Since January 2020 Elsevier has created a COVID-19 resource centre with free information in English and Mandarin on the novel coronavirus COVID-19. The COVID-19 resource centre is hosted on Elsevier Connect, the company's public news and information website.

Elsevier hereby grants permission to make all its COVID-19-related research that is available on the COVID-19 resource centre - including this research content - immediately available in PubMed Central and other publicly funded repositories, such as the WHO COVID database with rights for unrestricted research re-use and analyses in any form or by any means with acknowledgement of the original source. These permissions are granted for free by Elsevier for as long as the COVID-19 resource centre remains active.
A comprehensive analysis of genome composition and codon usage patterns of emerging coronaviruses

Fernando L. Torta,1, Matías Castells,1, Juan Cristina,b,*

1 Laboratorio de Virología Molecular, Sede Salto, Centro Universitario Regional, Litoral Norte, Universidad de la República, Gral. Rivera 1350, 50000, Salto, Uruguay
2 Laboratorio de Virología Molecular, Centro de Investigaciones Nucleares, Facultad de Ciencias, Universidad de la República, Iguá 4225, Montevideo, 11400, Uruguay

A R T I C L E   I N F O

Keywords: Coronavirus Evolution Codon usage Wuhan 2019-nCoV SARS-CoV-2

A B S T R A C T

An outbreak of atypical pneumonia caused by a novel Betacoronavirus (βCoV), named SARS-CoV-2 has been declared a public health emergency of international concern by the World Health Organization. In order to gain insight into the emergence, evolution and adaptation of SARS-CoV-2 viruses, a comprehensive analysis of genome composition and codon usage of βCoV circulating in China was performed. A biased nucleotide composition was found for SARS-CoV-2 genome. This bias in genomic composition is reflected in its codon and amino acid usage patterns. The overall codon usage in SARS-CoV-2 is similar among themselves and slightly biased. Most of the highly frequent codons are A- and U-ending, which strongly suggests that mutational bias is the main force shaping codon usage in this virus. Significant differences in relative synonymous codon usage frequencies among SARS-CoV-2 and human cells were found. These differences are due to codon usage preferences.

1. Introduction

The family Coronaviridae consists of four genera, namely, Alphacoronavirus (αCoV), Betacoronavirus (βCoV), Gammacoronavirus (γCoV) and Deltacoronavirus (δCoV) (Chen et al., 2020). Viruses from this family possess a single stranded, positive-sense RNA genome ranging from 26 to 32 kilobases in length (Su et al., 2016). Coronaviruses (CoV) have been identified in several avian hosts, as well as in mammals, including humans, bats, civets, mice, dogs, cats, cows and camels (Clark, 1993; Cavanagh, 2007; Zhou et al., 2018). Although several CoV are pathogenic to humans, most of them are associated with mild clinical symptoms (Su et al., 2016). Nevertheless, two notable exceptions have been described: severe acute respiratory syndrome (SARS) coronavirus (SARS-CoV), a novel βCoV that emerged in southern China in 2002 (Peiris et al., 2004) and Middle East respiratory syndrome (MERS) coronavirus (MERS-CoV), which was first detected in Saudi Arabia in 2012 (Zaki et al., 2012). Before the SARS epidemic, bats were not known to be hosts for CoVs. In the last 15 years, bats have been found to be hosts of more than 30 CoVs. Interactions among various bat species themselves, bat-animal and bat-human interactions, such as the presence of live bats in wildlife wet markets in China, have been proved to be important for interspecies transmission of CoVs (Wong et al., 2019). In fact, both SARS-CoV and MERS-CoV likely originated in bats, and genetically diverse CoVs that are related to these viruses were discovered in bats worldwide (Cai et al., 2019). Previous studies in different bat species from China permitted the identification of at least 41 new βCoV, all of them were Rhinolophus spp. bat-CoV (Lin et al., 2017).

At this moment, in Wuhan, the capital city of Hubei province of the People’s Republic of China, an outbreak of atypical pneumonia caused by a novel coronavirus (SARS-CoV-2) is currently underway. The outbreak appears to have started from a zoonotic transmission at a market in Wuhan where animals and meat were sold (Chan et al., 2020a). As February 26th, 2020, there have been 77,780 cases of SARS-CoV-2 confirmed in China, including 2666 deaths (WHO, 2020a). In addition, SARS-CoV-2 has been reported in 33 countries outside China, with 2459 cases confirmed and 34 deaths (WHO, 2020a). Very recent studies revealed that SARS-CoV-2 can be considered a new human-infecting βCoV (Lu et al., 2020).

