1	Full Title:
2	<b>Molecular Evolution of SARS-CoV-2 structural genes:</b>
3	Evidence of positive selection in Spike Glycoprotein
4	
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# 19 Abstract

20	SARS-CoV-2 caused a global pandemic in early 2020 and has resulted in more than 8,000,000
21	infections as well as 430,000 deaths in the world so far. Four structural proteins, envelope (E),
22	membrane (M), nucleocapsid (N) and spike (S) glycoprotein, play a key role in controlling the entry
23	into human cells and virion assembly of SARS-CoV-2. However, how these genes evolve during
24	its human to human transmission is largely unknown. In this study, we screened and analyzed
25	roughly 3090 SARS-CoV-2 isolates from GenBank database. The distribution of the four gene
26	alleles is determined:16 for E, 40 for M, 131 for N and 173 for S genes. Phylogenetic analysis shows
27	that global SARS-CoV-2 isolates can be clustered into three to four major clades based on the
28	protein sequences of these genes. Intragenic recombination event isn't detected among different
29	alleles. However, purifying selection has conducted on the evolution of these genes. By analyzing
30	full genomic sequences of these alleles using codon-substitution models (M8, M3 and M2a) and
31	likelihood ratio tests (LRTs) of codeML package, it reveals that codon 614 of S glycoprotein has
32	subjected to strong positive selection pressure and a persistent D614G mutation is identified. The
33	definitive positive selection of D614G mutation is further confirmed by internal fixed effects
34	likelihood (IFEL) and Evolutionary Fingerprinting methods implemented in Hyphy package. In
35	addition, another potential positive selection site at codon 5 in the signal sequence of the S protein
36	is also identified. The allele containing D614G mutation has undergone significant expansion during
37	SARS-CoV-2 global pandemic, implying a better adaptability of isolates with the mutation.
38	However, L5F allele expansion is relatively restricted. The D614G mutation is located at the
39	subdomain 2 (SD2) of C-terminal portion (CTP) of the S1 subunit. Protein structural modeling
40	shows that the D614G mutation may cause the disruption of salt bridge among S protein monomers

41	increase their flexibility, and in turn promote receptor binding domain (RBD) opening, virus
42	attachment and entry into host cells. Located at the signal sequence of S protein as it is, L5F mutation
43	may facilitate the protein folding, assembly, and secretion of the virus. This is the first evidence of
44	positive Darwinian selection in the <i>spike</i> gene of SARS-CoV-2, which contributes to a better
45	understanding of the adaptive mechanism of this virus and help to provide insights for developing
46	novel therapeutic approaches as well as effective vaccines by targeting on mutation sites.
47	

# 48 Introduction

49 Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), the causative agent of an 50 emerging coronavirus disease (COVID-19) that has caused more than 430,000 deaths, is still a 51 serious global pandemic currently. The genome of SARS-CoV-2 is consisting of a single-stranded 52 and positive-sense RNA of around 30 kb in length with a 5' cap and 3'-polyA tail. It shows that 53 SARS-CoV-2 genome possesses six major open reading frames (ORFs) that encodes 27 different 54 proteins, in which four are structural proteins named Envelope (E), Membrane (M), Nucleocapsid 55 (N) and Spike (S). Many studies have demonstrated important functions of these proteins in virus 56 entry, transcription and virion particle assembly of SARS-CoV-2. The E protein is a small envelope 57 protein with 75 amino acids. Given that a close genetic relationship between SARS-CoV-2 and 58 SARS-CoV, functions of this protein may include virion assembly and morphogenesis[1]. In 59 addition, induction of apoptosis of host cells might be another crucial function of SARS-CoV-2 E 60 protein, thus making it a potential determinant of viral pathogenesis [2]. M protein, consisting of 61 222 amino acids, is the most abundant component of the viral envelope and plays a key role in the 62 virion assembly[3]. N protein, composed of 419 amino acids, may form complexes with genomic 63 RNA, interact with the viral membrane protein, and play a critical role in enhancing the efficiency 64 of virus transcription and assembly[4]. S protein, consisting of 1,273 amino acids, is the most 65 important factor that mediates virus entry and a primary determinant of cell tropism and 66 pathogenesis of SARS-CoV-2[5]. 67 Many studies demonstrated SARS-CoV-2 underwent the evolution and some genetic evolutionary

Many studies demonstrated SARS-CoV-2 underwent the evolution and some genetic evolutionary
features have been reported[6]. The whole genomic sequence of SARS-CoV-2 has 79.6% identity
with SARS-CoV and 96% with a bat SARS-related coronavirus (SARSr-CoV), RaTG13. Although

70	no positive time evolution signal was found between SARS-CoV-2 and RaTG13, the SARS-CoV-
71	2 shows a strong positive temporal evolution relationship with bat-SL-CoVZC45, which has a
72	slightly less identical genomic sequence (87.5%) than RaTG13 [7]. Combining the phylogenetic
73	analysis of full-length genomes of coronaviruses, a potential bat origin of SARS-CoV2 is indicated
74	[8]. A recent study reported that <i>spike</i> (S) gene (coding gene of S protein) of SARSr-CoVs from
75	their natural reservoir host, the Chinese horseshoe bat ( <i>Rhinolophus sinicus</i> ), has coevolved with <i>R</i> .
76	sinicus angiotensin converting enzyme 2 (ACE2) via positive selection[9]. A single-stranded
77	positive-sense RNA virus as it is, SARS-CoV-2 causes global pandemic within half a year,
78	suggesting it may evolve rapidly. However, the evolution of SARS-CoV-2 based on structural genes
79	from human to human transmission has not been investigated in detail. The primary purpose of this
80	work is to study the evolutionary pattern of the four structural genes of SARS-CoV-2 derived from
81	a global isolate collection including the E, M, N and S. Various molecular evolution and selection
82	analysis approaches were employed to identify the phylogeny of the four structural proteins and
83	potential selection effects on these genes. Hereby, our study reveals that intragenic recombination
84	does not contribute to the evolution of these genes while purifying selection is the main evolutionary
85	force. Moreover, a D614G mutation in the S protein is operated by strong positive selection and
86	may be responsible for the quick spread of SARS-CoV-2 globally. Additionally, another potential
87	L5F mutation may also be operated by positive selection, but with relatively less strong pressure as
88	compared to D614G.

89

# 90 Materials and Methods

91 SARS-CoV-2 isolates

92	Complete full-length genomic sequences of SARS-CoV-2 were downloaded from 2019 Novel
93	Coronavirus Resource (2019nCoVR) in China National Center for Bioinformation. All of which
94	were also uploaded to the NCBI GenBank database. The sequences were manually checked and
95	finally a total of 3090 isolates were selected and verified for the present study. These isolates were
96	collected from December 24, 2019 to April 24, 2020 in the different geographical locations
97	including China, USA, Japan, Pakistan, Australia, Greece, German, Peru, Turkey, Kazakhstan, Iran,
98	Serbia, Thailand, Nederland, Sri Lanka, Czech, Malaysia, India etc. Detailed information of these
99	isolates including the GenBank accession number or biosample number is summarized in S1 Table.
100	

## 101 Sequence analysis of the four structural genes and proteins

102 The E, M, N, S gene sequences were extracted from SARS-CoV-2 global isolate collection and

aligned by the MEGA X package using Muscle (codons) parameters [10]. Because some regions of

104 genomic sequences of SARS-CoV-2 couldn't be exactly identified, in which nucleic acid bases are

shown as degenerate bases (e.g. N, R, Y), we were unable to obtain all of the four structural gene

sequences from an isolate sometimes. Allele type and DNA sequence polymorphism analyses were

107 performed using DnaSP 6.12.03[11]. The protein sequences and polymorphism loci of these isolates

108 were also aligned and analyzed with the MEGA X.

