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A comparative genomics-based study of positive strand RNA viruses emphasizing on SARS-CoV-2 utilizing dinucleotide signature, codon usage and codon context analyses

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\textbf{ARTICLE INFO}

Keywords:
SARS-CoV-2
Codon usage bias
Codon context
Positive strand RNA virus
Coronaviruses
COVID-19

\textbf{ABSTRACT}

The novel corona virus disease or COVID-19 caused by a positive strand RNA virus (PRV) called SARS-CoV-2 is plaguing the entire planet as we conduct this study. In this study a multifaceted analysis was carried out employing dinucleotide signature, codon usage and codon context to compare and unravel the genomic as well as genic characteristics of the SARS-CoV-2 isolates and how they compare to other PRVs which represents some of the most pathogenic human viruses. The main emphasis of this study was to comprehend the codon biology of the SARS-CoV-2 in the backdrop of the other PRVs like Poliovirus, Japanese encephalitis virus, Hepatitis C virus, Norovirus, Rubella virus, Semliki Forest virus, Zika virus, Dengue virus, Human rhinoviruses and the Betacoronaviruses since codon usage pattern along with the nucleotide composition prevalent within the viral genome helps to understand the biology and evolution of viruses. Our results suggest discrete genomic dinucleotide signature within the PRVs. Some of the genes from the different SARS-CoV-2 isolates were also found to demonstrate heterogeneity in terms of their dinucleotide signature. The SARS-CoV-2 isolates also demonstrated a codon context trend characteristically dissimilar to the other PRVs. The findings of this study are expected to contribute to the developing global knowledge base in countering COVID-19.

1. Introduction

The single strand RNA viruses with positive polarity (PRVs) are a group of viruses which are particularly infamous for their high degree of infectivity and incorporates more than one-third of all virus genera (Ahlquist et al., 2003). The viruses within this group include the Poliovirus, Dengue virus, Zika virus, Japanese encephalitis virus, Hepatitis C virus, Norovirus, Rubella virus, Coronavirus, and others. The most recent addition to this list is the novel Severe acute respiratory syndrome related coronavirus 2 (SARS-CoV-2).

This study was designed to make a comparative analysis of the dinucleotide signature and codon biology of the PRVs with special emphasis on the most recently discovered novel SARS-CoV-2 strain, since from the genomic view point, codon usage pattern plays a vital role in deciphering the basic biological and evolutionary processes (Wang and Chen, 2013). The study of codon usage bias in viruses have been carried out to understand host adaptation of the virus, host-pathogen interaction, host immune system evasion (Khandia et al., 2019; Butt

\textbf{Abbreviations:} PRV, Positive strand RNA Virus; SARS-CoV-2, Severe Acute Respiratory Syndrome Coronavirus 2; SARS, Severe Acute Respiratory Syndrome; MERS, Middle East Respiratory Syndrome; HCV, Hepatitis C Virus; CRS, Congenital Rubella Syndrome; CNS, Central Nervous System; PCA, Principal Component Analysis; Ne, Effective Number of Codons; GC1, Guanine and Cytosine content on the first position of the codon; GC2, Guanine and Cytosine content on the second position of the codon; GC3, Guanine and Cytosine content on the third position of the codon; RSCU, Relative Synonymous Codon Usage; CAI, Codon Adaptation Index; SCUO, Synonymous Codon Usage Order; Fop, Frequency of optimal codons; CUB, Codon Usage Bias; RCDI, Relative Codon De-Optimization Index; SiD, Similarity Index; MFE, Minimum Free Energy.

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https://doi.org/10.1016/j.genrep.2021.101055
Received 9 December 2020; Received in revised form 20 January 2021; Accepted 9 February 2021
Available online 17 February 2021
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The PRVs include some of the most pathogenic viruses, and PRVs like Dengue virus, Zika virus and Middle East respiratory syndrome-related coronavirus pose a serious threat to public health (Brechot et al., 2019; Neelam and Kikkert, 2019). PRVs either rapidly infect new host or develop mechanisms to tackle host defence machineries (Hilleman, 2004; Beachboard and Horner, 2016; Garcia-Sastre, 2017). These viruses are also thought to hide viral replication intermediates in intercellular replication factories to escape from host defence pathway (Igarah and Lohmann, 2015; Overby et al., 2010). Extensive studies have unravelled the fact that complete ability to escape from these defence mechanisms may be one of the potent reasons that helps PRVs like Dengue, Hepatitis C, Zika, and some coronaviruses to cause disease outbreaks in humans (Chen et al., 2017; Uno and Ross, 2018; Gokhale et al., 2014; Kindler et al., 2016; Pardy et al., 2019). The single strand RNA with positive polarity in both Middle East respiratory syndrome-related coronavirus and Severe acute respiratory syndrome-related coronaviruses encode structural proteins like membrane protein (M), envelope (E), nucleocapsid (N), non-structural polyproteins and spike (S) protein which along with glycoprotein and enzymes controls severity in disease exaggeration (Subbaram et al., 2017; Phan, 2020; Ortega et al., 2020).

The PRVs included in this study along with SARS-CoV-2 are very potent human pathogens. These include the Poliovirus, Japanese encephalitis virus, Hepatitis C virus, Norovirus, Rubella virus, Semliki Forest virus, Zika virus, Dengue virus, Human rhinoviruses and the Betacoronaviruses from sub genera Embecovirus, Hibecovirus, Merbecovirus, Nobecovirus and Sarbecovirus. Of all the known RNA viruses, coronaviruses possess the largest genomes (Woo et al., 2009; Almazan et al., 2014; Ruan et al., 2003), and its members are spread across the phylum Riboviriota within the different genera of the family Coronaviridae under the order Nidovira. (Almazan et al., 2014; Corman et al., 2018; Gorbalenya et al., 2020; Cui et al., 2019). Most of the Betacoronaviruses included in this study are zoonotic in origin out of which three have at some point of time jumped into humans (Rehman et al., 2020; Menczery et al., 2020; Woo et al., 2006). The Betacoronaviruses that have jumped into humans cause severe acute respiratory syndrome (SARS), Middle East respiratory syndrome (MERS) and COVID-19. These viruses are members of the sub-genera Sarbecovirus and Merbecovirus respectively which causes illness linked to the respiratory tract and in extreme cases death due to pneumonia and even multi organ failures (Zaki et al., 2012; Assiri et al., 2013; Farcas et al., 2005; Huang et al., 2005; Yin et al., 2004; Napoli, n.d.).

