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October 06-29

THE ULTIMATE WEBINAR SERIES IN GENE EXPRESSION STUDIES







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PHYLOGENETIC ANALYSIS OF SARS-COV-2 IN THE FIRST MONTHS SINCE ITS EMERGENCE

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This article has been accepted for publication and undergone full peer review but has not been through the copyediting, typesetting, pagination and proofreading process, which may lead to differences between this version and the Version of Record. Please cite this article as doi: 10.1002/jmv.26545.

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ABSTRACT

During the first months of SARS-CoV-2 evolution in a new host, contrasting hypotheses have been proposed about the way the virus has evolved and diversified worldwide. The aim of this study was to perform a comprehensive evolutionary analysis to describe the human outbreak and the evolutionary rate of different genomic regions of SARS-CoV-2.

The molecular evolution in nine genomic regions of SARS-CoV-2 was analyzed using three different approaches: phylogenetic signal assessment, emergence of amino acid substitutions, and Bayesian evolutionary rate estimation in eight successive fortnights since the virus emergence.

All observed phylogenetic signals were very low and tree topologies were in agreement with those signals. However, after four months of evolution, it was possible to identify regions revealing an incipient viral lineage formation despite the low phylogenetic signal, since fortnight 3. Finally, the SARS-CoV-2 evolutionary rate for regions nsp3 and S, the ones presenting greater variability, was estimated as 1.37×10^{-3} and 2.19×10^{-3} substitution/site/year, respectively.

In conclusion, results from this work about the variable diversity of crucial viral regions and determination of the evolutionary rate are consequently decisive to understand essential features of viral emergence. In turn, findings may allow the first

time characterization of the evolutionary rate of S protein, crucial for vaccine development.

KEYWORDS: SARS-CoV-2, Phylogeny, Evolution, Evolutionary Rate

Introduction

Coronaviruses belong to Coronaviridae family and have a single strand of positivesense RNA genome of 26 to 32 kb in length ^[1]. They have been identified in different avian hosts as well as in various mammals including bats, mice, dogs, etc. ^[2,3]. Periodically, new mammalian coronaviruses are identified. In late December 2019, Chinese health authorities identified groups of patients with pneumonia of unknown cause in Wuhan, Hubei Province, China^[4]. The pathogen, a new coronavirus called SARS-CoV-2^[5], was identified by local hospitals using a surveillance mechanism for "pneumonia of unknown etiology" ^[4,6,7]. The pandemic spread rapidly and more than 28 million confirmed cases and nearly 900,000 deaths was reported in just over an eight months period ^[8]. The rapid viral spread raised interesting questions about the way its evolution is driven during the pandemic. From the SARS-CoV-2 genome, 16 non-structural proteins (nsp1-16), 4 structural proteins [spike (S), envelope (E), membrane (M) and nucleoprotein (N)], and other proteins essential to complete the replication cycle have been translated ^[9,10]. The large amount of currently available information allows knowing, as never before, the real-time evolution history of a virus since its interspecies jump^[11]. Most studies published to date have characterized the viral genome and evolution by analyzing complete genome sequences ^[12,13,14,15]. Despite this, until now, the viral genomic region providing the most accurate information to characterize SARS-CoV-2, could not be established. This lack of information prevents from investigating its molecular evolution and monitoring of biological features affecting the development of antiviral drugs and vaccines. Therefore, the aim of this study was to perform a comprehensive viral evolutionary

analysis in order to describe the human outbreak and the molecular evolution rate of different genomic regions of SARS-CoV-2.