109

#### 110 Molecular evolution analysis

111 An unrooted phylogenetic tree of the four structural proteins was constructed using the MEGA X

112 package [10], and the evolutionary history was inferred using the Maximum Likelihood method,

113 based on the JTT matrix-based model for E protein sequences, General Reversible Chloroplast +

114 Freq. model for M, JTT matrix-based model for N and Jones et al. w/freq. model for S protein 115 sequences. Model selection was conducted in MEGA X. Bootstrap values were estimated by 1000 116 replications. Initial tree(s) for the heuristic search were obtained automatically by applying 117 Neighbor-Join and BioNJ algorithms to a matrix of pairwise distances estimated using each model 118 mentioned above. The tree is drawn to scale, and FigTree V1.4 was utilized to form cladogram 119 branches (http://tree.bio.ed.ac.uk/software/figtree/). The aligned DNA sequences were also 120 screened using RDP4 software to detect intragenic recombination among the alleles of each 121 structural gene[12]. Six methods implemented in the RDP4 were utilized. These methods are RDP 122 [12], GENECONV[13], BootScan [14], MaxChi[15], Chimaera [16], and SiScan [17]. Common 123 settings for all methods include considering sequences as linear and setting statistical significance 124 at the P < 0.05 with Bonferroni correction for multiple comparisons and requiring phylogenetic 125 evidence and polishing of breakpoints. Potential recombination events (PREs) were considered as 126 those identified by at least two methods. Reticulate network tree of alleles of the four structural 127 genes of SARS-CoV-2 was also generated by Splitstree4 [18]. Phi test implemented in Splitstree4 128 was used to define probable recombination events. Tajima's D, Fu and Li's D\* and F\* tests were 129 employed to test the mutation neutrality hypothesis of the whole gene as previously described by 130 our research group[19]. These analyses were carried out using DnaSP 6.12.03[11]. A statistical 131 significance level with P < 0.05 is acceptable. The false discovery rate and 1000 replications in a 132 coalescent simulation were applied for correcting multiple comparisons. Non-neutrality evolution 133 was considered when identified by at least two out of three tests. Nonsynonymous and synonymous 134 mutations of the alleles of the four structural genes were also calculated using MEGA X package 135 [10].

136

## 137 Analysis of positive selection based on codon

138 The selection pressure operating the four structural genes of SARS-CoV-2 was searched by using 139 the Maximum Likelihood (ML) method. Analyses were performed using a visual tool of codeml 140 program, named EasyCodeML algorithm with site model [20]. Three nested models (M3 vs. M0, 141 M2a vs. M1a, and M8 vs. M7) were compared and likelihood ratio tests (LRTs) were applied to 142 access a better fit of codes. Model fitting was also performed using multiple seed values for dN/dS143 and assuming the F3x4 model of codon frequencies. Positive selection is inferred when individual 144 site or codon with ratio of nonsynonymous to synonymous mutations (dN/dS ratios) is greater than 145 one ( $\omega$ >1). When the LRT is significant (p < 0.05), Bayes empirical Bayes (BEB) (M8 model) and 146 Naive Empirical Bayes (NEB) methods (M3 and M2a model) are further employed to identify 147 amino acid residues that likely evolve under positive selection based on a posterior probability 148 threshold of 0.95. Results from M8 model were taken as the standard as Yang et al. reported. M3 model was used for the frequency distribution of codon class analysis as Yang et al. 149 150 recommended[21]. HyPhy package was used to validate the result obtained by ML method[22].

151

## 152 Structural modeling of the protein with positive selection sites

Three-dimensional structures of proteins with positive selection sites were modeled using SWISSMODEL (<u>http://swissmodel.expasy.org</u>) according to the most fitted protein template. Model
quality was evaluated by QMEAN while the structure of the model was visualized by using PyMoL
[23].

157

## 158 **Results and Discussion**

#### 159 Characteristics of SARS-CoV-2 isolates, structural gene and protein

#### 160 sequences

161 The 3090 SARS-CoV-2 isolates harbor only 16 unique alleles of E and 40 alleles of M, but an 162 abundant number of alleles of N and S genes, which contain 131 and 173, respectively. These alleles 163 correspond to 10, 14, 88 and 99 different amino acid sequences of E, M, N, and S proteins, 164 respectively. Protein sequence comparisons of WH01 isolate with SARSr-CoV, bat-SL-CoVZC45 165 isolate show 100% (75/75) identity in E, 98.65% (219/222) identity in M, 94.27% (395/419) identity 166 in N and 80.06% (1171/1273) in S proteins, respectively. These results imply a close kinship 167 between SARS-CoV-2 and bat SARSr-CoV, especially on E and M proteins. On the other hand, it 168 indicates an extreme conservation of E and M proteins and their functions among coronaviruses[24]. 169 Further analysis revealed that there are 14 single nucleotide polymorphisms (SNPs) of E gene, but 170 only 5 single amino acid polymorphic (SAP) loci in the E protein. Similar result was observed on 171 M gene and protein, with 37 SNPs and 9 SAPs. In contrast, 126 SNPs and 75 SAPs are detected on 172 N gene and protein, respectively. S protein, the most important factor that mediates virus entry by 173 receptor binding and membrane fusion and determines the infection ability of SARS-CoV-2 [25], 174 harbors 155 SNPs on the alleles and 90 SAPs in the protein. Considering the size of nucleotides and 175 amino acid residues, N gene has the maximum sequence variability with 10.02% (126/1257) SNPs 176 and 17.90% (75/419) SAPs, respectively. However, S gene has most pairwise nucleotide differences 177 among the four structural genes, indicating a more genetic diversity of S gene (Table 1). A key 178 player in the virus transcription and assembly as N protein is [26, 27], high sequence variability of 179 the N protein may indicate a vast adaption of the virus during host transmission. Previous study

- 180 shows that high genetic variance has been found among bat SARSr-CoVs, particularly in the S
- 181 gene[9]. Similar, higher nucleotide diversity ( $\pi$ , a major parameter to define genetic diversity) of S
- 182 gene is also detected on SARS-CoV-2 isolates, suggesting this may benefit virus survival in the host
- 183 of human beings.
- 184

185 Table 1. Summary of genetic diversity of the 4 structural genes of the SARS-CoV-2 isolates

Gene	Sequence, n*	Sequence	h	π	S	θ	η
		length					
Е	2928	228	16	0.00012	14	0.00475	15
М	2891	669	40	0.00018	37	0.00665	40
Ν	2253	1260	131	0.00056	126	0.01081	130
S	2339	3825	173	0.00075	155	0.00753	169

186

- 187 *h*, Haplotypes,
- 188  $\pi$ , Nucleotide diversity
- 189 *S*, Polymorphic sites
- 190  $\theta$ , Theta (per site) from S, population mutation ration
- 191 *П*, Total number of mutations

192 \* Some bases of SARS-CoV-2 genomic sequences are not exactly identified; thus, the number of

193 gene sequences were less than 3090.

194

### 195 Distinct phylogenetic patterns of the four structural genes

The phylogenetic analysis revealed that all SARS-CoV-2 E proteins form three clusters. Similar to E protein, phylogenetic tree of SARS-CoV-2 M proteins is formed by three clusters with few branches (Figs 1A and 1B). The results suggest both E and M genes may display a relatively high conservation during coronavirus evolution. In contrast, SARS-CoV-2 N and S proteins show distinct phylogenetic pattern as compared with that of E and M. Four and three main phylogenetic clusters with various branches are identified in the N and S proteins, respectively (Figs 1C and D). Given the crucial roles of N and S proteins in virus transcription, assembly, and entry to host cells, whether

203 SARS-CoV-2 isolates harbor different N and S variants (such as those clustered into different clades)

204 may influence their infection efficiency remains unknown, and requires further study.