Among the other PRVs included in this study, Poliovirus, the causative agent of the communicable disease poliomyelitis (Kitamura et al., 1981; Xu et al., 2019) is a prominent member of Picornaviridae family with a RNA genome of approximately 7500 nucleotides, covalently linked to virus-coded VPg at 5' end and a poly(A) tail at the 3' end (Yogo and Wimmer, 1972; Flanagan et al., 1977; Lee et al., 1977; Racianni and Baltimore, 1981). Another PRV of approximately 7200 bp, included in this study are the human rhinoviruses of the Picornaviridae family. Some of the most significant health threats that arise from the family Flaviviridae and we have included these in our study. The flavivirus Japanese encephalitis virus is responsible for the mosquito borne zoonotic viral disease Japanese encephalitis and it is the most important causative agent behind epidemic viral encephalitis in the Southeast Asian and Western Pacific regions, China, and the Indian subcontinent (Li et al., 2019; Cherian and Walimbe, 2015). Zika virus is an arbovirus from the family Flaviviridae primarily transmitted by Aedes mosquito (Musso and Gubler, 2016; Plourde and Bloch, 2016) causing disease with commonly reported symptoms including rash, fever, arthralgia, myalgia, fatigue, headache, and conjunctivitis. Infection by Zika virus in human cortical neural progenitor cells results into stunted cell growth and transcriptional dysregulation (Tang et al., 2016), neonatal microcephaly (Oliveira Melo et al., 2016). Dengue virus is another mosquito borne Flavivirus transmitted by Aedes aegypti and Aedes albopictus. Although primary infection with Dengue virus may include rash and fever, but many infections are asymptomatic, while secondary infection causes severe complications with mortality rate up to 20% (Uno and Ross, 2018; Guzman and Harris, 2015). Another PRV of the Flaviviridae family, the Hepatitis C virus targets human hepatocytes (Zeisel et al., 2013; Chan and Ou, 2017), and leads to severe liver diseases including cirrhosis and hepatocellular carcinoma. The Hepatitis A virus is another important liver affecting virus which is highly contagious and belongs to the genus Hepatovirus of the family Picornaviridae (McKnight and Lemon, 2019). The Calcivirusidae is represented by Norovirus, infamous for causing acute infectious gastroenteritis with symptoms including rapid onset, abundant vomiting or diarrhoeal disease (Puruta et al., 2003; Maunula et al., 2012; Anttila et al., 2010). The human pathogen Rubella virus of the genus Rubivirus of family Togaviridae (Kanbayashi et al., 2018; Mangala Prasad and Klose, 2017) which is responsible for foetal death or congenital rubella syndrome (CRS) in pregnant women has also been included in this study. Semliki Forest virus, a PRV of the genus Alphavirus from Togaviridae family is reported to cause lethal encephalitis by infection of the central nervous system (CNS) (Atkins et al., 1999).

Now-a-days in the era of next generation sequencing, the rapid progress and robustness in sequence analysis techniques have provided an opportunity to compare genomic sequences on a large scale to unravel many hidden facets of the genome and its evolution. Simultaneously, this enriches the field of comparative genomics-based study to comprehend the pathogenic viruses to a large extent. A slight change in the codon and codon pair usage pattern may influence viral pathogenicity on a massive scale and hence these studies may provide critical pointer in the development of proper vaccine or therapy against the virus (Alexaki et al., 2019; Baker et al., 2015). Moreover, synonymous codon usage study may also unveil information regarding genic evolution and assist to detect the horizontal gene transfer events (Roy-Choudhury and Mukherjee, 2010). Furthermore, translational kinetics can be better understood, and protein structure may be predicted through such codon usage data analysis (Athey et al., 2017; Cheng et al., 2017). Although codon usage analysis of SARS-CoV-2 have been conducted in some isolates (Kandeel et al., 2020; Tort et al., 2020), but till now no study is available where the SARS-CoV-2 and other coronaviruses have been compared with other PRVs to find out how they relate to each other in terms of their genomic codon usage affinities. We have included the whole genomes of many SARS-CoV-2 isolates from different regions of the world available in the publicly available sequence databases to find out if there exist any dissimilarities within the codon and codon pair usage pattern among the different SARS-CoV-2 isolates both on the genomic scale as well as gene wise, and how these isolates relate in terms of dinucleotide signature, codon usage and codon pair usage with respect to the other PRVs. We have emphasised on the genomic signatures of all the viral genomes with reference to their dinucleotide composition, codon usage bias, amino acid usage trend and codon context pattern. We have also tried to comprehend the codon usage pattern of all the viruses included in this study by comparing them alongside with highly expressed human genes such as those encoding the human ribosomal proteins. We anticipate the findings of this study will be useful in understanding the biology of SARS-CoV-2 virus to a certain extent and contribute towards the knowledge base that is being developed to counter SARS-CoV-2.
2. Materials and methods

2.1. Retrieval of whole genome sequence data

The completed whole genome sequences of SARS-CoV-2 and 49 other PRV genomes (primarily reference genomes) were downloaded from GenBank (Benson et al., 2013) and NCBI Virus, a virus variation resource (Hatcher et al., 2017). Only complete genomes were included in the study, and those with incomplete sequencing status along with improper annotation, incomplete data regarding location and date of collection were discarded from the study. The different ORFs and coding sequences comprising the genome were also obtained from GenBank and NCBI Virus and the annotation data related to these were thoroughly scrutinised. The presence of additional meta data available with the whole genome data were also utilized for sorting the different ORFs and coding sequences of the different PRVs included in this study. Detailed information regarding the accession number and other features of the viral genomes is given in Supplementary Table 1.

