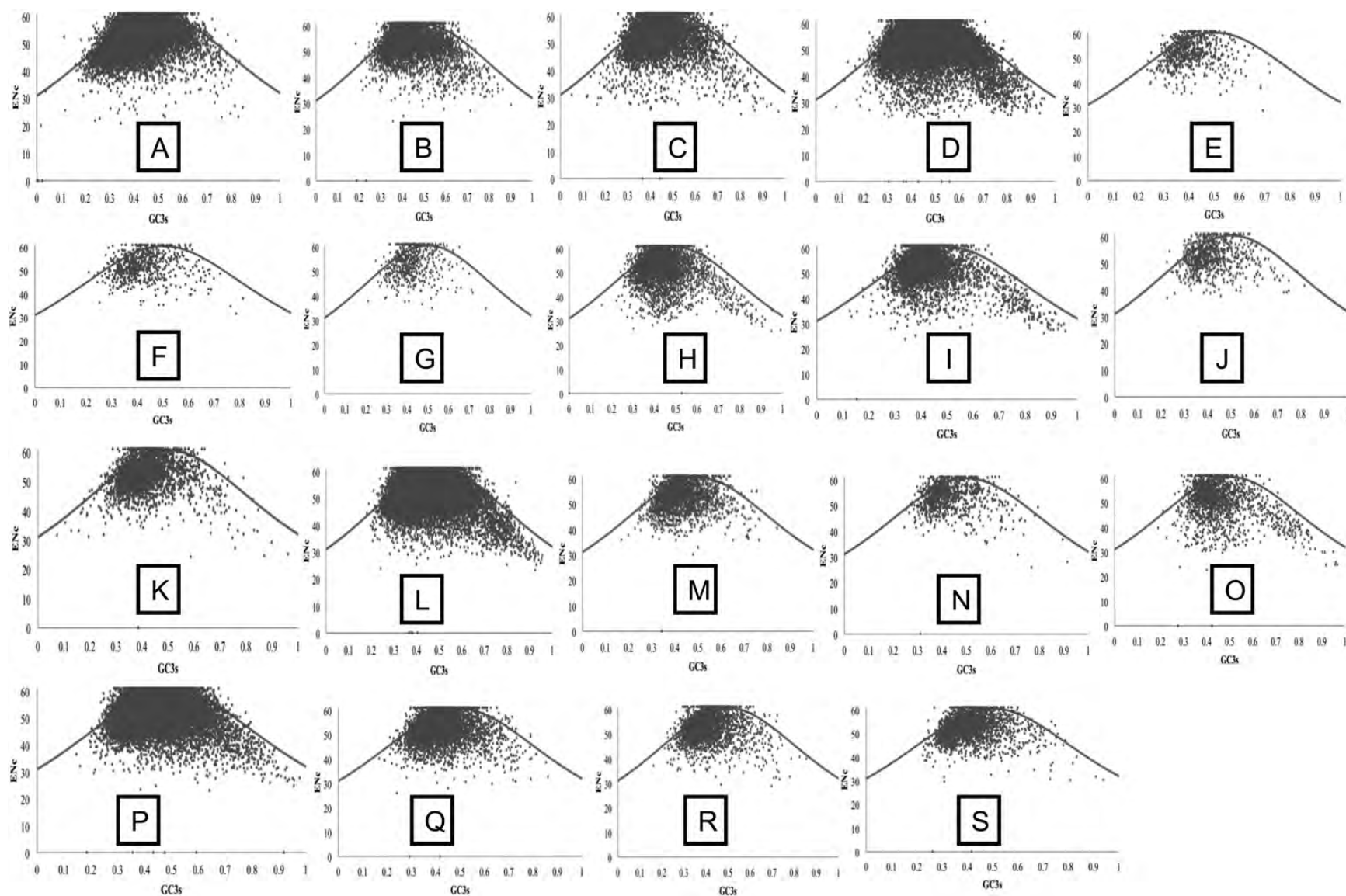


Figure 2. Nc versus GC_{3s} plot of *Citrus* species and *P. trifoliata* genes. The solid black line indicates the expected Nc value, if the codon bias is only due to GC_{3s}. Species order: A = *C. sinensis*; B = *C. aurantifolia*; C = *C. aurantium*; D = *C. clementina*; E = *C. clementina* × *C. tangerina*; F = *C. jambhiri*; G = *C. japonica* var. *margarita*; H = *C. limettoides*; I = *C. limonia*; J = *C. medica*; K = *C. reshni*; L = *C. reticulata*; M = *C. reticulata* × *C. temple*; N = *C. sinensis* × *P. trifoliata*; O = *C. sunki*; P = *P. trifoliata*; Q = *C. unshiu*; R = *C. paradisi*; and S = *C. paradisi* × *P. trifoliata*.