Materials and Methods

Datasets

In order to generate a dataset representing different geographic regions and time evolution of the SARS-CoV-2 pandemic from December 2019 to April 2020, data of all the complete genome sequences available at GISAID (https: //www.gisaid.org /) on April 18, 2020 were collected. Data inclusion criteria were: a.- complete genomes, b.- high coverage level, and c.- human hosts only (no other animals, cell culture, or environmental samples). Complete genomes were aligned using MAFFT against the Wuhan-Hu-1 reference genome (NC_045512.2, EPI_ISL_402125). The resulting multiple sequence alignment (Dataset 1) was split in nine datasets corresponding to nine coding regions: a.- four structural proteins [envelope (E), nucleocapsid (N), spike (S), Orf3a], b.- four nonstructural proteins (nsp1, nsp3, Orf6, and nsp14), and c.- an unknown function protein (Orf8).

More than six thousand SARS-CoV-2 publicly available nucleotide sequences were downloaded. After selection of data according to the inclusion criteria, 1616 SARS-CoV-2 complete genomes were included in Dataset 1. Sequences of this Dataset 1 came from 55 countries belonging to the five continents as follows: Africa: 39 sequences, Americas: 383 sequences, Asia: 387 sequences, Europe: 686 sequences, and Oceania: 121 sequences. After elimination of sequences with indeterminate or ambiguous positions, the number of analyzed sequences for each region was: nsp1, 1608; nsp3, 1511; nsp14, 1550; S, 1488; Orf3a, 1600; E, 1615; Orf6, 1616; Orf8, 1612; and N, 1610. Finally, nucleotide sequences were grouped by fortnight (FN) according to their collection date. Table 1 summarizes the number of sequences per fortnight since the beginning of the pandemic up to FN 8. On the

other hand, Dataset 2 was created using only variable sequences of each region analyzed in Dataset 1. Thus, Dataset 1 was used for the analysis of amino acid substitutions and Dataset 2 was used for the phylogenetic signal analysis and the Bayesian coalescent trees construction.

Phylogenetic signal

To determine the phylogenetic signal of each of the nine generated alignments, Likelihood Mapping analyzes were carried out ^[16], using the Tree Puzzle v5.3 program ^[17] and the Quartet puzzling algorithm. This algorithm allowed analyzing the tree topologies that can be completely solved from all possible quartets of the n alignment sequences using maximum likelihood. An alignment with defined tree values greater than 70-80% presents strong support from the statistical point of view ^[17]. Identical sequences were also removed with ElimDupes (available at https://www.hiv.lanl.gov/content/sequence/elimdupesv2/elimdupes.html) as they increase computation time and provide no additional information about dated phylogeny. The best-fit evolutionary model to each dataset was selected based on the Bayesian Information Criterion obtained with the JModelTest v2.1.10 software ^[18]. *Analysis of amino acid substitutions*

Entropy-One (available at

https://www.hiv.lanl.gov/content/sequence/ENTROPY/entropy_one.html) was used in determining the frequency of amino acids at each position for the nine genomic regions analyzed and evaluating their permanence in the eight investigated fortnights in Dataset 1.

Bayesian coalescence and phylogenetic analysis

To study the relationship among SARS-CoV-2 sequences, nine regions of the viral genome were investigated by Bayesian analyses. Phylogenetic trees were constructed using Bayesian inference with MrBayes v3.2.7a ^[19]. Each gene was

analyzed independently with the same dataset used for the phylogenetic signal analysis so that non-identical sequences were included in the analysis. Analyses were run for five million generations and sampled every 5000 generations. Convergence of parameters [effective sample size (ESS) \geq 200, with a 10% burn-in] was verified with Tracer v1.7.1 ^[20]. Phylogenetic trees were visualized with FigTree v1.4.4.