The World Health Organization declare this 2019-nCoV outbreak as a public health emergency of international concern on January 30th, 2020 (WHO, 2020b) and the disease caused by this specific virus species have recently been designated as COVID-19 (Coronavirus Infectious Diseas-19) (WHO, 2020a). In this sense, the Coronavirus Study Group of the International Committee on Taxonomy of Viruses (ICTV), formally recognizes this virus as a relative to severe acute respiratory...
syndrome SARS-CoVs and designates it as severe acute respiratory syndrome coronavirus 2: SARS-CoV-2 (Gorbalenya et al., 2020).

In order to gain insight into the emergence, evolution, adaptation and spread of the SARS-CoV-2 viruses, a comprehensive analysis of genome composition and codon usage of βCoV circulating in China was performed.

2. Material and methods

2.1. Sequences

Available and comparable complete genome sequences of 81 βCoV strains isolated in China, including SARS-CoV-2 as well as βCoV isolated from different hosts, were obtained from GenBank database (available at: http://www.ncbi.nlm.nih.gov). For accession number, strain name, host and date of isolation, see Supplementary material Table 1. For each strain ORFs1a + ORF1b + S + E + N + M were concatenated. Sequences were aligned using the MACSE program (Ranwez et al., 2011). MACSE algorithm is a useful tool for accommodating sequencing errors and other biological deviations from the coding frame (Ranwez et al., 2011). The alignment is available upon request. The dataset comprised a total of 725,433 codons.

2.2. Data analysis

Codon usage, amino acid usage, dinucleotide frequencies, base composition, the relative synonymous codon usage (RSCU) (Sharp and Li, 1986), total GC content, GC content in the third position of the codon (GC3s) and effective numbers of codons (ENC) were calculated using the program CodonW (written by John Peden) as implemented in the Galaxy server version 1.4.4 (Afgan et al., 2018).

The relationship between compositional variables and samples was obtained using multivariate statistical analyses. Principal component analysis (PCA) is a type of multivariate analysis that allows a dimensionality reduction. Singular Value Decomposition (SVD) method was used to calculate the principal components (PC). Unit variance was used as scaling method. By the same approach, Heatmaps were also constructed, which is a data matrix for visualizing values in the dataset by the use of a color gradient. This gives a good overview of the largest and smallest values in the matrix. Rows and/or columns of the matrix are clustered so that sets of rows or columns rather than individual ones can be interpreted. PCA and Heatmaps analysis were done using the ClustVis program (Metsalu and Vilo, 2015).

The RSCU values of human cells were obtained from Kazusa database (available at: http://www.kazusa.or.jp/codon/).

To study codon usage preferences in SARS-CoV-2 in relation to the codon usage of human cells, we employed the codon adaptation index (CAI) (Sharp and Li, 1987). CAI was calculated using the approach of Puigbo et al. (2008a) (available at: http://genomes.urv.es/CAIcalc) for human cells. This method allows to compare a given codon usage (in our case, SARS-CoV-2) to a predefined reference set (human). A statistically significant difference among CAI values was addressed.
applying a Wilcoxon & Mann-Whitney test (Wessa, 2012). In order to discern if statistically significant differences in the CAI values arise from codon preferences, we used e-CAI (Puigbo et al., 2008b) to calculate the expected value of CAI (eCAI) at the 95% confident interval. A Kolmogorov-Smirnov test for the expected CAI was also performed (Puigbo et al., 2008b).

3. Results

3.1. Trends in compositional properties across βCoV strains circulating in China

In order to study the genetic composition of SARS-CoV-2 emerging in China, 81 ORFs sequences from βCoV strains isolated in China (including SARS-CoV-2 and SARS-CoV from humans and βCoV isolated from bats, civets and ferrets) were aligned and the nucleotide frequencies at third codon position, total GC content, GC content at the third codon position, ENC and dinucleotide frequencies were established for all strains ORFs and PCA was performed. The results of these studies are shown in Fig. 1.

Heatmap analysis on genomic composition of all βCoV enrolled in these studies revealed a distinct genome composition of SARS-CoV-2 strains isolated in China, in relation to all other strains isolated from humans as well as bats, civets or ferrets (see Fig. 1).