P. trifoliata, we extracted the expression of ESTs count using the CAP3 assembly (.ace files) and compared the usage of codons in accordance with earlier reports.^{53–55} We also selected the ribosomal protein-encoding genes and quantified the association of the ribosomal encoding genes with axis 1 (COA/RSCU) and axis 2 (COA/RSCU) of correspondence analysis.

A star map showing optimal codons was constructed taking 10% of the genes from the extreme tails of multivariate analysis (Table 2), and several distinct trends of optimal codons were detected. In an earlier study,³⁸ optimal codons in *Citrus* were estimated based on the correspondence analysis of codon usage, and the relative frequency of synonymous codon using 177 CDS and A- or T-ending optimal codons (TAA, GCT, GAT, CTT, AGG, AGA, and GTT) was observed. Overall, we observed that the trend of optimal codons usage was not conserved across all analysed species. However, in our analyses, we have observed that GCT (~1.57; RSCU) and GGT (~1.09; RSCU) representing Alanine and Glycine were found to be evolutionary conserved across all the species. Because both these optimal codons are T-ending codons, which is an indication of the dominant role of the MB in the conservation of the A- or T-ending codons, it potentially represents the biased genome composition. However, for the other observed optimal codons, pattern genomic composition was not able to explain the deviation. We further observed that for most amino acids with 2- to 6-fold degeneracy level, there has been a general preference for the usage of two or more codons as optimal codons. For example, in Glycine, two highly distributed optimal codons GGA and GGT were identified and they could be classified as the primary and secondary optimal codons preferentially based on the RSCU (GGA, ~1.16 and GGT, ~1.09). A curious trend observed among the *Citrus* species and *P. trifoliata* is that Leucine is frequently encoded optimally by a G-ending codon (TTG; ~RSCU: 1.49) in four species (*C. sinensis*, *C. clementina*, *C. reticulata*, and *P. trifoliata*) rather than synonymous A- or T-ending codon. However, in other species, CTT (~RSCU: 1.49) and CTC (~RSCU: 0.89) (TTA, RSCU: 1.16; CTT RSCU: 1.51) were the optimal codons.

We found deviations in the usage of optimal codons encoding Lysine. For example, Lysine is frequently encoded by AAA (~RSCU: 0.98) as optimal codon in three species (*C. sinensis*, *C. reticulata*, and *P. trifoliata*), whereas AAG (~RSCU: 1.01) was found to be the optimal codon for Lysine in rest of the *Citrus* species. In Arginine, we observed that in two species (*C. aurantifolia* and *C. paradisi* × *P. trifoliata*), AGG represents the potential optimal codon instead of the synonymous AGA. But based on the RSCU values, AGA (~RSCU: 1.66) was assumed to be more

dominant over the AGG codon (~RSCU: 1.52) at the respective levels of significance P -value < 0.01 and P -value < 0.05. Using a mutation and selection model, Knight et al.⁵⁶ demonstrated that 'pairs of species with convergent GC content might also evolve convergent protein sequences, especially at functionally unconstrained positions'. For instance, the frequencies of both Lysine and Arginine are highly anti-correlated with GC content, and Lysine and Arginine can easily be substituted for one another in proteins. This observation explains the inclusion of Lysine and Arginine in our study and is well supported by an earlier study in nematodes.⁵⁷ It has been proposed that Arginine and Leucine have a tendency to show different codon usage patterns because of the prevalence of the synonymous GC substitutions in the first and the third codon position.⁵⁸ Recently, it has been postulated that codon optimization significantly enhanced *MIR* gene expression in *Solanum lycopersicum* cv. Microtom,⁵⁹ which potentially depicts the importance and the usage of the optimal codons in gene expression and transgenics. To our knowledge, this is the first time large-scale identification of optimal codons in *Citrus* species and *P. trifoliata*, which could serve as a model repertoire to enhance the transformation efficiency.

3.4. Role of other selective constraints on codon bias in *Citrus* species and *P. trifoliata*

A recent study has revealed that MB deeply influences the folding stability of proteins, making proteins on the average less hydrophobic and, therefore, less stable with respect to unfolding and also less susceptible to misfolding and aggregation.⁶⁰ To identify the potential effects of Gravy and aromaticity, we computed non-parametric correlation coefficients between axis 1 (COA/RSCU) and Gravy or aromaticity scores for all studied species. We observed that Gravy and aromaticity played a minor role in shaping the variation of codon usage across *Citrus* species and *P. trifoliata*. In the case of *P. trifoliata*, no significant correlation was observed for aromaticity, suggesting that aromaticity has no significant role in shaping the codon usage variation in this genus.