2.2. Dinucleotide frequency abundance and representation

The dinucleotide frequency or dinucleotide abundance in every genome bears a great significance in determining genome pattern and evolution which is specific for each organism. This can be regarded as a genomic signature (Prabha and Singh, 2014) since it varies little despite the diversity between species (Jernigan and Baran, 2002). Dinucleotide frequencies of all the possible 16 dinucleotide combinations were computed for all the genomes included in the study to determine whether there is a preference for specific dinucleotide pairs (Pandit et al., 2013). The statistical over- and underrepresentation of all the 16 dinucleotide combinations was computed for all the PRV genomes as well as the individual coding sequences of the SARS-CoV-2 isolates and compared utilizing a z-score statistic with a base model for the whole genomes and coding sequences, and a codon model and syncodon model for all the individual coding sequences (Palmeira et al., 2006; Gautier et al., 1985; Karlin and Cardon, 1994). The R package ‘seqinr’ was used to perform this analysis. The statistical over- and underrepresentation of dinucleotides in a sequence was further subjected to a multivariate data analysis technique called principal component analysis (PCA) to differentiate the PRV genomes based on their dinucleotide representation and detect a genomic signature both within the genomes and the coding sequences of SARS-CoV-2. The multivariate data analysis was carried out using the R package ‘factoextra’ and ‘FactoMiner’. PCA plots were constructed using the R package ‘ggplot2’.

2.3. Codon usage pattern analysis

To comprehend the codon usage pattern prevailing within the virus genomes, the parameters like effective number of codons (Ne) (Wright, 1990), guanine and cytosine content on the first, second and third position of the codon (GC1, GC2 and GC3 respectively) (Wright, 1990), relative synonymous codon usage (RSCU), gene length, hydrophobicity (Kyte and Doolittle, 1982), codon adaptation index (CAI) (Sharp and Li, 1987), synonymous codon usage order (SCUO) (Wan et al., 2004), and frequency of optimal codons (Pop) (Ikemura, 1985; Xu et al., 2013) were calculated using INCA 2.1 (Supek and Vlahovicek, 2004), CodonW (Peden, 1999), in house developed Perl scripts and the R packages ‘coRdon’ and ‘seqinr’.

The Ne is considered as one of the best quantitative measures to evaluate the degree of bias for the usage of codons. The Ne value ranges from 20 to 61 where lower value indicates higher codon usage bias while higher value denotes reduced codon usage bias (Wright, 1990). In the course of genome evolution, GC content on the third base of codon along with GC1 and GC2 has been reported as a key component in regulating gene expression (Genereux, 2002), while hydrophobicity actually predicts the nature of cellular protein encoded by the gene present within the genome (Saha et al., 2019). RSCU measures the non-uniform usage of synonymous codons in a coding sequence and represents the number of times a codon is used in comparison to the number of times that codon would be observed in case of uniform usage. CAI can also be envisaged as another effective measure for calculating the expression levels of gene sequences (Sharpe and Li, 1987; Prabha et al., 2017). This is an important index for CUB analysis, with values ranging from 0 to 1. This is a well-accepted parameter for estimating the relative adaptation of codon usage of a gene towards the codon usage of highly expressed genes (Ayon et al., 2014). CAI was calculated following an improved implementation (Xia, 2007). The human ribosomal protein coding genes were used as reference for measuring CAI. CAI is of great importance for estimating translational efficiency and predicting cellular protein levels (Baha et al., 2019). SCUO ranges from 0 to 1, and represents the synonymous codon usage bias (Wan et al., 2004) which is mainly based on Shannon Information theory explaining entropy of codon sequences of genes (Beluru and Severson, 2012). Larger value denotes a higher codon usage bias with less entropy. For comparative CUB analysis, this parameter is being widely applied (Prabha et al., 2017). Pop encodes the proportion of optimal codons accounting for all synonymous codons. The value varies from 0 to 1 (Ikemura, 1985). A gene with no optimal codon possess a Pop value of ‘0’ whereas the gene comprising of a significant number of optimal codons possess the value ‘1’ (Nakamura and Ikemura, 1995).

To further explore the codon usage bias in each of the strain and to depict the correlation between Ne and GC3, Ne plot was constructed for each of the gene present within the viral genome. Such a plot is generally used to elucidate the mechanistic forces influencing CUB (Pal et al., 2019). The neutrality plot explaining the interrelationship between different GC attributes like GC1, GC2, GC3 content of the prevailing genes within the genome was also utilized (Xiang et al., 2015). To decipher the genic organization within the viral genome and to investigate the mutation-selection equilibrium in shaping codon usage bias, this serves as an important tool. A plot regression with a slope of 1 indicates complete neutrality while a slope of 0 indicates no effect of directional mutation pressure (Kumar et al., 2016). The GC content in the first two codon position of each genic sequence was plotted against the respective GC3 content and regression values were estimated (Franzo et al., 2018).

2.4. Relative codon de-optimization index (RCDI)

To understand host-virus relationship this index has been proved to be useful as virus evolution occurs along with its host (Puigbo et al., 2010). This can be defined as a measure of codon usage deoptimization by comparing the codon usage of a gene with that of the reference genome. This value indicates the cumulative effects of codon biases on the expression of gene (D’Andrea et al., 2019). The concept of RCDI (Mueller et al., 2006), also explains the rate of viral gene translation where RCDI value closest to 1 indicates higher adaptation of a virus with its host. Similarly, it also predicts higher translation rate within the host genome (Khandia et al., 2019; Mueller et al., 2006).

2.5. Similarity index (SID)

The similarity index value (Zhou et al., 2013) is a codon usage parameter which evaluates the host potentiality and is often used to decipher the influence of codon usage bias on host genome (Zhou et al., 2019; Cao et al., 2018). SID is also suggested to be helpful in obtaining a more accurate assessment of adaptation between the virus and its host (Silverj and Rota-Stabelli, 2020). The value ranges from 0 to 1 representing the effect of codon usage on host, and a higher value is suggestive of low codon usage similarity (Silverj and Rota-Stabelli, 2020).
2.6. Minimum free energy (MFE) calculation of mRNA structure

Calculating the minimum free energy released by the mRNA secondary structure during the transcription process, has a great significance in assessing the mRNA stability. The value may range from negative to positive (Ringnér and Krogh, 2005). Higher value indicates comparatively greater loss of energy by the mRNA molecule sustaining a stable mRNA conformation (Deb and Uddin, 2020). The MFE was calculated using Quikfold hosted at DINAMelt Web Server (http://unafold.rna.albany.edu/?q=DINAMelt/Quickfold) (Markham and Zuker, 2005).