Evolutionary rate

The estimation of the nucleotide evolutionary rate was made with the Beast v1.10.4 program package ^[21]. Analyses were run at the CIPRES Science Gateway server ^[22]. Three hundred and twelve sequences without indeterminations corresponding to the nsp3 (5835nt) and S (3822nt) genes were randomly selected from Dataset 1. The sequences represent all the fortnights and most of the geographical locations sampled until April 17. Temporal calibration was performed by date of sampling. The appropriate evolutionary model was selected as described above for phylogenetic signal analysis. Nucleotide substitution TIM model was used for nsp3, and HKY model for S. Analysis were carried out under a relaxed (uncorrelated lognormal) molecular clock model as suggested by Duchene & col. ^[23] and with an exponential demographic, proper for early viral samples from an outbreak ^[24]. Independent runs were performed for each dataset and a Markov chain Monte Carlo (MCMC) with a length of 1.3x10⁹ steps, sampling every 1.3x10⁶ steps, was setup. The convergence of the "mean rate" parameter [effective sample size (ESS) \ge 200, burn-in 10%] was verified with Tracer v1.7.1 ^[20]. Additionally, in order to verify the obtained results, 15 independent replicates of the analysis were performed with the time calibration information (date of sampling) randomized as described by Rieux & Khatchikian, 2017^[25]. Finally, the obtained parameters for real data and the randomized replicates were compared.

Results

Phylogenetic signal

Using bioinformatics tools, a phylogenetic signal study was carried out in order to identify the most informative SARS-CoV-2 genomic regions. The likelihood mapping analysis showed that most genes has very poor phylogenetic signal with high values in the central region that represents the area of unresolved quartets (Figure 1). Accordingly, genes could be separated into three groups. A group with little or no phylogenetic signal (E, Orf6, Orf8, nsp1, and nsp14), a second group with low phylogenetic signal (Orf3a and N), and a last group with relatively more phylogenetic signal (S and nsp3) but still low to be considered a robust one (unresolved quartets >40%).

Analysis of amino acid substitutions

The analysis of amino acid substitutions by fortnights was useful to study the viral evolutionary dynamics in the context of the beginning of the pandemic. When analyzing amino acid sequences from different time periods, changes were observed in 5 out of 9 genomic regions and only in 14 out of the 4975 (0.28%) evaluated residues. In most of the regions, except nsp1, nsp14, E, and Orf6, 2 to 6 amino acids emerge since FN3 and remain unchanged until the end of the follow-up period (Table 2). Particularly, in Orf8 region, early selection of two amino acid substitutions (V62L and L84S) was observed on FN2. On the other hand, in the S region, the D614G substitution started with less than 2% in FN3 and FN4 and reached 88% in the last fortnight. In a similar way, the Q57H (Orf3a) substitution increased from 6% to 34% while L84S (Orf8) start to be selected in FN2 and reached 6% at FN8. The R203K and G204R substitutions from the N region emerged in FN4 and increased their population proportion to values greater than 20% towards the end of the follow-up period. Moreover, the emergence of a great number of sporadic substitutions that

remains in the population for a short period (1-3 fortnights) was observed in the nine analyzed regions. Indeed, 333 (6.83%) positions from the total analyzed presented at least one substitution throughout the eight fortnights. Table 3 summarizes the number of variable positions, number of mutations, and number of sequences with mutations by region.

Bayesian coalescence analysis

In this study, trees were performed by Bayesian analysis instead of by distance, likelihood, or parsimony methods. Consistently with the phylogenetic signal analysis, trees for nsp1, E, and Orf6 showed a star-like topology. Nevertheless, different proportions of clade formation could be observed in trees of Orf8, nsp14, Orf3a, N, S, and nsp3 regions (Figure 2). Finally, from the mentioned regions, nsp3 and S showed a better clade constitution. This analysis allowed to differentiate regions displaying a diversification process (nsp3, nsp14, Orf3a, S, Orf8, and N) from those that even after four months showed an incipient one (nsp1, E, and Orf6). Furthermore, this nucleotide analysis is complemented by the previous study of amino acid variations in each region. However, it is important to note that due to the low phylogenetic signal observed for each region, results can only be considered as preliminary.