3.5. Variations in GC₃ across *Citrus* species and *P. trifoliata*

Recent reports suggest that GC₃ composition and GC gradient are acting as major factors along the orientation of transcription in monocots. It has been also suggested that GC composition is vital for understanding chromatin remodelling, gene expression, and recombination.^{45,61} Some studies failed to depict GC gradient along the genes of dicot plants using *A. thaliana* as a model.⁶¹ The reason for this failure is that

Table 2. Star map showing optimal codon distribution in *Citrus* species and *P. trifoliata*

Codon/AA	A	B	C	D	E	F	G	H	I	J	K	L	M	N	O	P
TTT/Phe	*			*						*			*			
TTC/Phe		*	*			*	*	*			*	*		*	*	*
TTA/Leu	*			*						*			*			
TTG/Leu	*		*							*			*			
CTT/Leu	*	*	*	*	*			*	*	*	*	*	*	*	*	*
CTC/Leu		*	*	*	*			*	*		*	*		*	*	*
CTA/Leu	*			*		*				*			*			
CTG/Leu				*		*	*									
ATT/Ile	*							*		*	*		*			
ATC/Ile		*	*	*	*	*	*	*	*		*	*		*	*	*
ATA/Ile	*									*			*			
GTT/Val	*	*	*						*	*	*		*			*
GTC/Val		*	*	*	*	*	*	*	*		*	*		*	*	*
GTA/Val	*			*						*			*			
GTG/Val						*	*									
TCT/Ser	*	*	*	*	*			*	*	*	*	*	*	*	*	*
TCC/Ser		*	*		*			*	*		*			*	*	*
TCA/Ser	*	*	*	*		*	*			*	*	*	*	*		
TCG/Ser						*	*									
AGT/Ser	*					*	*					*	*			
AGC/Ser				*		*	*									
CCT/Pro	*	*	*	*		*	*	*	*	*	*	*	*	*	*	*
CCC/Pro					*			*	*		*	*			*	*
CCA/Pro	*	*	*	*	*	*	*		*	*	*	*	*	*	*	*
CCG/Pro						*	*									
ACT/Thr	*	*	*	*		*	*	*	*	*	*	*	*	*	*	*
ACC/Thr		*	*	*	*			*	*		*	*		*	*	*
ACA/Thr	*			*		*				*			*			
ACG/Thr						*	*									
GCT/Ala	*	*	*	*	*	*	*	*	*	*	*	*	*	*	*	*
GCC/Ala		*	*	*	*			*	*		*	*		*	*	*
GCA/Ala	*			*		*	*			*			*			
GCG/Ala																
TAT/Tyr	*									*			*			
TAC/Tyr		*	*	*		*	*				*	*				
CAT/His	*	*				*	*			*			*			
CAC/His			*	*												
CAA/Gln	*		*							*			*			
CAG/Gln		*		*		*	*					*				*
AAT/Asn	*	*	*	*		*		*			*	*			*	
AAA/Lys	*									*			*			
AAG/Lys		*	*	*	*	*	*	*			*	*		*	*	*
GAT/Asp	*	*	*	*		*						*			*	
GAA/Glu	*		*							*			*			
GAG/Glu		*		*		*	*					*				*
TGT/Cys	*									*			*			

Continued

Table 2. Continued

Codon/AA	A	B	C	D	E	F	G	H	I	J	K	L	M	N	O	P
TGC/Cys		*	*	*		*	*		*		*	*		*	*	
CGT/Arg		*	*	*	*	*	*	*	*		*	*		*	*	*
CGC/Arg		*	*	*	*	*	*		*		*	*		*	*	*
CGA/Arg	*			*		*	*				*		*			
CGG/Arg																
AGA/Arg	*		*	*		*	*			*			*			
AGG/Arg		*														*
GGT/Gly	*	*	*	*	*	*	*	*	*	*	*	*	*	*	*	*
GGC/Gly		*	*	*		*	*									
GGA/Gly	*		*	*		*	*			*	*		*		*	