2.7. Codon pair analysis and codon context study

This represents the basic preliminary feature of a gene which influence the mRNA decoding fidelity (Moura et al., 2007). Codon pair utilization bias present in each of the different organisms can be better explored through this parameter to infer on phylogeny (Moura et al., 2011). Codon context analysis was carried out using the software ANACONDA 2 to generate the frequency table of codon pair context (Baha et al., 2019), and followed by subsequent statistical analysis, to identify preferred set of codon pairs in a coding sequence (Pal et al., 2015; Nasrullah et al., 2015). Along with codon context analysis, the codon pair ratio or CPk was also calculated for each viral genome to find out how many codon pairs have been utilized in coding the different coding sequences relative to the number of codon pairs combinations that are feasible (Pal et al., 2015).

3. Results and discussion

In RNA viruses, mono and dinucleotide frequencies in genomes or mRNAs may significantly fluctuate (Belalov and Lukashev, 2013). To capture the variation in AU/GC content of all the PRV genomes included in this study (Supplementary Table 2), individual Mann-Whitney Rank Sum Tests utilizing the respective AU and GC content of the viruses was carried out after normality check failed using Shapiro-Wilk test (p < 0.05). The results demonstrated that the difference in the median values between the AU and GC is greater than would be expected by chance, suggesting a significant difference in AU as well as GC content between SARS-CoV-2 and the other PRVs (U = 1024.000; p < 0.01). The SARS-CoV-1 virus demonstrated an average 2.78% less AU content than the SARS-CoV-2 isolates. The highest AU content (67.94%) among all the PRVs considered in this study was observed in Human coronavirus HKU1 (NC_006577), whereas the highest GC content (69.60%) was observed in Rubella virus (NC_001545.2). The average AU and GC content of the SARS-CoV-2 isolates was found to be about 62% and 38% respectively. A graph depicting the relative fluctuations in AU and GC content among the PRVs is given in Fig. 1, and it suggests that in these viruses there is a strict positive correlation between AU and genome size (r = 0.72, p < 0.01) where larger genomes are dominated by AU nucleotides.

3.1. Genomic and genic dinucleotide signature analysis

A genome wide comparison of the relative under and over-representation of the different dinucleotide frequencies of the SARS-CoV-2 isolates and other PRVs was performed (Supplementary Table 3). The variation in dinucleotide composition in all the viruses was studied to find out if there is any bias towards specific dinucleotides which may have arisen due to host specific mutational pressure or inherent features of the virus. It was observed that in all the SARS-CoV-2 isolates there is a significant underrepresentation of CpG dinucleotides which is similar to other RNA viruses like the Influenza A virus (Gu et al., 2019b). This trend was also evident among the other PRVs included in this study except for Rubella virus which did not demonstrate CpG under-representation. Among the SARS-CoV-2 isolates we observed heterogeneity within the distribution of the CpG dinucleotides within the different genomes. The SARS-CoV-2 strains MT246488, MT159713, MT159717, MT246449 and MT066156 demonstrated substantial under-representation of CpG dinucleotides in their genome, whereas the isolates MT020881, MT152824, MT246458, MT246472, MT246478, MT118835, and the isolates from China demonstrated a relatively lesser underrepresentation of the CpG dinucleotides. The AU dinucleotide was also found to be underrepresented along with UA, UC, GG and GA. The dinucleotides AU, UA and UC were also found to be underrepresented in all the other RNA viruses included in this study.

To determine the dinucleotide genomic signature of the viral genomes we designed and executed many PCAs utilizing the relative over and underrepresentation of the 16 different dinucleotide combinations. A genome wise PCA utilizing the dinucleotide representation of all the PRV genomes considered in this study demonstrated the first two dimensions to contribute towards 34% and 20% of the total variance which was the highest among all the dimensions. Analysis of the contributions of the different dinucleotide combinations to the first and second dimensions clearly demonstrated that most of the GC containing dinucleotide combinations like GU, GG, GA and GC play a major role in differentiating the virus genomes (Supplementary Fig. 1). A bi-plot depicting the individual virus genomes and the different dinucleotide combinations (Fig. 2) shows that nearly all the SARS-CoV-2 genomes cluster together on the bi-plot except for a few one and segregate themselves quite prolifically from the rest of the other PRVs included in this study. The other PRVs represented by the genomes of Hepatitis A virus (NC_001489), Zika virus (NC_035889), Dengue virus (NC_001474), Semliki forest virus (NC_003215), Hepatitis C virus (NC_004102), Human rhinovirus (NC_038311), and Rubella virus (NC_001545) were found to occupy discrete locations on the left-hand side of the bi-plot, which is opposite to that of the SARS-CoV-2 genomes suggesting difference in genomic dinucleotide signature. To understand the uniqueness within the genomic dinucleotide signatures of the SARS-CoV-2 genomes, a PCA was carried out utilizing the 16 dinucleotide combinations and grouping the virus genomes based on their geographical location. The PCA bi-plot depicted in Fig. 3 shows that the first two dimensions contributes 16% and 11% of the total variance which is highest among all the dimensions. The dinucleotide combinations UA and UC was found to contribute the greatest in both the dimensions. To further comprehend the trend and uniqueness in the dinucleotide signatures within the genome of the SARS-CoV-2 isolates, a gene wise PCA analysis was carried out, since the sum of genes represents the genome in its near entirety in case of the PRVs. The statistical dinucleotide over- and under-representation of all the individual genes or coding sequences constituting the genome of SARS-CoV-2 were obtained based on the base model, codon model and syncodon model (Palmeira et al., 2006; Gautier et al., 1985; Karlin and Cardon, 1994). The data obtained (Supplementary Table 4) based on these models were analysed utilizing three separate PCAs. The PCA performed on the base model data (Supplementary Fig. 2) suggests that most of the SARS-CoV-2 genes has a characteristic dinucleotide signature. In this case, the first two dimensions accounts for 56.15% and 16.57% of the total variance with the dinucleotide combinations of UA, UC, AU and AG contributing the maximum to both the dimensions. These include the genes S, N, orf6, E, orf1ab and orf1b. The orf7a, orf3a and M clusters tightly together at the centre of the bi-plot which suggests that they share similar dinucleotide signature. The orf8 and orf10 also demonstrated some amount of overlap in terms of their gene wise signatures. The orf1ab and orf7b model demonstrated a similar trend, except for orf1ab which was found to be segregated into two different clusters, one being shaped by GU and AC whereas the other is influenced by UG. In depth analysis of the two clusters clearly demonstrated geographical separation where the isolates from South and South-East Asia such as Japan, India, South Korea, Taiwan and Vietnam were found to share the same cluster along with that of Sweden, Australia, Italy and Brazil (Supplementary Fig. 3). The orf1ab from the isolates of Sweden were found to be present in a separate
isolated from Wuhan province of China was found to cluster along with those from Asia suggesting similar dinucleotide signature. PCA analysis based on the syncodon model data that allows for random sequence generation through synonymous codon shuffling (Supplementary Fig. 4) also demonstrated similar trend for orf1ab. Based on the syncodon model data the dinucleotide signature of orf8, orf10, orf7a, orf7b and orf6 was found to be mostly influenced by CG, UA and UC.