# 205 Purifying selection drives the evolution at whole structural gene levels

## 206 of SARS-CoV-2 during its human to human transmission

207 Although many studies demonstrated that recombination plays an important role on the emergence 208 of SARS-CoV-2 and its contribution to admit SARS-CoV-2 as a human infectious pathogen [28-209 30], how this virus evolves during its global transmission has not been profiled yet. Therefore, we 210 first analyzed intragenic recombination events of each structural gene using RDP4. The results 211 indicate there were no recombination events occurred among the alleles of each gene (data not 212 shown). Recombination event were also assessed through reticulate network tree by phi test in 213 SplitsTree4. Although some internal nodes are noticed in N and S alleles, no significant evidence 214 for recombination is validated of each gene by Phi test (p>0.05) (Fig 2). It indicates a relative stable 215 state of SARS-CoV-2 during its transmission although a possible genetic interaction of different 216 isolates might have occurred when it became a global pandemic [31, 32]. In addition, Tajima's D, 217 Fu and Li's D\* and F\* statistics were calculated to examine the mutation neutrality hypothesis of 218 the four structural genes of SARS-CoV-2. The results reveal that the evolution of all four genes 219 does not match the neutral hypothesis, but favor purifying selection (Table 2 and Fig 3). The average 220 of all pairwise dN/dS ratios ( $\omega$ ) among the alleles of each structural gene of SARS-CoV-2 is 0.5443 in E, 0.1562 in M, 0.07978 in N, and 0.4980 in S gene, respectively. All together, these results 221 222 suggest that at the whole gene level, inconsistent purifying selection is the main evolution force 223 (Table 2). Li et al studied the origin of SARS-CoV-2 and showed evidence of strong purifying 224 selection in the S and other genes among bat, pangolin and human coronaviruses, indicating similar

225	strong evolutionary constraints in different host species [33]. Similarly, our results suggest purifying
226	selection drives the evolution at the whole structural gene level of SARS-CoV-2 during its
227	transmission from human to human. This result also implies that in general, the genetic variation on
228	these structural genes will not confer a significant disadvantage on the virus survival, and ratios
229	reflect general variability of these genes and proteins. Considering that no recombination happened,
230	nonsynonymous mutations would be removed at a great rate during the virus transmission [34].
231	

231

Table 2. Summary of neutrality for the four structural genes in SARS-CoV-2 isolates 232

Gene	Tajima'sD	Fu and Li's D* tes	Fu and Li's F	dN	dS	dN/dS	Selection
		t	* test			(ω)	
Е	-2.29974, P<0.01	-3.18477, P<0.02	-3.38505, P<0.02	0.006836	0.1256	0.5443	Purifying
							selection
Μ	-2.74611, P<0.001	-5.64276, P<0.02	-5.50855, P<0.02	0.001294	0.008296	0.1562	Purifying
							selection
Ν	-2.87598, P<0.001	-9.67153, P<0.02	-7.95879, P<0.02	0.000251	0.003146	0.07978	Purifying
							selection
S	-2.87646, P<0.001	-11.01171, P<0.02	-8.59037, P<0.02	0.000609	0.001223	0.4980	Purifying
							selection

233

#### SARS-CoV2 S gene is operated by positive selection at a definitive 234 codon located at the C-terminal portion of S1 subunit and a potential 235 codon located at the signal sequence 236

237 Guo et al. reported that the S gene of SARSr-CoV populations in their natural host, Chinese 238 horseshoe bat (*Rhinolophus sinicus*), has evolved through positive selection at some codons[9]. As 239 mentioned above, at the whole gene level, purifying selection is the main force driving the evolution 240 of studied genes. Whether positive selection pressure accelerates the diversification of the structural 241 genes of SARS-CoV-2 remains unclear. Therefore, we used codon-substitution models to estimate 242 the ratio of nonsynonymous over synonymous substitutions (dN/dS), also known as  $\omega$ . The role of 243 recombination in the polymorphism of four genes is excluded because no intragenic recombination 244 was detected (Fig 2). By using ML model, we don't find any codon of E and M gene subjecting to 245 positive selection obviously (data not shown). However, a potential positive selection site 208A in 246 N gene is identified by using M3 model, but not by any other models especially the M8 model, 247 suggesting a limited amount of evidence of positive selection in N gene (S1 Table). For the S gene, 248 we found the average  $\omega$  is 0.37199 calculated by M0 model of the codeML package, suggesting that 249 purifying selection was a major force operating the evolution of the S gene during its transmission 250 among human beings. In three LRTs, all alternative models (M3, M2a, M8) are significantly better 251 fit (P<10<sup>-4</sup>) than relevant null models (M0, M1a, M7), indicating that some sites of S were subjected 252 to strong positive selection ( $\omega$ =18.22175-20.61283) (Table 3). A single positive selection site (614D) 253 is identified in the S gene with posterior probability of 1.000 in all the three models [21], a clear 254 evidence showing that this site is still experiencing positive selection when the virus transmitted 255 from human to human. The result is also validated using internal fixed effects likelihood (IFEL) and 256 Evolutionary Fingerprinting methods implemented in HyPhy package (Fig 4) [35-37]. To our 257 surprise, the positive selection site is not located at the receptor binding domain (RBD) or receptor 258 binding motif (RBM) as we anticipated, which play the most important role in virus-receptor 259 interaction and virus entry into host cells [38]. This result suggests that a relatively genetic stability 260 of this motif would benefit the virus survival. Intriguingly, the site under positive selection pressure 261 always has a D614G (for the S gene is 1841A>G) mutation, implying such mutation may enhance 262 virus adaptability in human hosts. Another potential positive selection site at codon 5 is also 263 identified, and a L5F mutation (for the S gene is 13C>T) is always found, with posterior 264 probabilities greater than 0.95, 0.93 and 0.92 (critical values) calculated by M3, M2a and M8 models

- 265 (Table 3), respectively. Similar result was also confirmed by Evolutionary Fingerprinting method
- 266 (S1 Fig). Considering signal sequence (SS) is a short hydrophobic peptide that plays an important
- role in guiding viral protein into the endoplasmic reticulum (ER) for proper folding and assembly
- 268 [39], we postulate that L5F mutation may increase hydrophobicity of the SS, thus facilitating the
- entry of S protein into ER for folding and assembly, and in turn secretion of the virus.

270

Model	Ln L	Estimates of parameters	Model compared	LRT P-value	Positive sites
		p0=0.96797, p1=0.02883, p2=0.00320			<b>5 L 0.958</b> *,28 Y 0.850,221 S 0.901, <b>614 D</b>
M3 (discrete)	-6766.339162	ω0=00.26126, ω1= 2.70530, <b>ω2=20.61283</b>			<b>1.000</b> ***,677 Q 0.891
M0 (one ratio)	-6790.072925	ω0=0.37199	M0 vs. M3	0.000000001	Not Allowed
		p0=0.81731, p1=0.17872, p2=0.00397			5 L 0.9258,28 Y 0.812,221 S 0.832, <b>614 D</b>
M2a(selection)	-6766.432802	ω0=0.17504, ω1=1.00000, <b>ω2=18.76936</b>			<b>1.000</b> ***,677 Q 0.828
		p0=0.70461, p1=0.29539			
M1a (neutral)	-6778.770190	ω0=0.04395, ω1=1.00000	M1a vs. M2a	0.000004385	Not Allowed
		p0=0.99578, p=0.40368, q=0.82224			5 L 0.931,28 Y 0.817,221 S 0.831, <b>614 D</b>
M8(beta&ω)	-6768.829411	p1=0.00422, <b>ω= 18.22175</b>			<b>1.000</b> ***,677 Q 0.828
M7(beta)	-6779.230494	p=0.00857, q=0.02623	M7 vs.M8	0.000030400	Not Allowed

#### 271 Table 3. Log-likelihood values and parameter estimates for the SARS-CoV-2 S gene sequences

272

273 LnL is the log likelihood;  $\omega$  is ratio of dN/dS, LRT P-value indicates the value of chi-square test; Parameters indicating positive selection are presented in bold;

274 Positive selection sites were identified by the Bayes empirical Bayes (BEB) methods under M8 model. The posterior probabilities  $(p) \ge 0.80$  are shown,  $(p) \ge 0.95$ 

275 (p) ≥0.99, and (p)=1.000 are indicated by \*, \*\* and \*\*\*, respectively. Yang *et al.* recommended that results from M8 model were preferred to find sites under positive

selection pressure.