Species are in the following order: A = *C. sinensis*; B = *C. aurantifolia*; C = *C. aurantium*; D = *C. clementina*; E = *C. jambhiri*; F = *C. limettioides*; G = *C. limonia*; H = *C. medica*; I = *C. reshni*; J = *C. reticulata*; K = *C. reticulata* × *C. temple*; L = *C. sinensis* × *P. trifoliata*; M = *P. trifoliata*; N = *C. unshiu*; O = *C. paradisi*; and P = *C. paradisi* × *P. trifoliata*. AA represents amino acid.

high and low GC₃ genes have opposite gradients along the genes, and when high and low GC₃ genes are clumped together, the effect disappears. In *Citrus* species and *P. trifoliata*, unimodal bell-shaped distribution of GC₃, centred at 0.39, is considered to be a typical mode of GC₃ distribution for dicot species. We selected the top and bottom 5% of genes across each studied *Citrus* species and *P. trifoliata*, and strikingly distinct gradients of GC₃ and CG₃ skew for high and low GC₃ genes were revealed. Genes with high GC₃ showed higher level of GC₃ codons in their middle coding regions than in terminal coding regions. In addition, high GC₃ genes have a preference for C over G in the middle coding regions, and this preference was not found in the 3'-coding end ~100 bp in size (Fig. 4; Supplementary Fig. 2).

It was previously reported that stress-related genes in grasses are GC₃-rich.⁴⁵ We stratified all *Citrus* species and *P. trifoliata* genes into three groups by GC₃: rich (top 5%), poor (bottom 5%), and medium (middle 90%). We observed that among the four major economically important *Citrus* species, in *C. reticulata* (1718), *C. clementina* (1522), and *P. trifoliata* (951), most of the GC₃-rich genes were related to stress in comparison to the total number of observed stress-related genes, but in the case of *C. sinensis*, it was observed that a high number of medium GC₃ genes (1857) were also abundant in stress-related genes. The relative abundance of the medium GC₃ genes in *C. sinensis* may be due to the time-course adaptability of this species during the period of evolution, suggesting an adaptive evolution towards the stress.

A high abundance of stress-related genes in *C. reticulata* and *P. trifoliata* may be closely related to their high stress-tolerance trait and especially to *P. trifoliata*

that has the highest cold-resistant character among *Citrus* species and its wild relatives. The occurrence of the high GC₃ genes in all these species suggests an association between DNA methylation and GC₃ as it has been previously suggested that high GC₃ composition has been influenced by the GC mismatch-repair mechanism that is dominant in stress-associated genes as the absence of this positive bias may lead to the loss of the recombination repair mechanism and could be detrimental to the plant adaptation in the evolving stress conditions. Genomic regions under higher selective pressure are more frequently recombining, and as a result relative increase in GC₃ content can be observed.⁴⁵ As shown in Supplementary Table 2 and Fig. 5, in all *Citrus* species and *P. trifoliata*, the ratio of stress to non-stress-related genes in GC₃-rich group was elevated in comparison to the GC₃-medium group and depleted in the GC₃-poor group. It is worth noting that GC₃-poor group has fewer genes functionally described as stress-related when compared with the GC₃-rich group.

In all the studied species, GC₃-rich and -poor genes have different trends from 5' to 3' end of the genes. GC₃-rich genes become even more GC₃ rich, and GC₃-poor genes become more GC₃ poor. Firstly, the GC₃-rich group has more stress related genes than the GC₃-poor group. Secondly, the GC₃-rich group has a positive gradient of GC₃ from 5' and 3' flanks to the middle portion of the CDS. The low-GC₃ group has a negative gradient of GC₃ from 5' and 3' flanks to the middle portion of the CDS. Thirdly, rich- and poor-GC₃ groups showed different CG₃ skew trends along the CDS: GC₃-rich genes favour C₃ over G₃, and this preference is most pronounced in first 150 codons, whereas GC₃-poor genes favour G₃