3.2. Analysis of codon usage pattern of the PRVs

The results of the genomic codon usage (Nc\textsubscript{Genome}) pattern study (Supplementary Table 5) of the PRVs demonstrated that Rubella virus (Nc\textsubscript{Genome} = 38.41) and Hepatitis A virus (Nc\textsubscript{Genome} = 39.61) has the most codon biased genome, whereas Poliovirus (Nc\textsubscript{Genome} = 53.60), Zika virus (Nc\textsubscript{Genome} = 54.03), Norovirus (Nc\textsubscript{Genome} = 54.67), Semliki forest virus (Nc\textsubscript{Genome} = 55.66), Japanese encephalitis virus (Nc\textsubscript{Genome} = 55.96) and Hepatitis C virus (Nc\textsubscript{Genome} = 56.16) has the least biased genome in terms of codon usage. Among the coronaviruses, Rousettus bat coronavirus, a member of sub-genus Nobecovirus was found to demonstrate the least genomic codon bias (Nc\textsubscript{Genome} = 55.14), whereas Human coronavirus HKU1 (Nc\textsubscript{Genome} = 40.56), a member of sub-genus Embecovirus demonstrated the highest codon bias. The genomic codon bias of the SARS-CoV-2 isolates was found to range between 46.61 and 49.0 with the isolate MT049951 from China and MT233519 from Spain demonstrating the highest and lowest codon bias, respectively. This suggests that the SARS-CoV-2 along with the other coronaviruses has lower genomic codon bias, since an Nc value greater than 40 indicates low codon bias (Xu et al., 2013; Messier and Stewart, 1997). But when compared to the remaining PRVs except Rubella virus and Hepatitis A virus, the genomic codon bias is relatively higher.

The effect of GC3 on the codon usage pattern of the different viruses considered in this study was further explored with the help of a Nc plot. An Nc plot depicting the association between Nc and GC3 of different genes of the PRVs is shown in Fig. 4 which suggests that relative to the human ribosomal protein genes (which are generally highly expressed) (Lin et al., 2002; Luo et al., 2016) the positioning of the SARS-CoV-2 genes on the Nc plot is quite discrete. Nearly all the SARS-CoV-2 representatives on the Nc plot showed similar clustering pattern. Relative to SARS-CoV-1, the SARS-CoV-2 isolates demonstrated a different clustering pattern. A detailed study of Nc plot clearly shows that there is little overlap in the codon usage pattern among the coronaviruses, except for Middle East respiratory syndrome-related virus which resembles the aggregation pattern of SARS-CoV-2 on the Nc plot to a large extent. The PRVs (excluding all the different coronaviruses) included in this study demonstrated a Nc-GC3 relationship which was found to be significantly different from all the Betacoronaviruses but resembled that of the human ribosomal genes to a greater extent.

3.2.1. Intra and inter genic codon usage analysis of different SARS-CoV-2 genomes

The codon usage pattern of all the genes constituting the SARS-CoV-2 genomes were analysed intricately (Supplementary Table 6). The codon bias of the genes coding for the structural components of the virus demonstrated that the structural component coding genes have low codon bias with gene E being the least biased one (Nc = 61) followed by gene M, N and S (Nc ~ 44.1). A similar trend was also observed for the other ORFs. A correlation of Nc with sequence length demonstrated a significant negative correlation in the case of orf1ab (ρ = −0.672, p < 0.01). This suggests that codon bias is positively correlated with length in these ORFs. The correlation between Nc and GC1 demonstrated a significant negative correlation in the case of orf1ab (ρ = −0.737, p < 0.01), S (ρ = −0.42, p < 0.01). A positive correlation was observed in case of orf8 (ρ = 0.869, p < 0.01) which is entirely contrary to the prevalent trend. In terms of association between Nc and GC3, significant positive correlation was observed in most of the cases for orf3a and orf8.

![Fig. 1. A parallel plot depicting the relation between genome size and AU/GC content of the positive strand RNA viruses (PRVs) included in this study.](image-url)
The relationship between \( \text{Nc} \) and GC3 was further explored to better understand the role of GC3 in determining the codon usage bias trend in all the 12 coding sequences constituting the genome of SARS-CoV-2. All the ORFs except orf6 and orf1α were found to cluster together in proximity on the Nc plot shown in Fig. 5. Apart from gene E and orf10, the other genes were found to lie below the null hypothesis curve suggesting the mechanistic effect of translational selection as a determinant of codon bias. The coding sequences of gene E was found to cluster tightly and lie above the null hypothesis curve along with orf10 indicating the presence of mutational pressure.

To have a better insight on the effect of GC3 on codon usage bias, the relation between GC3 and SCU0 was analysed (Fig. 6), since GC3 has been found to depict a quantitative relationship with SCU0 in microbial and archaeal genomes, and in some mammals (Wan et al., 2004; Pal et al., 2015; Zeeberg, 2002). Although a linear relationship between GC3 and SCU0 was evident in case of orf1α (\( R^2 = 1.0, F = 14,398,631.11 \)) and orf7a (\( R^2 = 0.80, F = 482.56 \)), such a trend was absent in most of the other coding sequences. In case of orf6 (\( R^2 = 0.99, F = 623.84 \)) and orf10 (\( R^2 = 1.0, F = 5.535E+14 \)), a quadratic equation was found to best describe the association between GC3 and SCU0. A linear relationship was also found to exist in case of gene S (\( R^2 = 0.5, F = 124.85 \)) and N (\( R^2 = 0.42, F = 88.02 \)). No distinct association was found to exist between GC3 and SCU0 of orf8, and M.