#### 277 Evolutionary relationship of S gene alleles with or without D614G and

#### 278 L5F mutation

279 Phylogenetic tree of S gene alleles was derived to test the evolutionary relationship among the alleles 280 with or without D614G mutation. As shown in Fig 5A, the 173 alleles of the S gene could be 281 clustered into four clades. Alleles with D614G mutation could be found in all 4 clades, among which 282 a dominant one contains 79 out of 85 alleles with such mutation. The remaining 6 mutated S alleles 283 are distributed in other 3 clades. The result suggests a potential common ancestor for the majority 284 of S alleles with D614G mutation, while some other maybe derived from alternative ancestors. This 285 result is also supported by the parsimony network of S gene alleles using PopART 286 (http://popart.otago.ac.nz) [40]. Two central alleles (representative virus isolates are WH01 and 287 GZMU0019) and associated alleles around them form a star scattering network, suggesting that the 288 S gene may have two potential origins (Fig 5B). All S alleles with D614G mutation are closely 289 related (with a few point mutations), and comprise a scattered star structure, suggesting the 290 expansion of SARS-CoV-2 population with D614G mutation on S gene. In contrast, alleles of the 291 N gene show a single ancestor analyzed by parsimony network though 3 phylogenetic clades are 292 identified (S2 Fig).

A total of 5 alleles with L5F mutation are found and all of them are in one clade, accounting for 83.33% of all alleles in the clade (S3A Fig). Further parsimony network analysis reveals that S alleles with L5F mutation are not closely related, but distribute in both WH01 and GZMU0019 haplotype groups (S3B Fig). No scattered star structure of these alleles can be formed, indicating L5F mutation might arise from independent origins other than that of D614G mutants. Limited

number of alleles with L5F mutation identified so far also suggests that L5F might subject torelatively less strength of the pressure and is still at early stage of positive selection.

300

### 301 Frequency of S allele with D614G mutation increased in SARS-CoV-2

#### 302 isolates during human to human transmission

303 Considering that mutation of a positive selection site should be beneficial to the survival of the 304 individuals carrying the mutation, we postulate that the D614G (1841A>G) mutation may help the 305 spread of SARS-CoV-2. Some evidence has been obtained from the haplotype network of S alleles 306 mentioned above (Fig 5B). S gene haplotypes (alleles) with D614G mutation (representative isolate 307 GZMU0019) have evolved many subtypes and comprise a star structure with GZMU0019 in the 308 center. This starburst pattern with one haplotype in the center and many other haplotypes 309 surrounding the central haplotype suggests a signature of rapid population expansion [41]. To 310 further study whether SARS-CoV-2 isolates with D614G mutation have advantage in survival 311 during its transmission among human beings, we calculated the frequencies of S alleles carrying 312 D614G mutation in each week from the collected SARS-CoV-2 isolates from December 24, 2019 313 to April 20, 2020 (17 weeks). Detailed information of these isolates including collection date, 314 collection region and accession or biosample numbers is summarized on S3 and S4 Tables. 315 In 173 S gene alleles, 85 carry D614G mutation, accounting for 49.13% of all. Similarly, 47 out 99 316 S proteins carry D614G mutation, accounting for 47.47% of all. The first two isolates, 317 GWHABKF00000001 and WH01 (isolated in December 24, 2019 and December 26, 2019, 318 respectively), carry 614D in the S protein, while the first SARS-CoV-2 isolate with a D614G 319 mutation is GZMU0019 in our collected dataset, isolated from a patient with COVID-19 in

320 Guangzhou, Guangdong Province of China on February 5, 2020 (week 7 in our dataset). After that, 321 except for week 9 and week 10 (possibly due to the small number of samples and sampling 322 deviation), a spread trend that more and more proportion of isolates carry the D614G mutation in 323 the S protein stands out. In the week 17, the last week of our dataset, 91.11% of SARS-CoV-2 324 isolates carry this mutation (S3 Table, Fig 6A). Further analysis reveals that the frequency of D614G 325 mutation in the S gene was steadily increasing when combining data from week 6 to 17 (S3 Table, 326 Fig 6B). To exclude the influence of sample size on the result (in some weeks, only 4-6 isolates 327 were collected in the dataset), we reorganized the dataset by taking both the sample size and 328 sampling time into account. Various panels of 200-300 isolates were studied and similar results 329 were observed (S4 Table, Figs 6C and D). Taken together, these results suggest that SARS-CoV-2 330 isolates with D614G mutation may increase their ability to transmit, and contribute to the rapid 331 spread of this virus to the world.

332

# 333 D614G mutation of S gene may destabilize S protein trimer and 334 promote receptor binding and membrane fusion

The positive selected D614G mutation might play an important role for the adaptability of SARS-CoV-2 in both the host and the virus population[42]. Another explanation is that the mutation is driven by specific interaction between high level of virus sequence divergence and polymorphic host receptors or interacting proteins[43]. S protein is the key determinant for the tissue tropism and host range and specificity of coronavirus such as SARS-CoV-2. The virus infects host cells through the interaction between the S protein and its cellular receptor, named ACE2 [8]. In this process, virus entry requires the precursor S protein cleaved by cellular proteases including trypsin, furin,

342	transmembrane serine protease 2 (TMPRSS2), or endosomal cathepsin L, which generate the
343	receptor binding subunit S1 and the membrane fusion S2 [44-46]. From structural studies in both
344	SARS-CoV and SARS-CoV-2, receptor binding domain (RBD) located at the C-terminal of S1 and
345	the adjacent N-terminal domain (NTD) are relatively flexible, which is the feature required for
346	receptor recognition and subsequent membrane fusion[47, 48]. We found that the D614G mutation
347	is located at the subdomain 2 (SD2) that at the C-terminal of RBD and close to the two potential
348	cleavage sites between S1 and S2 [48] (Fig 7A). Considering that positive selection is usually
349	beneficial to the survival of the individual carrying the mutation, we speculate that the D614G
350	mutation may facilitate structural conformation change to promote receptor binding or membrane
351	fusion[5, 44], and in turn improving the infection efficiency. From the latest cryo-electron
352	microscopy (cryo-EM) structure of SARS-CoV-2 S protein, the negatively charged sidechain of
353	D614 points towards the positively charged sidechain of K854 from the neighboring monomer (Fig
354	7B) [48]. The distance between the closest atoms of the two residues is 2.6 Å, which is an optimal
355	distance to form salt bridge (Fig 7C). From the modelled structure with D614G mutation, the
356	distance is increased to 5.2 Å (Fig 7D), which would potentially abolish the salt bridge and
357	destabilize the integrity of the S trimer in wild type. It has been reported that human receptor ACE2
358	binds to an "open" conformation of S protein, where RBD move away from the core structure and
359	expose its receptor binding surface. The entire S trimer then undergoes a serial of dramatic
360	conformation changes, including cleavages between S1 and S2, disassociation of S1 and post-fusion
361	transformation of S2 [49, 50]. Changes including mutations at cleavage sites and adding internal
362	crosslinks in S trimer would keep the protein in a stable and "closed" conformation where the
363	receptor binding surface of RBD is inaccessible [48, 51]. Therefore, we hypothesize that the

364	highly transmissible D614G mutation driven by the positive selection through evolution promotes
365	accessibility of RBD by losing a critical salt bridge between the S protein monomers, which
366	subsequently triggers membrane fusion upon ACE2 binding.