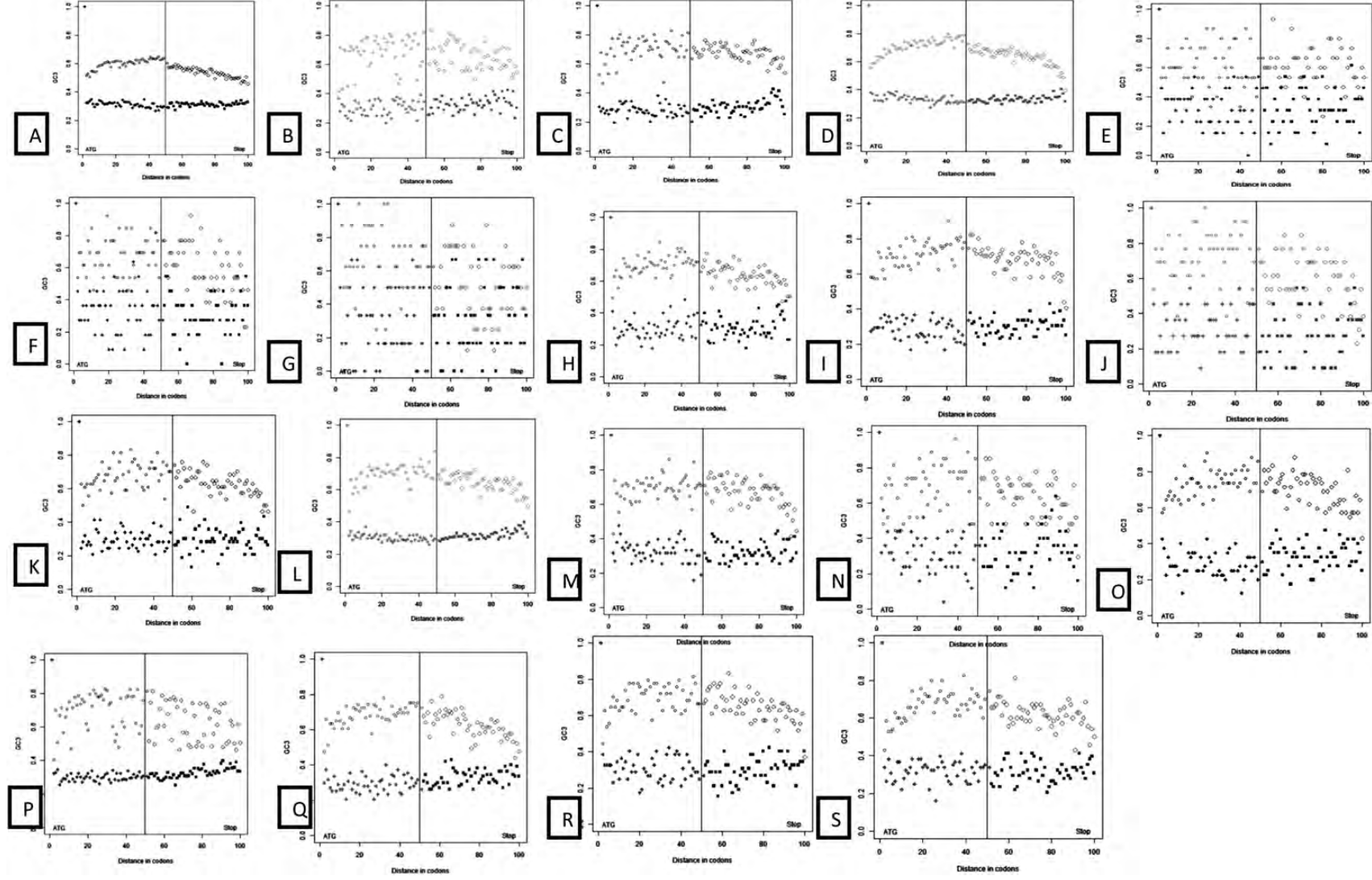


Figure 4. GC₃ gradient across *Citrus* species and *P. trifoliata*. Species order: A = *C. sinensis*; B = *C. aurantifolia*; C = *C. aurantium*; D = *C. clementina*; E = *C. clementina* × *C. tangerina*; F = *C. jambhiri*; G = *C. japonica* var. *margarita*; H = *C. limettoides*; I = *C. limonia*; J = *C. medica*; K = *C. reshni*; L = *C. reticulata*; M = *C. reticulata* × *C. temple*; N = *C. sinensis* × *P. trifoliata*; O = *C. sunki*; P = *P. trifoliata*; Q = *C. unshiu*; R = *C. paradisi*; and S = *C. paradisi* × *P. trifoliata*. These plots show gradients of GC₃ for 5% of GC₃-rich and GC₃-poor genes for all analysed genomes. Gradients from 5' and 3' ends of the CDS are shown in the same plot, separated by a vertical line. For all genomes, GC₃-rich genes become more GC₃ rich towards the middle of the CDS.