To study if these differences in genomic composition can be observed at codon and amino acid usage, these variables were established for all βCoV strains enrolled in these studies and their relations were observed by Heatmap analysis. The results of these studies are shown in Fig. 2. Significant differences in codon and amino acid usage was observed among SARS-CoV-2 strains and all other βCoV strains included in these studies (see Fig. 2). Among β-CoV strains isolated from bats, a significant variation was also observed. Linkage analysis suggests a closer relation among SARS-CoV-2 and β-CoV isolated from bats, and a more distant relation with SARS-CoV-2 and other β-CoV isolated from civet and ferret enrolled in these analyses (Fig. 2).

3.2. The nucleotide frequencies in SARS-CoV-2 genome

To gain insight into the genomic composition of SARS-CoV-2 strains, the nucleotide frequencies found for this virus where compared to the nucleotide frequencies found for other human CoVs. The results of these studies are shown in Fig. 3.

Some general characteristics of CoVs were observed, since the U-frequencies is significantly above average, while the C-frequencies are below the expected frequencies. In the case of purines, A is preferred over G (Fig. 3). As it can be seen in Fig. 3, most variation occurs in the C/U and not the A/G section. Moreover, the results of these studies also revealed the presence of species-specific trends, since the C/U ratio differs profoundly per coronavirus type (see Fig. 3). Comparison of SARS-CoV-2 with the other two recent zoonotic transmission to the human population (SARS and MERS-CoVs) reveals that SARS-CoV-2 is quite extreme with a C-count of 18.4% (Fig. 3). On the other hand, SARS-CoV-2 has the highest A-count among CoVs enrolled in these studies (29.9%), mean A-count of 27.6 ±1.1 for all human CoVs enrolled in these studies (see Fig. 3).

3.3. General codon usage patterns in βCoV

To study the extent of codon usage bias in SARS-CoV-2, the ENC’s values were calculated for all SARS-CoV-2 strains enrolled in these studies. A mean value of 48.54 ± 2.34 was obtained. Due to the fact that all values obtained were > 40, the results of these studies suggest that the overall codon usage among SARS-CoV-2 is similar among themselves and slightly biased. Mean ENC values of 49.11 ± 0.02, 49.66 ± 0.52 and 49.21 ± 0.05 were found for SARS and βCoV isolated from bats and civets, respectively. ENC quantifies how far a codon usage departs from equal usage of synonymous codons and is a measure of codon usage bias in genomes that ranges from 20 (maximal bias) to 61 (unbiased) (Wright, 1990). Although βCoV ENC values are roughly similar, in the case of SARS-CoV-2 strains we observed a range of ENC values from 45.18 to 50.09.

Since codon usage by its very nature is multivariate, it is necessary to analyze the data using different and complementary approaches. An ENC–GC3S plot (ENC plotted against GC content at the third codon position) can be used as a method that quantifies how far the codon usage of a gene departs from equal usage of synonymous codons (Wright, 1990). If GC3S is the only determinant factor shaping the codon usage pattern, the values of ENC would fall on a continuous curve, which represents random codon usage. If G + C compositional constraint influences the codon usage, then the GC3S and ENC correlated spots would lie on or below the expected curve (Tsai et al., 2007).

When the ENC–GC3S plot was constructed with values obtained for all 81 βCoV strains enrolled in this analysis (including SARS-CoV-2 strains), all spots lie below the expected curve, indicating that G + C compositional constraints may play a role in all βCoVs codon usage (see Fig. 4).
3.4. Codon usage preferences in SARS-CoV-2

In order to compare the codon usage preferences of SARS-CoV-2 with those of human cells, the RSCU values of the codons in SARS-CoV-2 ORFs were calculated and compared with those of human. The results of these studies are shown in Table 1.

The frequencies of RSCU in SARS-CoV-2 ORFs are significantly different in relation to human cells. Highly biased frequencies were found for UUU (Phe), UUA (Leu), CUU (Leu), AUU (Ile), AUA (Ile), GUU (Val), UAU (Tyr), CAU (His), CAA (Gln), AAU (Asp), GAA (Glu), UCU (Ser), CCU (Pro), CCA (Pro), GCU (Ala), UGU (Cys), CGU (Arg), AGA (Arg), and GGU (Gly). As it can be seen, most of the highly preferred codons are A and U-ending, while most of the highly underrepresented codons are C and G-ending codons, particularly CG containing codons. These results strongly suggest that mutational bias is a main force shaping codon usage in SARS-CoV-2 (see Table 1). The occurrence of dinucleotides is established, they are not random and no dinucleotide is present at the expected frequencies (see Supplementary material Table 2). The relative abundances of CpG and CpC and GpG showed a strong deviation from the expected frequencies (i.e. 1.0) (mean ± S.D. = 0.22 ± 0.00, 0.48 ± 0.00 and 0.66 ± 0.00, respectively) and were markedly under-represented. Cytosine deamination and selection against CpG motifs have been proposed as two independent selection forces that shape codon usage bias in CoV (Woo et al., 2007), suggesting that immune selection may play a role in SARS-CoV-2 codon usage bias. On the other hand, UpU, ApA, UpG are markedly over-used (mean ± S.D. = 1.96 ± 0.00, 1.55 ± 0.00 and 1.40 ± 0.00, respectively) (Supplementary material Table 2). These results indicate that the composition of dinucleotides also determines the variation in synonymous codon usage among SARS-CoV-2 strains.