Evolutionary rate

Nsp3 and S sequences were selected to perform the evolutionary rate analysis since both regions provided the best phylogenetic information among studied regions. The observed evolutionary rate for SARS-CoV-2 nsp3 protein was estimated as 1.37×10^{-3} ³ nucleotide substitutions per site per year (s/s/y) (95% HPD interval 9.16 $\times 10^{-4}$ to 1.91 $\times 10^{-3}$). On the other hand, the corresponding figures for S were estimated in 2.19 $\times 10^{-3}$ nucleotide s/s/y (95% HPD interval 3.19 $\times 10^{-3}$ to 1.29 $\times 10^{-3}$). In both genomic regions, date-randomization analyses showed no overlapping between the

95% HPD substitution-rate intervals obtained from real data and date-randomized datasets. This fact suggests that the original dataset has enough temporal signal to perform analyses with temporal calibration based on tip-dates (Figure 3).

Discussion

The phylogenetic characterization of an emerging virus is crucial to understand the way the virus and the pandemic will evolve. Thereby, a detailed study of the SARS CoV-2 genome allows, on the one hand, to contribute to the knowledge of viral diversity in order to detect the most suitable regions to be used as antivirals or vaccines targets. On the other hand, the large amount of information that has been continuously generated since SARS CoV-2 emergence in human beings is allowing studying its genome and describing the real-time evolution of a new virus like never before.

In the present study, the molecular evolution and viral lineages of SARS-CoV-2 in nine genomic regions, during eight successive fortnights, were analyzed using three different approaches: phylogenetic signal assessment, the emergence of amino acid substitutions, and Bayesian evolutionary rate estimation. In this context, the observed phylogenetic signals of nine coding regions were very low and the obtained trees were consistent with this finding, showing star-like topologies in some viral regions (nsp1, E, and Orf6). However, after a four months evolution period, it was possible to identify regions (nsp3, S, Orf3a, Orf8, and N) revealing an incipient formation of viral lineages, despite the phylogenetic signal, both at the nucleotide and amino acid levels from FN3. Based on these findings, the SARS-CoV-2 evolutionary rate was estimated, for the first time, for the two regions showing higher variability (S and nsp3).

In respect for the phylogenetic signal, several simulation studies has proven that for a set of sequences to be considered robust, the central and lateral areas representing

the unresolved quartets, must not be greater than 40% ^[16]. In this regard, none of the nine analyzed regions has met this requirement. Three regions (E, nsp1, and Orf6) presented values of 100% unresolved quartets. Most regions (nsp14, Orf3a, Orf8, and N) reached values higher than 85%. Only in regions nsp3 and S, the number of unresolved quartets dropped to ~ 60%. Thus, despite being a virus with an RNA genome, the short time elapsed since its emergence, and possibly genetic restrictions, have led to a constrained evolution of SARS-CoV-2 in these months. For this reason, it is expected that trees generated from SARS-CoV-2 partial sequences in the first months of the pandemic are unreliable for defining clades. Therefore, they should be analyzed with caution.

Since Bayesian analysis allows to infer phylogenetic patterns from tree distributions, it represents a more reliable tool to compare different evolutionary behaviors. Bayesian analysis helps to obtain a tree topology that is closer to reality in the current conditions of SARS-CoV-2 pandemic ^[26]. The phylogenetic analysis for nsp1, E, and Orf6 regions confirmed the star-like topologies in accordance to a lower diversification of these regions using the sequences available up to FN8 (Figure 2). Trees generated from nsp14 and Orf8 are at an intermediate point, where the formation of small clusters can be observed. In fact, a mutation at position 28,144 (Orf8: L84S) has been proposed as a possible marker for viral classification ^[27,28]. Finally, trees obtained from regions Orf3a, N, nsp3, and S showed the best clade formation. Indeed, in the most variable regions nsp3 and S, it can be clearly seen that sequences are separated into two large groups. Despite the aforementioned for the nsp3 and S regions, even clusters with very high support values should be taken with precaution and longer periods should be considered to obtain more accurate phylogenetic data. However, even when data are not the most accurate to study the

spread or clade formation ^[29, 30], they provide a good representation of the way the virus is evolving.