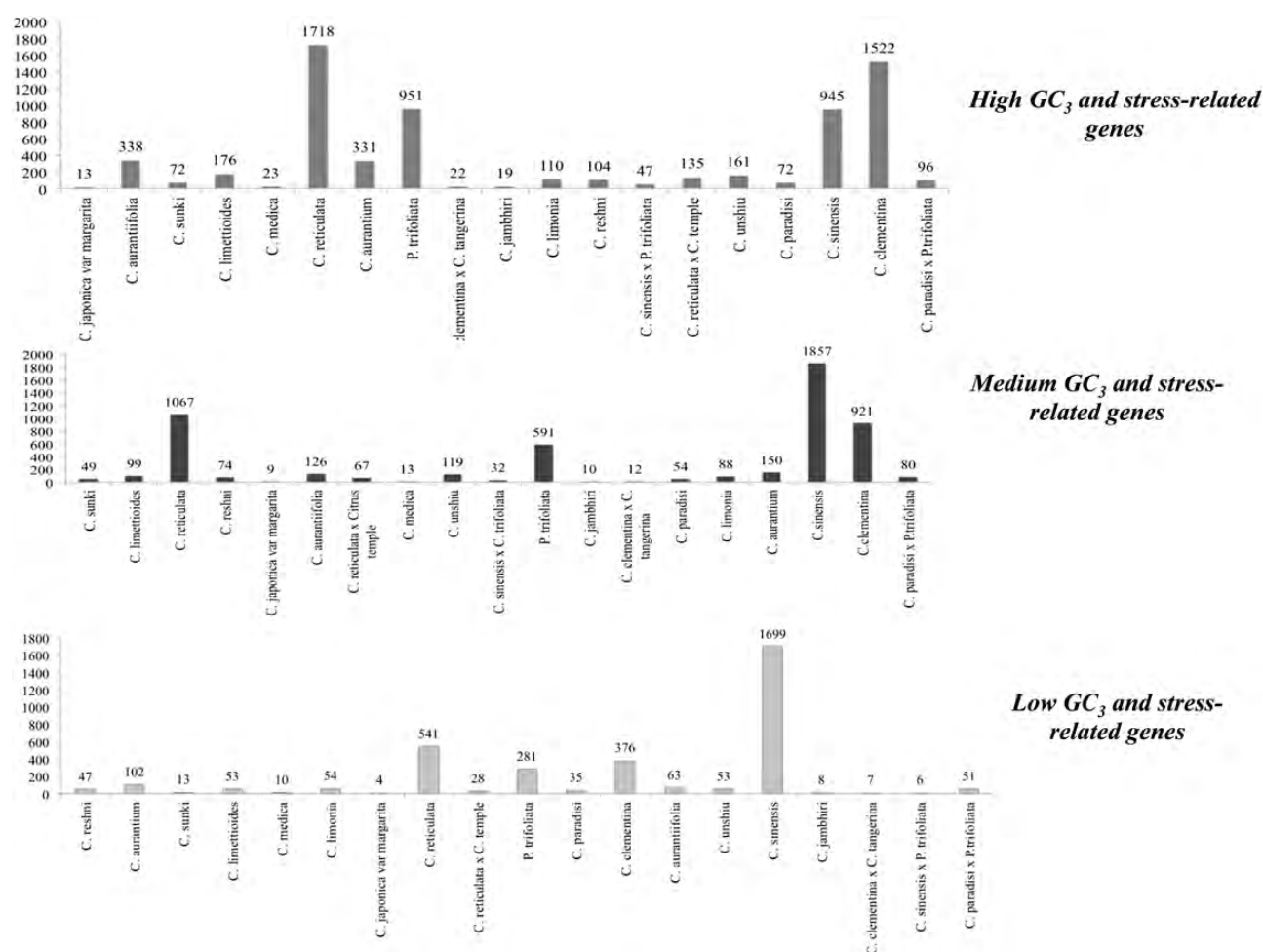


Figure 5. Distribution of stress-related genes in *Citrus* species and *P. trifoliata* according to the GO of *Arabidopsis*.

over C₃ (Supplementary Fig. 2). It was suggested that transcriptional and translational optimization of stress-related and tissue-specific genes is the major force responsible for maintaining high GC₃ content.⁴⁶ The replacement of the AT pair with the GC pair at the third codon position enhances the transcriptional activity that in turn enhances the array of ribosomal-binding proteins. Stayssman et al.⁶² provide a strong positive correlation between the methylation of internal unmethylated regions and expression of the host gene and postulated that the genes with high GC₃ provide more targets for methylation. Genes involved in response to various stresses need to produce a protein with a faster rate as a response to external stimulus. This results in shortened length of transcript and preference for G and C (and especially, C) in the third position of the codon (to avoid abortive transcription and ribosome congestion). In addition, high GC₃ genes have more methylation targets that allow for fine-tuning of transcriptional regulation.⁵⁸