367

## 368 Conclusions

369 We present modern molecular evolution analyses on a large and comparative set of SARS-CoV-2 370 structural gene sequences, derived from an international collection of SARS-CoV-2 isolates. 371 Distinct phylogenetic patterns of four structural proteins of SARS-CoV-2 are depicted. Protein 372 sequence comparisons show E and M genes exhibit a relatively close relationship to bat SARSr-373 CoV, suggesting the evolution conservation of these two genes. In contrast, relatively high genetic 374 variation is observed in N and S proteins among SARS-CoV-2 isolates, implying extensive 375 adaptability of N and S genes. No clear intragenic recombination is detected of these four genes, 376 suggesting that it is not the major force to drive the evolution of the four genes. However, our 377 analyses show purifying selection pressure may be the main force operating the evolution at whole 378 gene levels of SARS-CoV-2 during its human to human transmission. We also identify a codon in 379 S gene definitively experiencing positive selection pressure, and always leads to the D614G 380 mutation in S proteins. S alleles with D614G mutation have expanded rapidly among SARS-CoV-381 2 isolates. D614G mutation significantly extends the distance between monomers in the S protein trimer, which may disrupt the salt bridge formed by D614 and K854 between monomers, promote 382 383 RBD opening, and facilitate the entry of the virus into host cells, thus contributing to the diffusion 384 of this mutated alleles. Codon 5 of S gene is another potential positive selection site. Although a 385 limited number of alleles with L5F mutation is identified, it may potentially affect the assembly and

392	Acknowledgements
391	
390	and how they affect the expansion of these mutated alleles on SARS-CoV-2.
389	mechanism remains unclear, further study should focus on the exact function of these mutation sites
388	diagnostics, the D614G mutation of S should be paid more attention. Owning that the exact
387	occurs. As S protein is a key target for SARS-CoV-2 vaccines, therapeutic antibodies, and
386	secretion of SARS-CoV-2. A close eye on L5F mutation may be required in case another expansion

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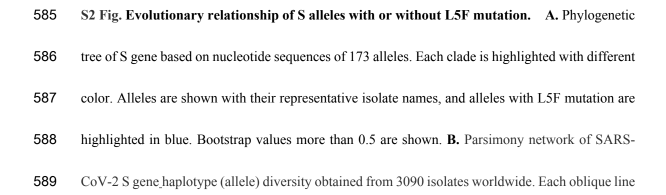
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567	

# 568 Supporting information

- 569 S1 Table. SARS-CoV-2 isolates information.
- 570 S2 Table. Log-likelihood values and parameter estimates for the SARS-CoV-2 N gene sequences.
- 571 S3 Table. Detailed information of SARS-CoV-2 isolates with full length sequence of S gene. The
- 572 data are organized by weekly.
- 573 S4 Table. Detailed information of SARS-CoV-2 isolates with full length sequence of S gene. The
- data are organized by panels. Each panel contains 200-300 isolates by combining isolates from
- 575 several days.

576 S1 Fig. The evolutionary relationship of N alleles. A. Phylogenetic tree of N gene based on 577 nucleotide sequences of 131 alleles. The evolutionary history is inferred using the Maximum 578 Likelihood method and Tamura-Nei model. The tree is drawn to scale, with branch lengths measured 579 in the number of substitutions per site. Bootstrap values more than 0.5 are shown. B. Parsimony 580 network of SARS-CoV-2 N gene haplotype (allele) diversity obtained from 3090 isolates worldwide. Each oblique line linking between haplotypes (haplotype name is shown as its representative isolate 581 582 name) represents one mutational difference. The ancestral haplotype, or root of the network, is 583 labeled with a square, and represent haplotype name is marked red. 584



590	linking between haplotypes (haplotype name is shown as its representative isolate name) represents
591	one mutational difference. Unlabeled nodes (Gray circle) indicate inferred steps have not found in
592	the sampled populations yet. The ancestral haplotype, or root of the network, is labeled with a square,
593	and represent haplotype name is marked green or red. The blue nodes indicate haplotypes with L5F
594	mutation. Dotted boxes indicate major haplotype groups. Haplotypes include in red dotted boxes
595	are with D614G mutation while those included in black dotted boxes are without D614G mutation.
506	
596	
596	S3 Fig. Positive selection analysis of S gene codon 5 by Evolutionary Fingerprinting method.
	<b>S3 Fig. Positive selection analysis of S gene codon 5 by Evolutionary Fingerprinting method.</b> Log (Bayes Factor) for positive selection at codon 5 of S gene and its frequencies. The cut-off value
597	
597 598	Log (Bayes Factor) for positive selection at codon 5 of S gene and its frequencies. The cut-off value
597 598 599	Log (Bayes Factor) for positive selection at codon 5 of S gene and its frequencies. The cut-off value for the Bayes factor (BF) in the Evolutionary Fingerprinting method was set at 25 to reflect a positive

603

# 604 Figure legends

605

- 606 Figure 1. Phylogenetic tree of E (A), M (B), N (C), and S(D) proteins of SARS-CoV-2. Major
- 607 clades are highlighted with different color. The tree shows topology of the protein of each allele,
- 608 named by their representative isolates.
- 609
- 610 Figure 2. Reticulate network trees of E (A), M (B), N (C) and S (D) alleles of SARS-CoV-2
- 611 analyzed by the neighbor-net algorithm of SplitsTree4. Scale bars indicate number of substitutions
- 612 per site. All internal nodes represent hypothetical ancestral alleles and edges that correspond to
- 613 reticulate events such as recombination. Red arrows indicate edges. Because there are too few
- 614 informative characters to use the Phi test for E and M genes, *p-values* of Phi test of N and S genes
- are shown.
- 616

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617 Figure 3. Tajima's D, Fu and Li's D* and F* test for the four structural gene alleles of SARS-
618 CoV-2. *p < 0.05; **p < 0.01; ***p<0.001</li>
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621 Figure 4. Positive selection analysis of S gene codons by IFEL and Evolutionary 622 Fingerprinting methods. A. Diagram of selection analysis result of S codons by IFEL method. 623 Asterisk indicates the positive selection site with statistical significance (p < 0.01). B. Log (Bayes 624 Factor) for positive selection at codon 614 of S gene and its frequencies. The cut-off value for the 625 Bayes factor (BF) in the Evolutionary Fingerprinting method was set at 25 to reflect a positive

626 selection

selection at a given site (Posterior probability>0.95). Pr {BF>25} indicates posterior probability of

**627** Bayes Factor >25.

628

629 Figure 5. Evolutionary relationship of S alleles with or without D614G mutation. A. 630 Phylogenetic tree of S gene based on nucleotide sequences of 173 alleles. The evolutionary history 631 is inferred using the Maximum Likelihood method and Tamura-Nei model. The tree is drawn to 632 scale, with branch lengths measured in the number of substitutions per site. Each clade is highlighted 633 with different color. Alleles are shown with their representative isolate names, and alleles with 634 D614G mutation are highlighted in red. Bootstrap values more than 0.5 are shown. B. Parsimony 635 network of SARS-CoV-2 S gene haplotype (allele) diversity obtained from 3090 isolates worldwide. 636 Each oblique line linking between haplotypes (haplotype name is shown as its representative isolate 637 name) represents one mutational difference. Unlabeled nodes (Gray circle) indicate inferred steps 638 have not found in the sampled populations yet. The ancestral haplotype, or root of the network, is 639 labeled with a square, and represent haplotype name is marked green or red. The red nodes indicate 640 haplotypes with D614G mutation, while green or black nodes indicate haplotypes without D614G 641 mutation. Dotted boxes indicate major haplotype groups.

