

1 **Following the Trail of One Million Genomes: Footprints of SARS-CoV-2 Adaptation to Humans**

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10 **Abstract**

11 Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) has accumulated ge-
12 nomic mutations at an approximately linear rate since it first infected human populations in late
13 2019. Controversies remain regarding the identity, proportion, and effects of adaptive muta-
14 tions as SARS-CoV-2 evolves from a bat- to a human-adapted virus. The potential for vaccine-
15 escape mutations poses additional challenges in pandemic control. Despite being of great in-
16 terest to therapeutic and vaccine development, human-adaptive mutations in SARS-CoV-2 are
17 masked by a genome-wide linkage disequilibrium under which neutral and even deleterious
18 mutations can reach fixation by chance or through hitchhiking. Furthermore, genome-wide
19 linkage equilibrium imposes clonal interference by which multiple adaptive mutations compete
20 against one another. Informed by insights from microbial experimental evolution, we analyzed
21 close to one million SARS-CoV-2 genomes sequenced during the first year of the COVID-19
22 pandemic and identified putative human-adaptive mutations according to the rates of synony-
23 mous and missense mutations, temporal linkage, and mutation recurrence. Furthermore, we
24 developed a forward-evolution simulator with the realistic SARS-CoV-2 genome structure and
25 base substitution probabilities able to predict viral genome diversity under neutral, background
26 selection, and adaptive evolutionary models. We conclude that adaptive mutations have
27 emerged early, rapidly, and constantly to dominate SARS-CoV-2 populations despite clonal
28 interference and purifying selection. Our analysis underscores a need for genomic surveillance
29 of mutation trajectories at the local level for early detection of adaptive and immune-escape
30 variants. Putative human-adaptive mutations are over-represented in viral proteins interfering
31 with host immunity and binding host-cell receptors and thus may serve as priority targets for de-
32 signing therapeutics and vaccines against human-adapted forms of SARS-CoV-2.

33

Introduction

34 Evolution in action: a trail of one million viral genomes

35 The 2002-2004 severe acute respiratory syndrome (SARS) coronavirus outbreaks had
36 multiple origins (Chinese SARS Molecular Epidemiology Consortium 2004). In contrast, severe
37 acute respiratory syndrome coronavirus 2 (SARS-CoV-2), the causative agent of the COVID-
38 19 pandemic, showed nearly 100% sequence identity among the first outbreak strains from
39 China, suggesting a single point of viral breach (Lu *et al.* 2020; Zhou *et al.* 2020). However,
40 sequence diversity quickly accumulated as COVID-19 spread globally and remained uncon-
41 trolled a year later (Andersen *et al.* 2020; To *et al.* 2021). This high-stake case of evolution in
42 action has brought unprecedented health, economic, and social devastation in modern times
43 (Peeri *et al.* 2020; Kissler *et al.* 2020; To *et al.* 2021). Many of the evolutionary mechanisms
44 driving SARS-CoV-2 genome diversification are unknown and urgently require elucidation. For
45 example, to what extent has SARS-CoV-2 adapted to its new human hosts after one year of
46 genome evolution (Phan 2020; Cagliani *et al.* 2020; Bai *et al.* 2020; Yang *et al.* 2020)? Moreo-
47 ver, how long can the global vaccine campaigns, most of which rely on vaccines formulated on
48 basis of the bat-adapted viral genome, maintain effectiveness against the waves of new viral
49 variants emerging worldwide (Koyama *et al.* 2020; Burton and Topol 2021)?

50 The vast number of publicly available SARS-CoV-2 genomes – expected to surpass a
51 million before June 1, 2021 – offers unique opportunities for understanding the evolutionary
52 processes accompanying the rapid emergence of a new viral pathogen, while challenging the
53 ability to translate evolutionary understandings into the control and prevention of current and
54 future pandemics (de Wit *et al.* 2016; Hadfield *et al.* 2018; Cui *et al.* 2019; Benvenuto *et al.*
55 2020; Andersen *et al.* 2020; Cagliani *et al.* 2020). Here we tested the hypothesis of rapid adap-
56 tation of SARS-CoV-2 genomes to human populations during the first year of the global
57 COVID-19 pandemic. We focused on developing methods and computational tools for identify-
58 ing human-adaptive mutations in the genomes of zoonotic viral pathogens. Identifying human-
59 adaptive mutations is essential to uncovering the molecular mechanisms underlying the transi-
60 tion of SARS-CoV-2 from bat to human hosts, as well as the viral mechanisms of human path-
61 ogenesis and virulence (Cagliani *et al.* 2020). For disease treatment and prevention, human-
62 adaptive mutations are prime targets for the development of therapeutics against human-
63 adapted SARS-CoV-2 variants as well as the development of broadly effective escape-proof
64 vaccines (Burton and Topol 2021; Cohen *et al.* 2021).