3.6. Evolution and coevolution of nucleotide and amino acid composition

We analysed the relationship between amino acid sequence divergence and variation in GC₃, to test the effect of evolutionary divergence on codon usage. We observed that there is an overall positive correlation (0.43) between relative change in GC₃ and normalized Hamming distance (Fig. 6). This was expected because more diverse amino acid sequences are likely to have more diverse nucleotide sequences. The trend is not uniform: for the genes with GC₃ in the range between 0.5 and 0.6, the correlation is 0.596 and it drops to 0.32 for GC₃ above 0.8 or below 0.4. There are some pairs of organisms that have negative correlation between GC₃ and Hamming distance (*C. jambhiri*: *C. limettioides*, *C. limonia*: *C. sinensis* x *P. trifoliata*, *C. aurantiifolia*: *C. jambhiri*, *C. jambhiri*: *C. paradisi*, *C. aurantiifolia*: *C. paradisi*, *C. jambhiri*: *P. trifoliata*, *C. medica*: *C. sinensis* x *P. trifoliata*, *C. jambhiri*: *C. unshiu*, *C. jambhiri*: *C. reshni*, *C. japonica* var. *margarita*: *C. paradisi*,

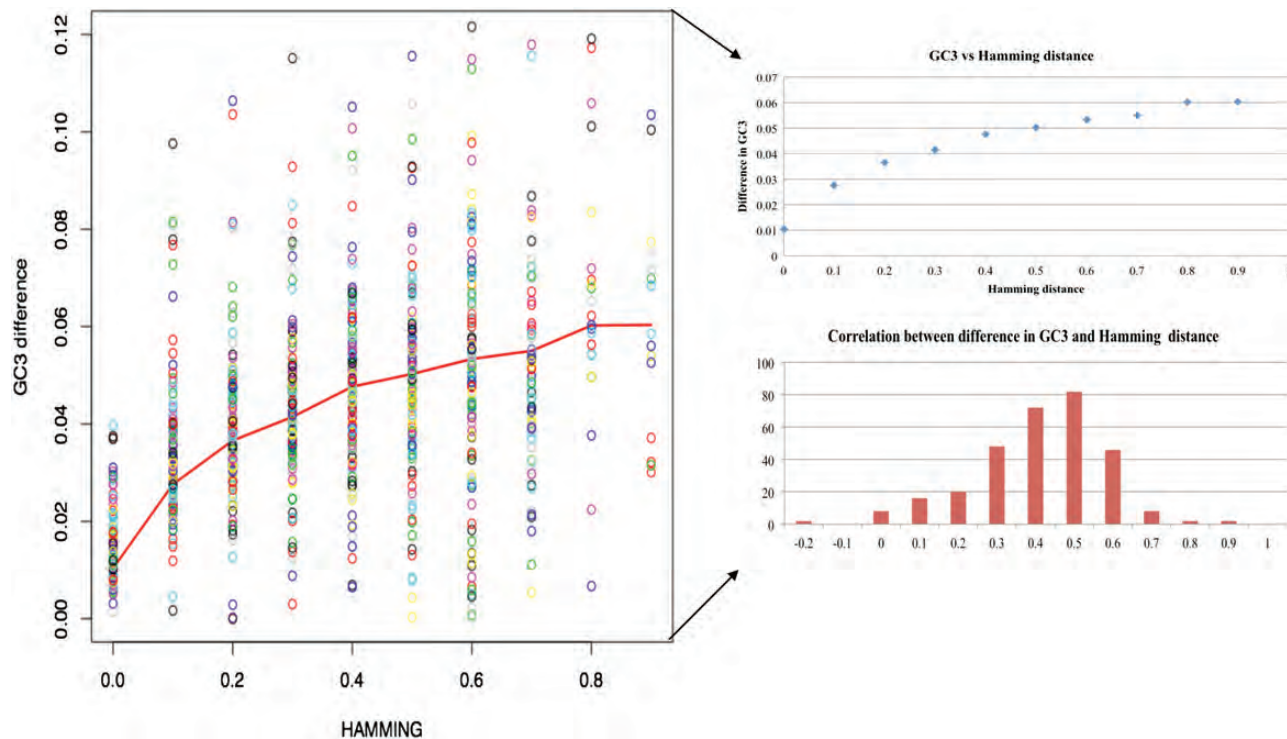


Figure 6. Hamming distance versus change in GC₃ visualization across *Citrus* species and *P. trifoliata*. Relative difference GC₃ composition between genes A and B, defined as $GC_3(A) - GC_3(B) / [GC_3(A) + GC_3(B)]$ is positively correlated with the Hamming distance between corresponding protein sequences, calculated as $1 - [I(AB) + I(BA)] / [I(AA) + I(BB)]$, where $I(AB)$ is the number of identities in alignment of protein A to protein B. The resulting Hamming distance was rounded to 1 DP, and an average difference in GC₃ was computed for each category.