Comparison of SARS-CoV-2 RSCU values with the ones of SARS-CoV (Gu et al., 2004) and MERS-CoV (Chen et al., 2017) revealed a similar pattern for most of the codons (Table 1). Highly underrepresented codons ACG (Thr), GCG (Ala), GGG (Gly) were observed in all three viruses in relation to human cells (Table 1). On the other hand, significant differences in frequencies of two glutamine codons (GAG and GAA) were observed among SARS-CoV-2 and SARS and MERS-CoVs (Table 1). While GAG codon frequency is significantly different from the expected frequency (i.e. 1.0) in SARS-CoV-2 (0.55), it is not in SARS-CoV and MERS-CoV viruses (0.96 and 0.95, respectively). In the case of GAA, this codon is highly expressed in SARS-CoV-2 (1.45) and its frequency is very near the expected frequency in SARS-CoV and...
MERS-CoVs (1.04 and 1.05, respectively). These results suggest that although a general codon usage pattern among human βCoV can be found, each virus may evolve to a unique codon usage in its adaptation to the host cells.

3.5. Codon usage adaptation in SARS-CoV-2

In order to compare the codon usage preferences of SARS-CoV-2 with those of humans, CAI values for all triplets were calculated, using human codon usage as reference set. The results of these studies are shown in Table 2.

CAI index ranges from 0 to 1, being 1 if the frequency of codon usage by SARS-CoV-2 equals the frequency of usage of the reference set. A mean value of 0.710 was obtained for SARS-CoV-2 genes in relation to human (see Table 2). To evaluate if the differences were statistically significant, we performed a Wilcoxon & Mann-Whitney test. The results of this test revealed that the differences in CAI values were statistically significant (T = 256, p-value < 0.001).

To discern if the statistically significant differences in CAI values arise from codon preferences (Puigbo et al., 2008a), the expected CAI (e-CAI) values were calculated for SARS-CoV-2 sequences in relation to human codon usage reference set. The e-CAI algorithm (Puigbo et al., 2008a) generated 500 random sequences with the same nucleotide and amino acid composition as the sequences of interest (in this case SARS-CoV-2 sequences). Then, we calculated the CAI values for all of them, and applied a Kolmogorov-Smirnov test for the e-CAI of these random sequences (Kolmogorov-Smirnov test of e-CAI) values were calculated for SARS-CoV-2 sequences in relation to human codon usage reference set. The e-CAI algorithm (Puigbo et al., 2008a) generated 500 random sequences with the same nucleotide and amino acid composition as the sequences of interest (in this case SARS-CoV-2 sequences). Then, we calculated the CAI values for all of them, and applied a Kolmogorov-Smirnov test for the e-CAI of these random sequences in order to show whether the generated sequences follow a normal distribution. The results of these studies revealed an e-CAI value of 0.719 (p < 0.05). Kolmogorov-Smirnov test revealed a normal distribution of the generated sequences (Kolmogorov-Smirnov test of e-CAI value of 0.028, which is below critical value of 0.061). Taking all these results together, our studies revealed that the CAI values for SARS-CoV-2 genes are different from the CAI values obtained from human codon usage reference set. The e-CAI algorithm (Puigbo et al., 2008b) generated 500 random sequences with the same nucleotide and amino acid composition as the sequences of interest (in this case SARS-CoV-2 sequences). Then, we calculated the CAI values for all of them, and applied a Kolmogorov-Smirnov test for the e-CAI of these random sequences in order to show whether the generated sequences follow a normal distribution. The results of these studies revealed an e-CAI value of 0.719 (p < 0.05). Kolmogorov-Smirnov test revealed a normal distribution of the generated sequences (Kolmogorov-Smirnov test of e-CAI value of 0.028, which is below critical value of 0.061). Taking all these results together, our studies revealed that the CAI values for SARS-CoV-2 genes are different from the CAI values obtained from human cells. Again, these results suggest that these differences are related to codon usage preferences.