The analysis of amino acid frequencies allowed identifying different degree of region conservation throughout the viral genome because of positive and negative pressures. In particular, nsp3, S, Orf8, and N showed some substitutions in high frequencies. This would indicate, as other authors have previously reported, the frequent circulation of polymorphisms due to significant positive pressure ^[13,27,31]. Additionally, since S and N are among the candidates to be used in the formulation of vaccines and antibody treatment, it will be important to monitor these substitutions in different geographic regions in order to improve treatment and vaccination efficacy ^[32,33,34]. In particular, the appearance of the D614G variant in the third week and its rapid increase until reaching an 88% prevalence in the eighth week could reflect an improvement in viral fitness, as it has been previously reported ^[35]. This is supported by studies in SARS CoV showing that predicted S protein domains underwent the most extensive amino acid substitutions and the strongest positive selection ^[36]. Contrarily, in regions nsp1, nsp14, E, and Orf6 no substitutions were selected during

the first 4 months of the pandemic. This would suggest that these regions present constraints to change due to a great negative selection pressure, as it has been recently reported ^[13].

In the present study, the evolutionary rate for SARS-CoV-2 genes was estimated by analyzing a large number of sequences, which were carefully curated and had a good temporal and spatial structure. Additionally, the most phylogenetically informative regions of the genome (nsp3 and S) were used for analysis, reinforcing the results confidence. Previous studies on SARS-CoV-2 have reported similar data ranging from 1.79×10^{-3} to 6.58×10^{-3} s/s/y for the complete genome ^[6,37]. However, in both articles, small datasets of complete genomes were used (N=32 and 54,

respectively). As studies were performed early in the outbreak and due to datasets temporal structure, analysis could have led to less precise estimates of the evolutionary rate ^[23]. Alternatively, another study from van Dorp et al. (2020), analyzing 7,666 sequences, has obtained different results with a remarkably low evolutionary rate (6 x 10⁻⁴ nucleotide/genome/year) ^[15]. However, it is important to consider that van Dorp et al. (2020) estimate the evolutionary rate using the complete genome, including several highly conserved genomic regions, while in our work, the estimation was performed with the most variable regions of the genome. Additionally, tests randomizing the dates of nsp3 and S datasets were carried out; they showed that these partial genomic regions have enough temporal structure and that they are informative, allowing the estimation of evolutionary rates. In this context, our results $(1.37 \times 10^{-3} \text{ s/s/y for NSp3} \text{ and } 2.19 \times 10^{-3} \text{ s/s/y for S})$ are in close agreement with those published for SARS-CoV genome, which have been estimated to range between 0.80 to 3.01 x 10^{-3} s/s/y ^[36,38,39]. In particular, Zhao et al. (2004) estimated a similar evolutionary rate for SARS-CoV S gene ^[39]. Moreover, our estimated values are in the same order magnitude as other RNA viruses ^[40]. Even though we should be cautious with these results interpretation, our date-randomization analysis indicated a robust temporal signal.

In addition, the importance of separately studying the evolutionary rate of the S genomic region arises from the fact that it represents the main target for antiviral agents and vaccines since it includes the SARS-CoV-2 binding receptor domain (RBD), a crucial structure for the virus to enter host cells, and binding site for neutralizing antibodies ^[41]. "Furthermore, a re-infection case occurring 142 days after the first infection episode has been reported. The second infection virus sequence showed 4 changes out of 14 amino acid in the spike protein and two changes in nsp3 ^[42], the two genome genes considered phylogenetically most informative in our work.