C. aurantiifolia: *C. clementina* × *C. tangerina*, *C. aurantium*: *C. jambhiri*, *C. clementina* × *C. tangerina*: *C. jambhiri*). Note that in the list of negative correlations, *C. jambhiri* occurs eight times.

Genes that have low GC₃ content in *C. jambhiri* (<0.4) tend to have negative correlation between difference in GC₃ and Hamming distance, and genes with high GC₃ content (>0.6) are slightly positively correlated. This atypical behaviour can be explained by larger evolutionary distance between *C. jambhiri* and other species analysed in the present investigation. The highest correlation was observed between *C. japonica* var. *margarita* and *C. limonia* ($\rho = 0.86$, number of orthologous pairs $N = 12$, t -statistics $t = 5.22$, P -value = 0.0002). *C. japonica* var. *margarita* and *C. limonia* are evolutionary distant, and the number of orthologous pairs is only $N = 12$. Hence, a high value of correlation coefficient may be a result of an evolutionary pressure to conserve codon usage for a selected subset of conserved proteins. We observed that pairs of species having positive correlation between GC₃ and Hamming distance are enriched in translation, transport, proteolysis, photorespiration, and genes involved in photosynthesis; pairs of species with negative correlation are enriched in stress-response genes.

Similar patterns of hierarchical clustering were depicted; when we clustered the species based on co-orthologous genes using genome-predicted and frame-corrected reconstructed proteins of the *Citrus* species and *P. trifoliata*. The phylogenetic clades were rerooted using *P. trifoliata* as an outgroup, and it was observed that *C. sinensis*, *C. clementina*, and *C. reticulata* all belong to the same clade, which are in accordance with the previous reports using major intrinsic proteins (XIP subfamily of aquaporins) and further support the conserved homology between these two species.⁶³ These supportive views suggest that the identified frame-corrected coding regions are trustworthy and accurate to be used for the predictions in the species, with no genome information available till now (Supplementary Fig. 3A and B).

Because a nucleotide bias can lead to an overall bias in amino acid composition of proteins, it is possible that a genome with nucleotide bias may have introduced atypical amino acid substitutions in its proteome. Hence, AT-rich coding sequences would encode proteins rich in FYMINK amino acids (Phenylalanine, Tyrosine, Methionine, Isoleucine, Asparagine, and Lysine), whereas GC-rich coding sequences would produce proteins containing high levels of GARP amino acids (Glycine, Alanine, Arginine, and Proline).

Because *Citrus* species and *P. trifoliata* are AT-rich, we found higher proportion of FYMINK [(FYMINK and CDS_{GC} ($y = 4.1431x + 22.341$, $R^2 = 0.9099$); GARP and CDS_{GC} ($y = 3.0687x + 45.414$, $R^2 = 0.7803$); P -value $< 10^{-9}$)].

Because the third synonymous position does not influence the protein sequence, we computed the correlation coefficient between the FYMINK and GARP parameters and the GC₃. This correlation is an important factor in describing nucleotide bias at synonymous and non-synonymous sites. We observed a high significant correlation between nucleotide composition and amino acid composition at the third synonymous position [(FYMINK and CDS_{GC3} ($y = 10.862x + 142.4$, $R^2 = 0.9296$); GARP and CDS_{GC3} ($y = 9.842x + 212.15$, $R^2 = 0.7495$)], suggesting that nucleotide bias has an influence on amino acid composition in *Citrus* species and *P. trifoliata*.

In summary, we found prevalence of A- or T-ending codons with an exception for Lysine and Arginine in *P. trifoliata*. We suggest that although the patterns of optimal codons were not conserved, two codons (GCT and GGT) were found to be conserved across all the *Citrus* species and *P. trifoliata*. We analysed GC₃-rich and -poor genes and their association with stress, and our results provided novel insights into stress evolution of stress adaptation in *Citrus* species and *P. trifoliata* in accordance with the GC₃ biology. This research is critical for all *Citrus* species, as it might facilitate understanding of genome dynamics and evolution in *Citrus* and *P. trifoliata*, transformation of interested target genes, designing multitargeting gene systems, and ultimately genetically improving these important fruit crops.

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