Analysis of neutrality plots depicting the relationship between GC1/ GC2 with GC3 of the genes constituting the SARS-CoV-2 genome (Fig. 7) demonstrated a linear relationship in the case of orf1α and orf1α. No such prominent relationship was found to exist in the case of the other genes (Supplementary Table 7). In a neutrality plot, a slope of 1 indicates complete neutrality while a slope of 0 indicates no effect of directional mutation pressure (Kumar et al., 2016).

The CAI which is a measure of the potential expression of genes was found to be relatively low for the genes N, M, E and orf10 whereas the gene coding for the surface glycoprotein and other non-structural genes had higher CAI values. To find out the effect of codon usage bias on gene expression, Spearman rank correlation was performed between CAI and the other codon usage bias determining parameters. Our results demonstrated significant negative correlation between Nc and CAI for most of the sequences such as orf7a, orf7b, orf1α, M and N. On the other hand, an inverse trend was observed in the case of orf1α, orf6, orf8, orf3a and S. In terms of association, CAI was found to be significantly positively correlated for majority of the coding sequences like orf7a (\( \rho = 0.819; p < 0.01 \)), orf6 and E (\( \rho = 0.77; p < 0.01 \)), orf3a (\( \rho = 0.656; p < 0.01 \)) and orf7b (\( \rho = 0.61; p < 0.01 \)). Significant negative correlation was observed in case of orf1α (\( \rho = -0.62; p < 0.01 \)) and M (\( \rho = -0.70; p < 0.01 \)). In our study with the SARS-CoV-2 genomes, Fop was found to correlate the best with CAI. Apart from orf1α and orf1α, Fop was found to correlate significantly in a positive manner with CAI for gene S, orf8, E, orf6, orf3a, etc. The only significant negative association between Fop and CAI was evident in case of orf1α (\( \rho = -0.66; p < 0.01 \)). A significant positive correlation was observed between CAI and hydrophobicity in orf7a, orf1α, S, orf3a and N while the rest of the sequences were found to be negatively correlated. The orf10 and orf1α did not depict any statistically significant correlation. The GC content at different codon positions like GC1, GC2 and GC3 was found to correlate significantly with CAI in orf1α whereas in rest of the coding sequences there was no substantial association between these parameters suggesting GC1, GC2 and GC3 do not play a significant role in determining the potential expression levels of the coding sequences.

### 3.2.2. Association between dinucleotide frequency and codon usage pattern

The correlation between CAI and dinucleotide abundance was performed to detect the effect of the latter on the potential expression level...
of a gene. The dinucleotide combinations UA ($\rho = -0.266; p = 0.003$), UC ($\rho = -0.266; p = 0.003$), UU ($\rho = -0.240; p = 0.007$), CC ($\rho = -0.217; p = 0.016$) and GC ($\rho = -0.216; p = 0.017$) was found to be negatively associated with CAI whereas CG and GC was found to positively influence the CAI levels. This suggests that the abundance of certain dinucleotide combinations within the coding sequences are responsible to a certain extent in determining the codon usage pattern that in turn influences gene expression.

3.2.3. Codon usage bias and mRNA minimum free energy (MFE)

The mRNA MFE is an indicator of stability of the mRNA molecule and plays an important role in translation and product formation. Our results suggest that on a gene-to-gene basis in the SARS-CoV-2 isolates, the genic GC content has a direct bearing on the stability of the mRNA positively in case of gene M ($\rho = 0.81, p < 0.01$) and orf8 CDS ($\rho = 0.36, p < 0.01$), whereas in orf3a ($\rho = -0.90, p < 0.01$) and gene E ($\rho = -0.70, p < 0.01$) an anti-correlation was observed. The other coding sequences did not show any relation with GC content in terms of mRNA stability. The mRNA MFE was also not found to correlate significantly with CAI in about all the coding sequences except orf6, orf7a and orf10. For orf3a ($\rho = 0.34, p < 0.01$), orf8 ($\rho = 0.32, p < 0.01$) and E ($\rho = 0.58, p < 0.01$) a significant positive correlation was detected whereas the structural genes S ($\rho = -0.31, p < 0.01$) and M ($\rho = -0.81, p < 0.01$) depicted a negative correlation. Except for orf8, the Nc ($\rho = 0.44, p < 0.01$) was not found to correlate significantly with MFE, but the SCUO was found to dictate codon usage bias to some extent in orf3a ($\rho = 0.52, p < 0.01$), orf8 ($\rho = -0.42, p < 0.01$), gene S ($\rho = -0.30, p < 0.01$) and M ($\rho = -0.24, p < 0.01$).

3.3. Analysis of RSCU

3.3.1. RSCU analysis of SARS-CoV-2 genes

The analysis of RSCU data (Supplementary Table 8) of the genes coding for the structural proteins of the SARS-CoV-2 isolates showed gene specific preference towards certain codons coding for the different amino acids. For glycine, the gene E and S demonstrated a strong bias towards GGU whereas GGC and GGA was preferred in gene N and M. In case of alanine, GCU was universally found to be the most and preferred codon in genes S, M, and N. Out of the six synonymous codons of leucine, CUU was found to be the most preferred by all the structural genes except N. In case of isoleucine, out of the 3 synonymous codons of leucine, CUU was found to be the most preferred by all the structural genes except N. In case of isoleucine, out of the 3 synonymous codons, preference towards the use of AUU in all the structural genes was visualized barring E where all the three synonymous codons were equally employed without any bias. Of the two synonymous codons of phenylalanine, UUC was the most preferred whereas UUU was the least preferred codon in E, M and N, while S exhibited an entirely opposite scenario. For tyrosine, UAC was found to preferred in E, M and N gene whereas this was UAU in S. In case of arginine, the different structural component coding genes depicted a heterogenous choice of codons. In case of the polar neutral amino acids like serine, threonine, asparagine, glutamine and cysteine, somewhat similar codon usage pattern has been observed in the structural genes.