642

643 Figure 6. Expansion of S alleles with D614G mutation during SARS-CoV-2 human to human
644 transmission. A. Percentage of SARS-CoV-2 isolates carrying the alleles of D614G mutation in
645 each week collected. B. Frequencies of D614G mutation in the S gene in each period of time (Four
646 to five weeks' data are combined). C. Percentage of SARS-CoV-2 isolates carrying the alleles with

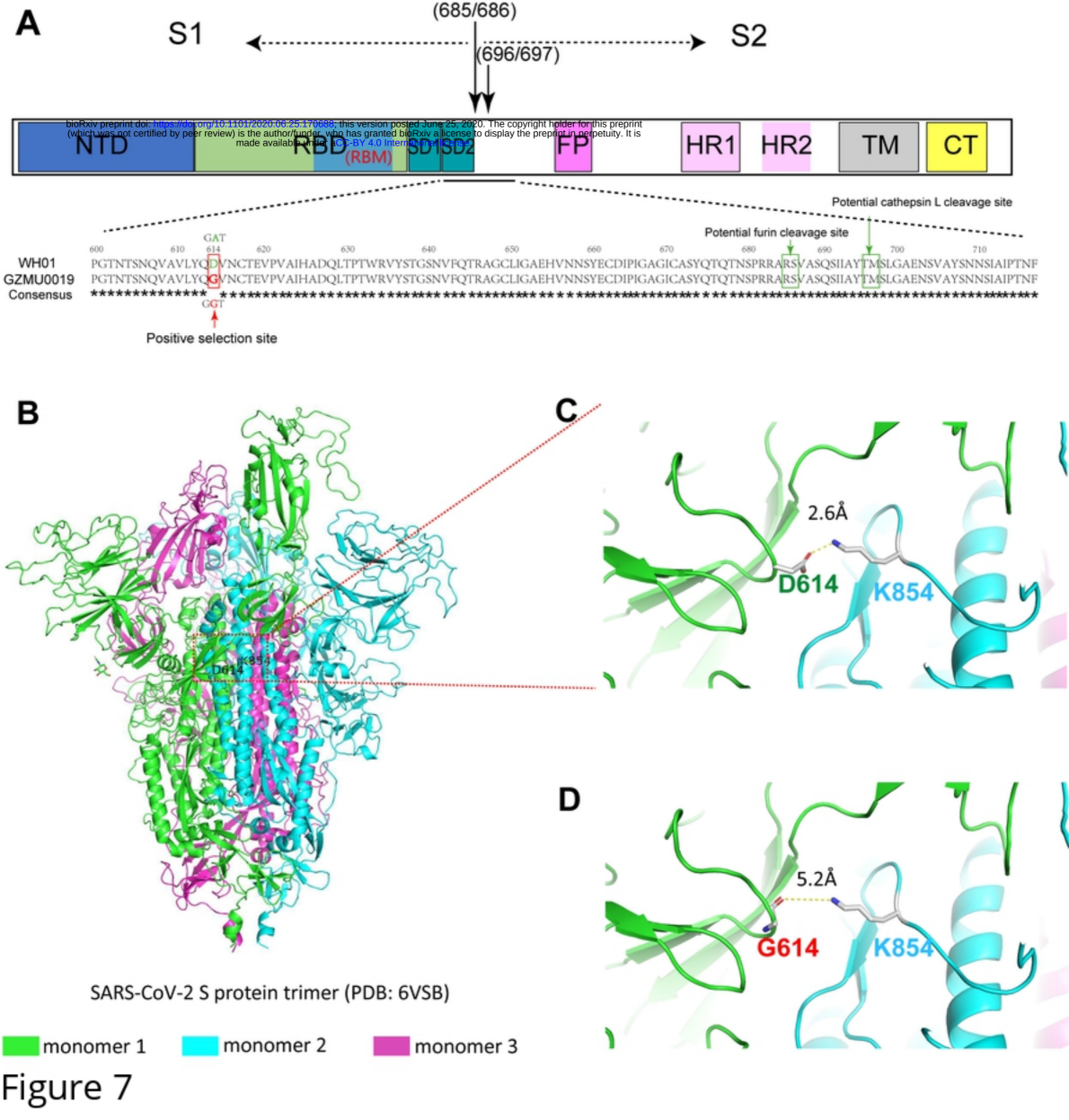
647 D614G mutation in each period of time. **D.** Frequencies of D614G mutation in the S gene in each
648 period of time. \*p < 0.05; \*\*p < 0.01.</li>

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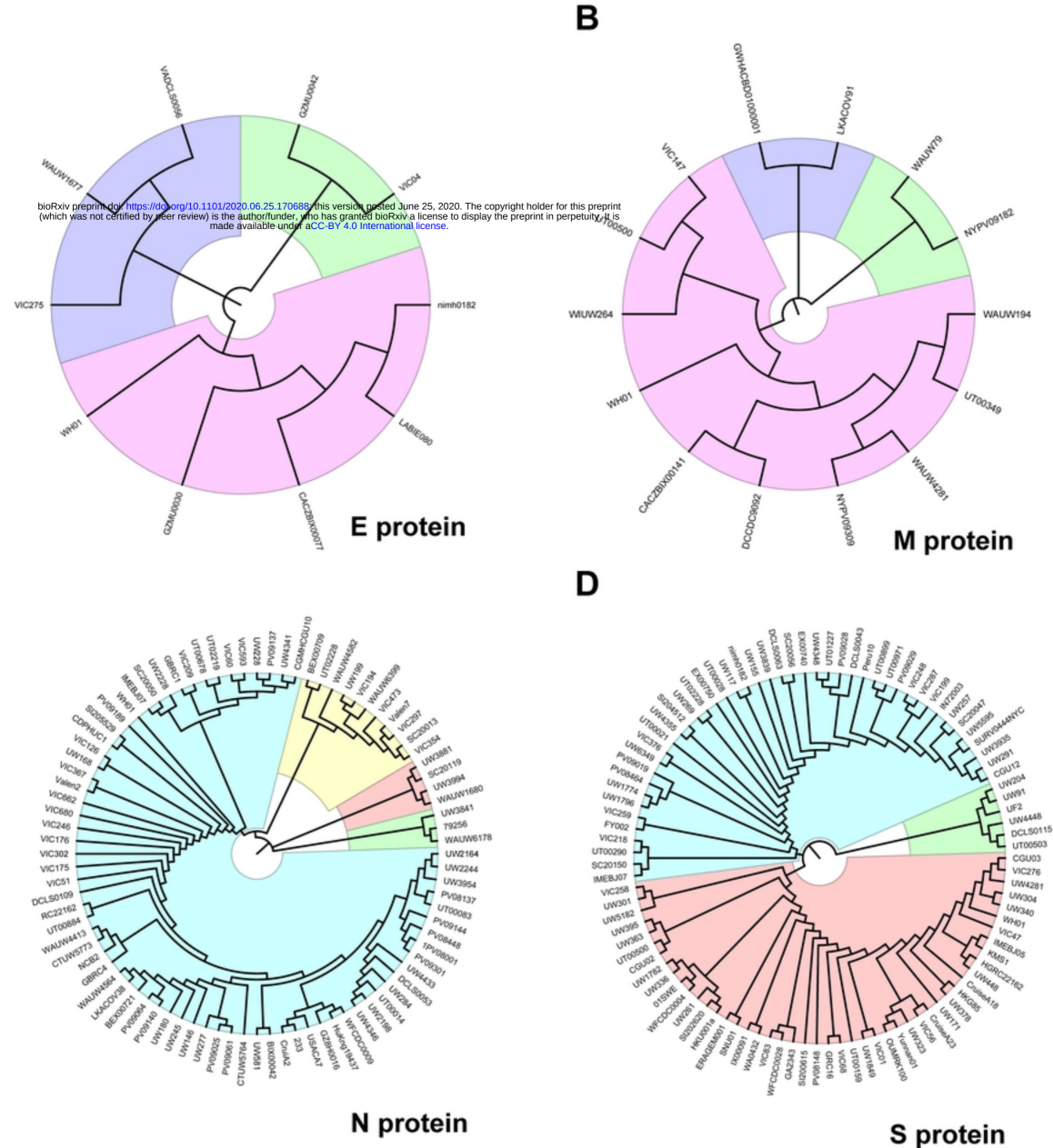
651	Figure 7. The structure of the S protein of SARS-CoV-2 and potential influence of D614G mutation
652	on its structural change. A. Schematic of the primary structure of SARS-CoV-2 S protein colored
653	by domains. Some boundary-residues are listed. The S1/S2 cleavage sites are indicated by arrows.
654	RBD: receptor binding domain; RBM: receptor of binding motif; FP: fusion peptide, HR1/2: heptad
655	repeat 1/2; TM: transmembrane domain; CT: cytoplasmic tail; NTD: N-terminal domain; CTD: C-
656	terminal domain; SD1: subdomain 1; SD2: subdomain 2. The structure of the S protein trimer of
657	SARS-CoV-2 and potential influence of D614G mutation on its structural change. B.
658	Experimentally determined structure of SARS-CoV-2 S protein trimer (PDB ID is 6VSB and the
659	amino acid sequences is the same as WH01 isolate). C. D614-K854 inter-monomer salt bridge. D.
660	G614-K854 inter-monomer salt bridge. The distance of the salt bridge is increased from 2.6 to 5.2
661	Å in D614G mutation as shown.