Since neutralizing antibodies are targeted against the spike protein ^[43], a high evolutionary rate in this gene can imply changes in the circulating virus and therefore turning it less susceptible to neutralizing antibodies generated during a first infection. In fact, certain mutations in the spike protein, more precisely in the receptor binding and in the N-terminal domain, have been reported to confer a reduced susceptibility to neutralizing antibodies ^[44,45]. For this reason, the evolutionary rate of S and nsp3 genes, reported separately here for the first time, is a crucial issue as it may have implications for vaccines development, vaccine efficacy, or natural re-infections."

Despite limitations of the evolutionary study of an emerging virus, where the selection pressures are still low and therefore its variability low, this work has a strength: the extremely careful selection of a big sequence dataset to be analyzed. First, sequences were selected considering their good temporal signal and their balanced spatial (geographic) distribution. Secondly, attention was paid to eliminate sequences with low coverage and indeterminacies that could generate bias in the phylogenetic analysis of a virus that is beginning to evolve in a new host.

The appearance of a virus means an adaptation challenge. In this sense, both SARS-CoV and SARS-CoV-2 have shown a rapid emergence of several lineages in a short period ^[36,46], reflecting a high adaptability. However, the spike of SARS-CoV-2 binds to the host cell receptor with a 10 to 20-fold greater affinity compared to SARS-CoV and contains a polybasic (furin) cleavage site insertion, which may enhance the virus infectivity ^[47]. Thereby, changes in the S protein constitute an important contribution, turning SARS-CoV 2 to spill over stage and show a significantly higher spread than SARS-CoV and MERS-CoV. Due to this fact, SARS-CoV 2 becomes the most important pandemic of the century. In this context, results obtained in this work about the uneven diversity of nine crucial viral regions and the determination of the evolutionary rate, are decisive to understanding essential features of viral

emergence. Nevertheless, monitoring SARS-CoV-2 population will be required to determine the evolutionary dynamics of new mutations as well as to understand the way they affect viral fitness in human hosts.

Competing interest: On behalf of all authors, the corresponding author states that there is no conflict of interest.

Funding: None

Declaration of Author Contributions

MJP: Data curation, acquisition of data, analysis and interpretation of data, drafting the article, final approval of the version to be submitted.

LM: Data curation, acquisition of data, analysis and interpretation of data, revising the article critically for important intellectual content, final approval of the version to be submitted.

APM: Data curation, Validation, revising the article critically for important intellectual content, final approval of the version to be submitted.

DMF: Data curation, Validation, drafting the article, final approval of the version to be submitted.

GG: Data curation, acquisition of data, analysis and interpretation of data, drafting the article, final approval of the version to be submitted.

FAD: Conception and design of the study, acquisition of data, analysis and interpretation of data, drafting the article, final approval of the version to be submitted.

Acknowledgements

MJP, LM, DMF, and FAD are members of the National Research Council (CONICET). We would like to thank to the researchers who generated and shared the sequencing data from GISAID (https://www.gisaid.org/) and Mrs. Silvina Heisecke from CEMIC-CONICET for providing language assistance.

Data Availability Statement: Data derived from public domain resources

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Fortnight	Date	Median of analyzed sequences		
		(Q1-Q3)		
FN1	12/24/2019 to	15		
	12/31/2019			
FN2	01/01/2020 to	19		
	01/15/2020			
FN3	01/16/2020 to	145 (136-145.5)		
	01/31/2020			
FN4	02/01/2020 to	119 (113-120)		
	02/15/2020			
FN5	02/16/2020 to	258 (247-259)		
	03/02/2020			
FN6	03/03/2020 to	403 (390-406)		
	03/17/2020			
FN7	03/18/2020 to	447 (416-450)		
	04/01/2020			
FN8	04/02/2020 to	199 (197-201)		
	04/17/2020			
TOTAL		1488 to 1616		

 Table 1. Number of SARS-CoV-2 sequences by fortnight (Temporal structure)