Fig. 3. A PCA bi-plot showing the segregation of the SARS-CoV-2 isolates based on the over and underrepresentation of the 16 dinucleotide combinations. The x-axis represents the 1st dimension which accounts for 16% of the total variance and the y-axis represents the 2nd dimension accounting for 11.1% of the total variance.
Fig. 4. A Nc-plot depicting the relationship between GC content at the 3rd position of codon or GC3 (x-axis) with effective number of codons or Nc (y-axis) of all the genes present in the genomes of the positive strand RNA viruses (PRVs) included in this study. The dashed blue line represents the null hypothesis curve which suggests that codon usage bias is solely due to mutation and not selection (Wright, 1990).

Fig. 5. A Nc-plot depicting the relationship between GC content at the 3rd position of codon or GC3 (x-axis) with effective number of codons or Nc (y-axis) of all the genes present in the genomes of the SARS-CoV-2 isolates included in this study. The dashed blue line represents the null hypothesis curve which suggests that codon usage bias is solely due to mutation and not selection (Wright, 1990).
### 3.3.2. Comparative RSCU analysis of SARS-CoV-2 with human ribosomal protein coding genes

A comparison of the highly expressed ribosomal protein coding genes of humans with that of SARS-CoV-2 demonstrated that for glycine, GGC is the preferred codon in most of the human ribosomal protein coding genes which was found to be in accordance with that of orf10, orf7a and N gene of SARS-CoV-2. Similarly, for phenylalanine and tyrosine, a bias towards the use of codons UUC and UAC respectively was observed in most of the human ribosomal protein coding genes and SARS-CoV-2 structural component coding genes. This trend was also evident in the usage of the codon CAC for histidine. For the amino acids proline, leucine, alanine, isoleucine, valine, and arginine, no significant similarity between the codon preference of human ribosomal protein coding genes and SARS-CoV-2 genes was found to exist. Furthermore, we observed that the most preferred codons for lysine, aspartate, glutamate, and cysteine in the different SARS-CoV-2 genes are least preferred in the different human ribosomal protein coding genes. These inferences were found to be in accordance with the different tRNA species predictions of the human genome (Chan and Lowe, 2008; Chan and Lowe, 2015). The preference of certain codons like GGC for glycine, UUC for phenylalanine, UAC for tyrosine and CAC for histidine was observed in both human and SARS-CoV-2 genes.

### 3.3.3. Comparative RSCU analysis of SARS-CoV-2 with other PRVs

In the rest of the PRVs considered in this study, except Hepatitis C virus, *Norovirus*, and Rubella virus the codon GGA was found to be preferred for glycine by all the seven PRVs (like Dengue virus, Zika virus, Hepatitis A virus, and Japanese encephalitis virus) which is in line with that of M gene of SARS-CoV-2. Similarly, the preferential utilization of the codon GGC for glycine in Hepatitis C virus, *Norovirus*, and Rubella virus was also found to be inline with that of orf10, orf7a and N gene of SARS-CoV-2. In case of alanine, the optimal usage of GCU in Hepatitis A virus (54%) and Human rhinovirus (45%) was noticed to be in accordance with that of S, M, and N gene of SARS-CoV-2. The preferential usage of AUU for isoleucine in Hepatitis A virus (66%) and *Norovirus* (53%) was also in accordance with the three structural genes of SARS-CoV-2. Besides, 80% of the studied PRVs were found to prefer UUC codon for phenylalanine and UAC for tyrosine, while 60% of the PRVs were observed to preferentially utilize CAC for histidine which is in line with our observation for SARS-CoV-2. Within the studied PRVs, only Hepatitis A virus was found to completely avoid GCC for alanine and CGG for arginine and in most situations, and its preferential codon usage pattern was found to differ somewhat with respect to the other PRVs.

### 3.4. Analysis of SiD values in the different PRVs

The analyses of SiD values of all the viruses in this study clearly indicates the presence of the SARS-CoV2 among with the other viruses which demonstrates optimisation of codon pattern to tune itself with the host human genome. The SiD value of Rubella virus was the lowest among all the PRVs while Human rhinovirus demonstrated the highest SiD score.
3.5. Analysis of codon context pattern

The SARS-CoV-2 isolates were found to be biased towards the utilization of valine-initiated (GUU-UUA, GUU-GAA, GUU-UAU, GUU-GUA) and glycine-initiated codon pairs (GGU-GUU, GGU-GAU, GGU-AAA, GGU-GGU). Glutamate (GAA-GAA, GAA-GCU, GAA-ACU, GAA-GGU), alanine (GCU-GUU, GCU-UUA, GCU-UUU, GCU-UGU) and isoleucine (AUU-GUU, AUU-GCU, AUU-AAA, AUU-CUU, AUU-CAA) initiated codon pairs were also observed in higher frequency. In general, guanine initiated and uracil ending codon pairs dominated the entire genome sequence. A genome wise comparison of codon pair data of the other PRVs revealed near universal preference for ACU-GAU, AAG-AAA, AAU-GAU, AAG-AAG and UAC-AAG codon pairs. Specific lysine-initiated codon pairs like AAA-CAU, AAA-CAA, AAA-AUU, AAA-AUG; asparagine-initiated codon pairs like AAU-GGU, AAU-UUU; valine ending codon pairs like GGU-GUU, GAA-GUU, UCU-GUU, GAA-GUU;
glutamate dominated codon pairs like GAA-GUU, GAA-GAU, GAA-GAA and GUU-UAU, UAU-UAU were found to be preferred in most of the coronaviruses. UCG-initiated codon pairs were least preferred among all these viral genomes. “Asparagine-pattern” (Pal et al., 2015) and valine initiated codon context pairs like GUU-UAU, GUU-GUU, GUU-GUG, GUU-GGU, GUU-GCU, GUU-GAU, GUU-AAC, GUU-AAU, GUU-AAA were manifested in most of the coronavirus species. Some aspartate-initiated codon pairs such as GAU-UUU was found be preferred by Bovine coronavirus, Human coronavirus OC43 and Human coronavirus HKU1. The GAU-GUU codon context pair was found to be extensively used by Bovine coronavirus, Human coronavirus HKU1, Rabbit coronavirus HKU14, Rat coronavirus Parker, Middle East respiratory syndrome-related coronavirus and Human rhinovirus. In SARS-CoV-1, the leucine-initiated codon pairs (CUC-AAA, CUC-AAC, CUC-AAG, CUC-ACU, CUC-AUG, CUC-AAU, CUC-AAU, CUC-AAA) were found to be preferred. Threonine and phenylalanine-initiating codon pairs were also found to be widely preferred by the coronaviruses.