Table 1
Codon usage in SARS-CoV-2, displayed as RSCU values.

| AA | Cod | HC | CoV-2 | SARS | MERS | AA | Cod | HC | CoV-2 | SARS | MERS |
|----|-----|----|-------|------|------|----|-----|----|-------|------|------|
| Phe | UUU | 0.92 | 1.42 | 1.23 | 1.28 | Ser | UCU | 1.14 | 2.00 | 1.96 | 2.11 |
| | UUC | 1.08 | 0.58 | 0.77 | 0.71 | UCC | 1.32 | 0.44 | 0.42 | 0.71 |
| Leu | UUA | 0.48 | 1.66 | 1.04 | 1.21 | GGA | 0.90 | 1.63 | 1.70 | 1.21 |
| | UUG | 0.78 | 1.06 | 1.10 | 1.43 | UGG | 0.30 | 0.11 | 0.23 | 0.18 |

Table 2
Codon adaptation of SARS-CoV-2 genes in relation to human codon usage, displayed as CAI values.

| SARS-CoV-2 genes | CAI-Hs | CAI | Human genes |
|------------------|--------|----|-------------|
| 0.710 ± 0.003    |        |    | 0.809 ± 0.038 |

* RSCU, relative synonymous codon usage; AA, amino acid; Cod, codons; HC, human cells; CoV-2, SARS-CoV-2; SARS-CoV; MERS, MERS-CoV; **, termination codons. Highly increased codons in SARS-CoV-2 with respect to human (Δ ≥ 0.30) are shown underlined and in italics. Underrepresented codons are shown in bold.

On January 30th 2020, the World Health Organization declared the current SARS-CoV-2 outbreak a public health emergency of international concern (WHO, 2020a). To gain insight into the biology and
evolution of emerging SARS-CoV-2, a comprehensive analysis of genome composition, codon and amino acid usage of βCoV strains isolated in China from humans, bats, civets and ferret hosts was performed, including SARS-CoV-2 strains recently isolated from current outbreak.

The results of these studies revealed that SARS-CoV-2 strains enrolled in these analyses have a distinct genome composition in relation to other βCoV strains isolated from human (SARS-CoV), bats, civets and ferrets (see Fig. 1). This is in agreement with very recent studies revealing that SARS-CoV-2 is sufficiently divergent from SARS-CoV to be considered a new human-infecting βCoV (Lu et al., 2020; Zhu et al., 2020). This distinct genomic composition is also reflected in its codon and amino acid usage patterns (see Fig. 2). Moreover, correlation distances and average linkage suggests a closer relation among SARS-CoV-2 and bats βCoV isolated in China and a more distant relation to SARS-CoV or SARS-like βCoV isolated from civets or ferrets (see Fig. 2). This is in agreement with recent results suggesting that bats might be the original host of this virus, an animal sold at the seafood market in Wuhan (Lu et al., 2020; Chan et al., 2020b). This speaks of the importance of bats as a reservoir of potential emerging CoV. Moreover, significant numbers of new βCoV have been discovered in Chinese bats species, particularly Rhinolophus affinis (Lin et al., 2017). Interestingly, a clear degree of variation in codon and amino acid usage was observed among βCoV isolated from bats included in these studies (see Fig. 2). Nevertheless, bats might represent an intermediate host facilitating the emergence of the SARS-CoV-2 in humans (Wong et al., 2019). More studies will be needed to address this important issue.

In these studies, a biased nucleotide composition was found for SARS-CoV-2 genome (Fig. 3). This bias can also have a major influence on derived parameters, as previously demonstrated for other CoVs (Berkhout and Van Hemert, 2015). This is in agreement with the results of this work, since the nucleotide composition of SARS-CoV-2 has a strong influence in the codons that are used by this virus for the translation of its RNA genome. Previous studies on codon usage in RNA viruses have shown that mutational pressure is the major factor in shaping codon usage patterns in comparison with natural selection (Jenkins and Holmes, 2003; Wang et al., 2011). Although mutational pressure is still a major driving force, it is certainly not the only evolutionary force that might be considered in RNA viruses. In these studies, a significant A genomic content was found in SARS-CoV-2 in comparison with other human CoVs (Fig. 3). Previous studies done in Human Immunodeficiency virus (HIV) found that this virus has an A-rich genome and this property has been proposed to help the virus to avoid recognition by the innate immune system (Vabret et al., 2012). This could provide a strong selective pressure on retroviruses as well as many other RNA viruses, including CoVs (Kinder and Thiel, 2014; van Hemert et al., 2014). Moreover, previous studies done in HIV suggest that an RNA genome with A-rich domains may provide a molecular signature that is recognized during virus replication (van Hemert et al., 2013). In the context of CoV, A-rich regions, like the Transcription Regulation Sequence (TRS), which is involved in transcription of subgenomic RNAs have been established (Pyrc et al., 2004). This suggests that nucleotide bias may serve distinct biological function in SARS-CoV-2 as well as in other CoVs and is in direct relation to the characteristic codon usage of these viruses (Berkhout and Van Hemert, 2015).