FN: Fortnight; Q1=quartile 1, Q3=quartile 3. The total number of sequences is variable depending on the analyzed region (nsp1, 1608; nsp3, 1511; nsp14, 1550; S, 1488; Orf3a, 1600; E, 1615; Orf6, 1616; Orf8, 1612; and N, 1610)

Decien	Amino acid	Amino acid percentage by FN							
Region	substitution	FN1	FN2	FN3	FN4	FN5	FN6	FN7	FN8
nsp3	A58T	0	0	0	1.0	6.0	3.0	3.0	2.5
	P135L	0	0	0.8	0	0	1.5	0.5	2.5
S	D614G	0	0	1.5	1.8	37.0	64.0	75.0	88.0
Orf3a	Q75H	0	0	0	0	6.0	22.0	23.0	34.0
	G196V	0	0	0	0	0.8	4.0	0.9	0.5
	G251V	0	0	8.0	24.0	8.0	9.0	10.0	3.0
Orf8	V62L	0	5.0	1.0	3.3	0.0	1.5	1.3	3.0
	L84S	0	42.0	37.0	21.0	21.0	18.0	7.0	6.0
Ν	P13L	0	0	0	0	1.0	1.0	2.5	0.5
	S197L	0	0	0	0	1.1	5.0	0.9	0.5
	S202N	0	0	3.5	4.2	0	0.5	2.2	2.5
	R203K	0	0	0	0	17.0	19.0	24.0	23.0
	G204R	0	0	0	0	17.0	19.0	24.0	23.0
	I292T	0	0	0	0	2.0	0.2	0.2	0.5

Table 2. Amino acids selected by region and fortnight. The number indicates the amino acids location in its protein.

Only regions where amino acid change was selected and remained until the last analyzed fortnight are shown. FN: Fortnight; aa: amino acid

Table 3. Number of variable positions, number of mutations, and number of sequences with mutation by region

Pagion	N° of variable aa N° of aa	N⁰ of sequences with			
Region	positions (%)	substitutions	aa substitutions (%)		
nsp1 (180aa)	3 (1.7)	37	37 (2.4)		

nsp3 (1945aa)	158 (8.1)	322	294 (19.3)
nsp14 (527aa)	6 (1.4)	83	83 (5.5)
S (1273aa)	76 (5.9)	1013	904 (59.4)
Orf3a (275aa)	11 (4)	491	468 (30.7)
E (75aa)	5 (6.7)	6	6 (0.4)
Orf6 (60aa)	7 (11.6)	9	9 (0.6)
Orf8 (121aa)	14 (11.6)	312	288 (18.9)
N (419aa)	53 (12.6)	760	470 (30.9)
Total (4875aa)	333 (6.8)	3033	-

aa: amino acid

FIGURE LEGENDS

Figure 1 Phylogenetic signal for SARS-CoV-2 datasets. Presence of phylogenetic signal was evaluated by likelihood mapping, unresolved quartets (center) and partly resolved quartets (edges) for genomes available on April 17 for the nine analyzed regions: nsp1 (29 sequences), nsp3 (225 sequences), nsp14 (65 sequences), S (183 sequences), Orf3a (74 sequences), E (11 sequences), Orf6 (12 sequences), Orf8 (23 sequences), and N (113 sequences). Presence of strong phylogenetic signal (<40% unresolved quartets) was not reached for any region.



Figure 2 Bayesian trees of 29 sequences of nsp1 (540nt), 225 sequences of nsp3 (5835nt), 65 sequences of nsp14 (1581nt), 183 sequences of S (3822nt), 74 sequences of Orf3a (828nt), 11 sequences of E (228nt), 12 sequences of Orf6 (186nt), 23 sequences of Orf8 (366nt), and113 sequences of N (1260nt). Scale bar represents substitutions per site.



Figure 3 Comparison of the evolutionary rates estimated using BEAST for the original dataset and the date-randomized datasets (312 sequences). This analysis was performed for regions nsp3 (5835nt) and S (3822nt). s.s.y = substitutions/site/year.