The codon pair analysis data (Supplementary Table 9) of the SARS-CoV-2 isolates demonstrated the presence of five different types of CP_R clusters based on a k-means clustering (Supplementary Fig. 5). This suggest that there is a certain amount of heterogeneity within the SARS-CoV-2 isolates in the utilization pattern of the different codon pairs. When compared to the other PRVs it was observed that the coronaviruses have a relatively higher CP_R compared to all the other PRVs. Lower CP_R values was evident in Hepatitis A virus (0.3199), Human rhinovirus (0.325) and Rubella virus (0.3632). This is in line with the overall codon usage nature of these viruses which is much more constricted or biased compared to SARS-CoV-2.

To exploit the heterogeneity in codon pair usage of the SARS-CoV-2 isolates, multivariate data analysis in the form of PCA was performed to differentiate the genomes based on their genomic codon pair signature. The PCA analysis of the different SARS-CoV-2 isolates demonstrated that the first two dimensions account for 55.54% and 30.89% of the total variance. The PCA plot (Supplementary Fig. 6) clearly depicts five distinct codon pair signatures that are encircled by different colours. The isolates MT 240479 and MT 121215 were found to occupy characteristically isolated positions on the plot suggesting distinctively different codon pair signature. The isolates NC_045512 and MT 049951 from China was found to club together with the USA based isolates MT 163716 to MT 163719 on the plot, suggesting similarity in terms of their overall codon context. The remaining two clusters were found to be dense, populated by many of the remaining isolates. The magenta cluster was predominated by the isolates from USA (MT 246451 to MT 246490) and Spain (MT 233519, MT 233522 and MT 233523) suggesting they share a similar genomic codon context. Thus, these isolates apart from depicting a similar dinucleotide signature also shares similar codon context pattern. The blue cluster was found to be the most heterogeneous assemblage with the maximum number of isolates from the other geographical regions clubbed together into it. Thus, we find that there is a differentiation in terms of codon context within the genome of SARS-CoV-2 and that is quite in in line with their dinucleotide signature.

To investigate the similarities and dissimilarities in the codon context pattern of the SARS-CoV-2 isolates with respect to the other PRVs included in this study, representatives from the five different clusters obtained previously was taken together with the 26 other PRVs included in this study. The x-axis represents the 1st dimension which accounts for 23.3% of the total variance and the y-axis represents the 2nd dimension accounting for 11.7% of the total variance. The red dashed square depicts the distribution of the SARS-CoV-2 isolates relative to the other coronaviruses (in the blue square) and the other PRVs (depicted by the green square).
included in this study and a PCA was performed. The PCA plot obtained (Fig. 9) demonstrated an interesting trend where it was observed that all the five different SARS-CoV-2 isolates demonstrated a codon context trend characteristically dissimilar to the other PRVs. While many of the other coronaviruses displayed a similar clustering on the plot, the SARS-CoV2 isolates did not overlap even with the other coronaviruses, neither with SARS-CoV-1 or the Middle East respiratory syndrome-related coronavirus. This is a very interesting finding since SARS-CoV-2 is a member of the coronavirus family but displays a codon context signature different from the other PRVs including coronaviruses.

4. Conclusion

The SARS-CoV-2 is a PRV and one among the three coronaviruses that have successfully jumped into human beings so far and have wreaked a havoc. This study was carried out to compare and comprehend the codon biology of the SARS-CoV-2 with respect to the other PRVs. The presence and continuous generation of a colossal amount of genome sequence data of SARS-CoV-2 is making it possible to intricately analyse the SARS-CoV-2 genome and gene features. But in comparison, the amount of genomic sequence data available for the other PRVs is quite low making it difficult to intricately analyse the same. Based on the limited amount of genome sequence data of the other PRVs in the public databases we observed that the codon usage bias analyses of many SARS-CoV-2 genomes and genes overall shows a great deal of heterogeneity in comparison to the other PRVs. In this comparative dinucleotide and codon biology-based analysis of the SARS-CoV-2 virus including other PRVs we found that among the SARS-CoV-2 isolates heterogeneity within the distribution of the CpG dinucleotides exist and SARS-CoV-2 genomes demonstrate a genomic dinucleotide signature different from the rest of the PRVs. In terms of codon usage, SARS-CoV-2 along with the other coronaviruses had relatively greater genomic codon bias compared to most of the other PRVs. Gene specific codon bias was observed in SARS-CoV-2, and the preference of certain codons were observed in both human ribosomal protein and SARS-CoV-2 coding sequences. The SARS-CoV-2 isolates were also found to demonstrate a codon context trend characteristically dissimilar to the other PRVs. RNA viruses are notorious in terms of mutation leading to continuous and ever-changing variability and hence a continuous vigil is required to monitor the genetic changes that occur in these. More studies concentrating on the other PRVs is of utmost necessity since if kept untracked, these viruses can resurface and catch us off-guard causing pandemics and wreaking havoc in the future.

Supplementary data to this article can be found online at https://doi.org/10.1016/j.genrep.2021.101055.

Ethical approval

This article does not contain any studies with human participants or animals performed by any of the authors.

CRediT authorship contribution statement

Jayanti Saha: Conceptualization, Investigation, Methodology, Formal analysis, Writing – original draft, Writing – review & editing. Sukanya Bhattacharjee: Investigation, Methodology, Software, Formal analysis, Writing – original draft, Data curation, Writing – review & editing. Barnan Kumar Saha: Investigation, Formal analysis, Data curation, Writing – original draft. Parimal Mandal: Conceptualization, Investigation, Writing – original draft. Abhik Chatterjee: Supervision, Conceptualization, Methodology, Validation, Software, Writing – original draft. Ayon Pal: Supervision, Conceptualization, Methodology, Validation, Software, Formal analysis, Writing – original draft, Writing – review & editing.

Declaration of competing interest

All the authors declare that they have no conflict of interest.

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