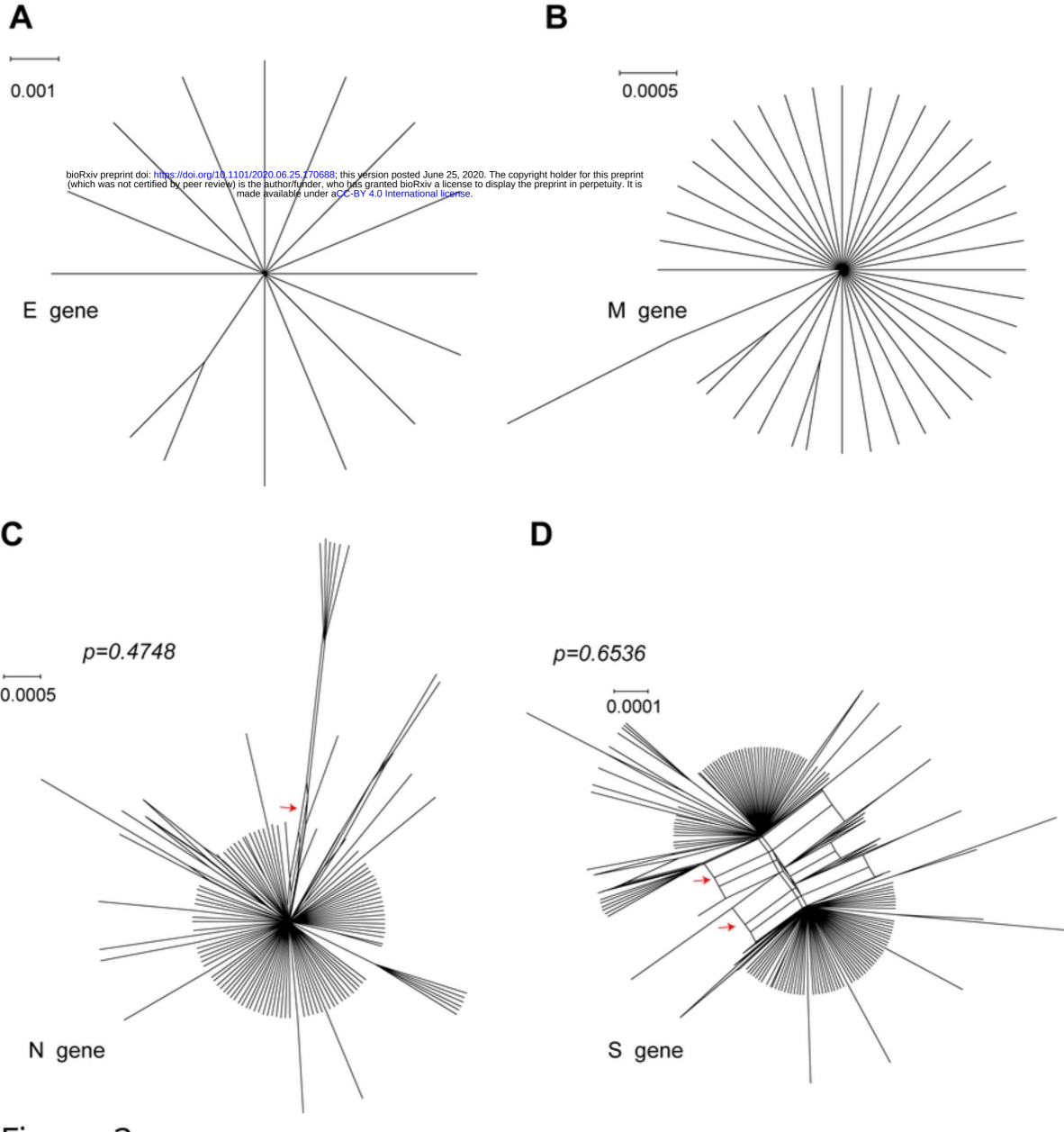
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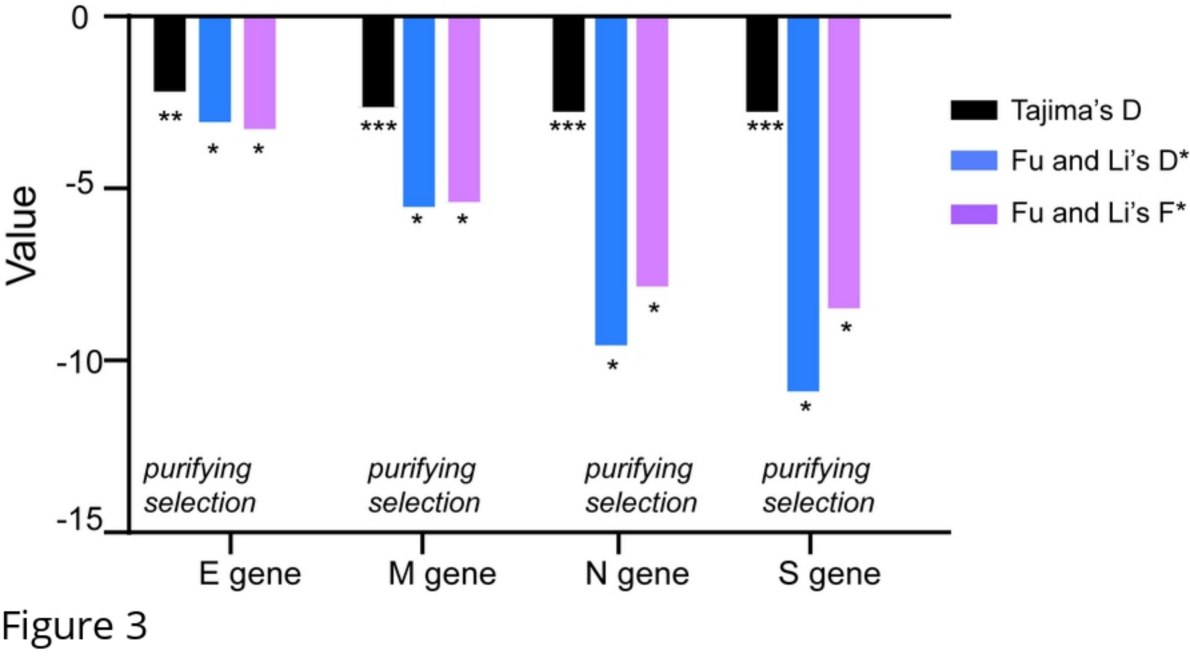