A mean ENC value of 48.54 ± 2.34 was obtained for SARS-CoV-2 strains enrolled in these studies, suggesting that the overall codon usage among SARS-CoV-2 is similar among themselves and slightly biased. This is in agreement with mean ENC values obtained for other CoV, like SARS-CoV (ENC = 48.99) (Gu et al., 2004); Bovine Coronavirus (BCoV) (ENC = 43.78) (Castells et al., 2017); MERS-CoV (ENC = 55.50) (Alnazawi et al., 2017) or Avian CoV (ENC = 51.33) (NSP2; Brandao, 2013). ENC-GC3s plot of the values obtained for SARS-CoV-2 revealed that all spots cluster below the expected curve, suggesting that G + C compositional constraints play a role in SARS-CoV-2 (see Fig. 3).

In these studies, significant differences in RSCU frequencies among SARS-CoV-2 and human cells were found (Table 1). A strong bias toward A and U ending codons was found. This in agreement with very recent studies revealing a significant predominance of A and U at third codon positions in CoV genomes (Sheikhi et al., 2020; Kandeel et al., 2020). These results also suggest that these differences are related to codon usage preferences. Previous studies have shown that both cytosine deamination and selection of CpG-suppressed clones are the major factors that shape codon bias in CoV genomes (Woo et al., 2007).

MERS-CoV codon usage revealed a bias among hydrophobic amino acids, being CCG (Pro) and GUU (Val) the least and most frequently used codons (Chen et al., 2017). The same results were found in this work for SARS-CoV-2 ORFs, since CCG y GUU resulted to be the least and most frequently hydrophobic codons (11 and 352 times, respectively). Regarding hydrophilic amino acids, the least and most frequently used codons in MERS-CoV were CCG (Arg) and GAU (Asp), respectively (Chen et al., 2017). These codons were also the least and most frequently hydrophilic codons used in SARS-CoV-2 ORF’s (11 and 310 times, respectively). Similarly, CgP and UpU dinucleotide frequencies resulted to be the lowest and highest frequencies found in MERS-CoV (Chen et al., 2017). The same results were obtained in these studies on SARS-CoV-2, since CgP and UpU were the lowest and highest dinucleotide frequencies found (0.22 and 1.96, respectively) (see Supplementary material Table 2). Again, these analyses revealed that genomic composition affects the codon usage pattern in both, MERS-CoV and SARS-CoV-2 viruses.

5. Conclusions

The results of these studies revealed that SARS-CoV-2 strains enrolled in these analyses have a distinct genome composition in relation to other βCoV strains. This distinct genomic composition is also reflected in its codon and amino acid usage patterns. Most of the highly frequent codons are A- and U-ending, which strongly suggests that mutational bias is the main force shaping codon usage in this virus. Significant differences in RSCU frequencies among SARS-CoV-2 and human cells were found. These differences are due to codon usage preferences.

Funding

This research was funded by Agencia Nacional de Investigación e Innovación; Comisión Sectorial de Investigaciones Científicas, Universidad de la República, Uruguay, through Grupos I + D grant and PEDECIBA, Uruguay.

CRediT authorship contribution statement

Fernando L. Tort: Visualization, Investigation, Data curation.
Matías Castells: Visualization, Investigation, Data curation.
Juan Cristina: Conceptualization, Methodology, Data curation, Writing - review & editing.

Declaration of Competing Interest

None.

Acknowledgements

We acknowledge Drs. Pilar Moreno, Gonzalo Moratorio and Rodney Colina for critical reading of this work.

Appendix A. Supplementary data

Supplementary material related to this article can be found, in the online version, at doi:https://doi.org/10.1016/j.virusres.2020.197976.