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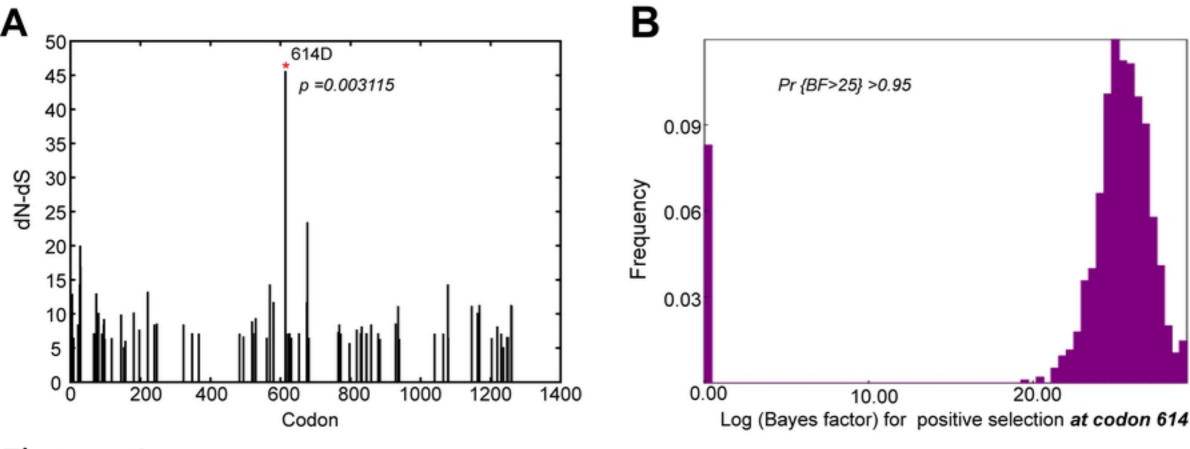
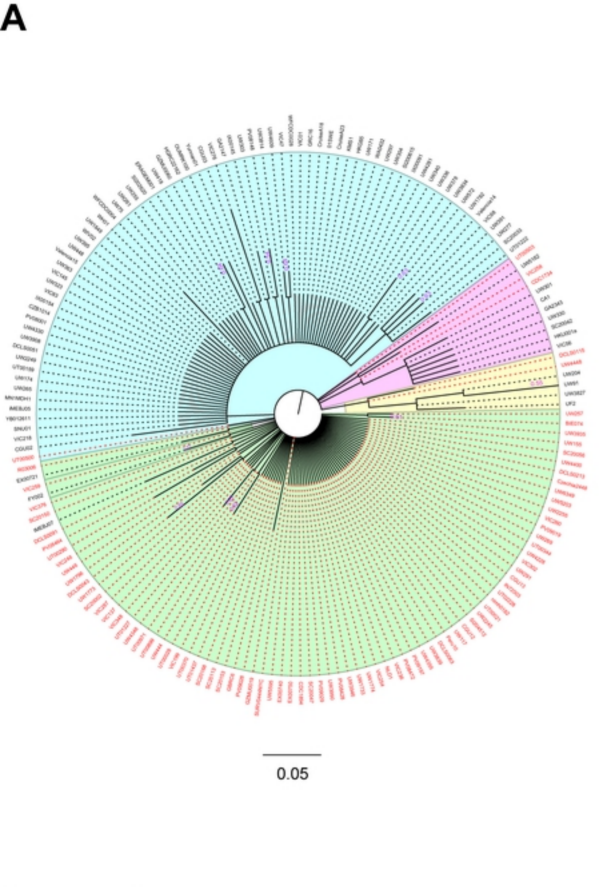
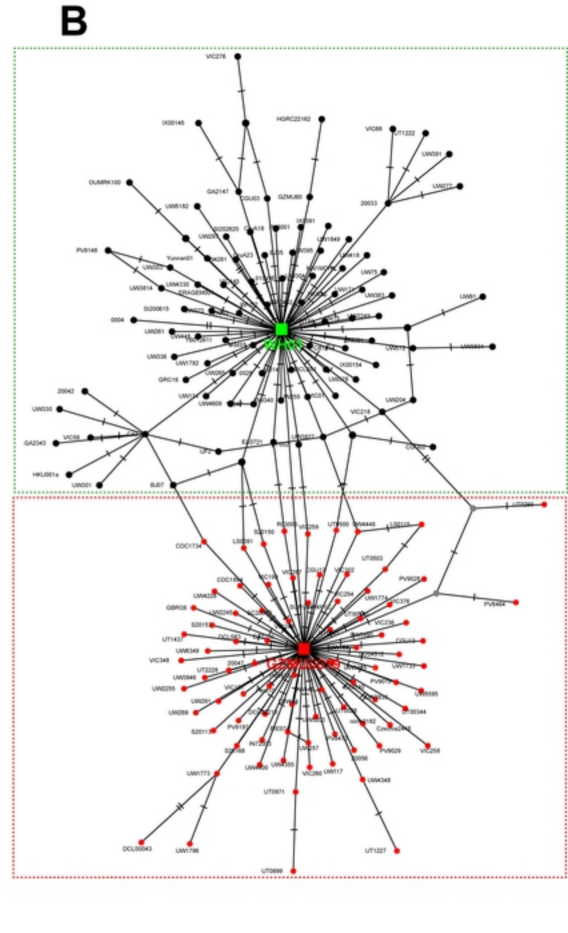
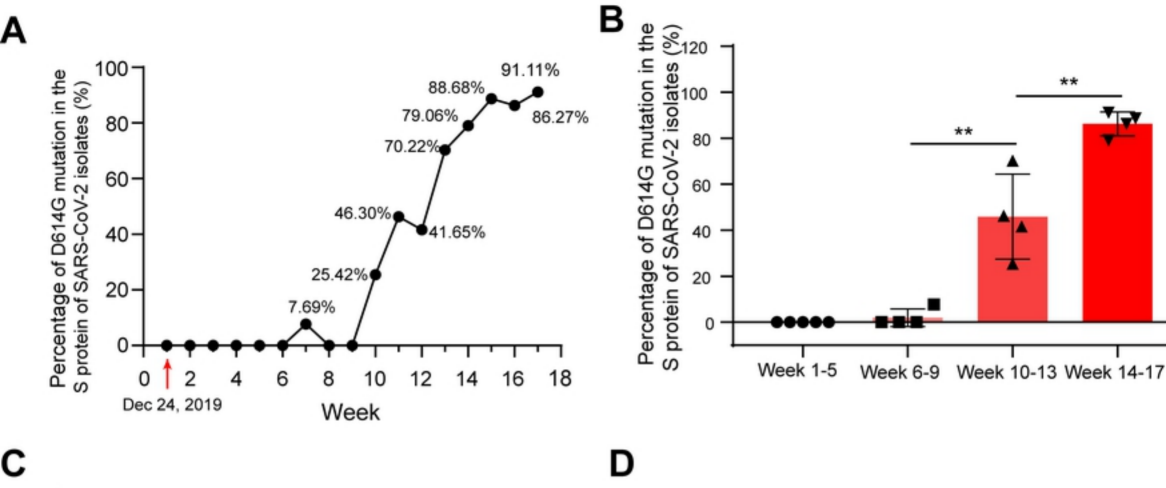
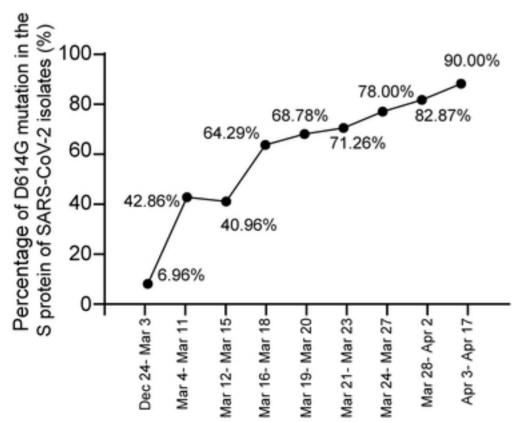


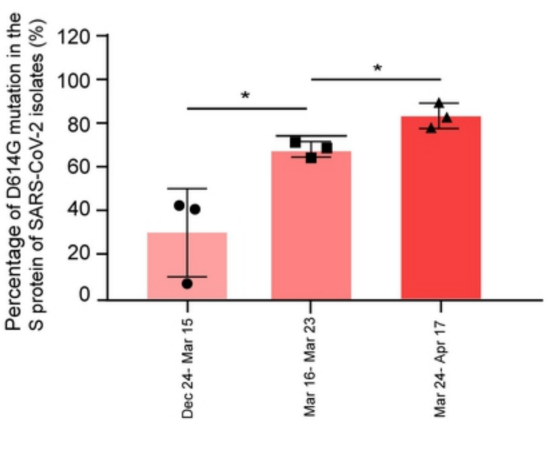
Figure 4











**T**.7