65 Challenges of identifying adaptive mutations in an asexual microbial population

66 Despite the benefits of a small genome size (~30,000 base pairs) and an abundance of
67 geographically and longitudinally marked genome samples, identifying signatures of natural
68 selection in SARS-CoV-2 is hindered by the challenge of a compact, gene-rich genome with
69 few non-coding sequences, as is typical for microorganisms (DeLong 2004; Rocha 2018). Se-
70 quence evolution at non-coding loci in eukaryotic species hews closely to the standard Neutral
71 Theory of molecular evolution, thus providing a powerful control for testing the presence of
72 natural selection in functional genomic regions (Garud *et al.* 2015; Koropoulis *et al.* 2020). For
73 example, presence of balancing (i.e., diversifying) selection at the *Adh* locus in *Drosophila* was
74 discovered by an excess of nucleotide polymorphisms in the coding region relative to the 5'-
75 flanking sequences (HKA test) (Hudson *et al.* 1987). The unexpected decrease in non-coding
76 sequence diversity in genomic regions with low recombination rates has led to the discovery of
77 pervasive purifying (i.e., negative) selection in *Drosophila* and humans (Hudson and Kaplan
78 1995; Charlesworth 2013; Campos and Charlesworth 2019). Likewise, adaptive mutations
79 cause selective sweeps and reduce genetic diversity at linked non-coding loci (Sabeti *et al.*
80 2002; Garud *et al.* 2015).

81 Genome-wide linkage disequilibrium (LD) imposes an additional, more severe constraint
82 for detecting adaptive mutations during SARS-CoV-2 evolution in human populations. In bacte-
83 rial species, recombination is infrequent, yet it occurs at rates high enough to uncouple the
84 evolution of loci under diversifying selection (e.g., loci encoding surface antigens) from evolution
85 of housekeeping loci under purifying selection (Milkman and Bridges 1990; Haven *et al.* 2011;
86 Bobay *et al.* 2015). In sexual populations, proportions of adaptive amino-acid divergence could
87 be estimated at a protein-coding locus by contrasting levels of synonymous and nonsynony-
88 mous substitution rates within and between species (MK test) (McDonald and Kreitman 1991;
89 Charlesworth and Eyre-Walker 2006). However, the standard MK test severely underestimates
90 adaptive divergence in asexual populations due to accumulation of slightly deleterious non-
91 synonymous mutations (Charlesworth and Eyre-Walker 2008; Messer and Petrov 2013). Fur-
92 thermore, both background selection and selective sweeps (“genetic draft”) reduce the effec-
93 tive population size and elevate the chance of random fixation of neutral and deleterious non-
94 synonymous mutations in an asexual population, thus biasing the estimation of adaptive muta-
95 tion rates (Gillespie 2000; Messer and Petrov 2013).

96 Similarities between microbial experimental evolution and SARS-CoV-2 evolution

97 Experimental evolution under controlled laboratory conditions using microorganisms

98 provides perhaps the most pertinent model for understanding SARS-CoV-2 evolution in hu-
99 mans (Lenski 2017; Good *et al.* 2017; Cvijović *et al.* 2018; Bergh *et al.* 2018). Although SARS-
100 CoV-2 is a non-free-living organism evolving under open and diverse environmental condi-
101 tions, SARS-CoV-2 populations share several key evolutionary characteristics with microbial
102 populations in long term evolution experiments (LTEEs) (Lenski 2017; Cvijović *et al.* 2018).
103 First, both evolving systems were seeded with a single genetically identical clone. Second,
104 species in both systems were microorganisms containing a compact and gene-rich genome
105 with few non-coding loci. Third, both systems had large populations in which natural selection
106 was expected to prevail over genetic drift. For example, in a population with $N_e = 1000$ individ-
107 uals, any mutation with a selection coefficient $|s| > 0.001$ would cross the neutral barrier
108 $N_e s = 1$ and evolve deterministically towards fixation or extinction. Fourth, although capable of
109 recombination, populations in both systems evolved clonally without detectable levels of genet-
110 ic exchange among coexisting individuals. Thus, both systems evolved under genome-wide LD
111 and were expected to show strong clonal interference (Lang *et al.* 2013; Lenski 2017; Good *et*
112 *al.* 2017). Fifth, populations in both evolving systems were tracked in great genetic detail
113 through whole-genome sequencing of temporally sampled isolates with spatial replication. Re-
114 sembling the replicated populations in LTEEs, SARS-CoV-2 subpopulations in six continents
115 (Asia, Africa, Europe, North America, South America, and Oceania) allowed for detection of
116 adaptive changes based on recurring genetic events.

117 As expected given the strong similarities in key evolutionary characteristics, we found
118 that SARS-CoV-2 populations during the COVID-19 showed similar adaptive dynamics as the
119 *E. coli* populations in LTEE, including the early rise and rapid fixation of adaptive mutations,
120 competing adaptive mutations, and recurrent genetic changes at key gene loci. Previous anal-
121 yses of genome evolution of SARS coronaviruses have relied mainly on phylogenetic ap-
122 proaches to identify adaptive genes, haplotypes, and lineages (Chinese SARS Molecular Epi-
123 demiology Consortium 2004; Phan 2020; Cagliani *et al.* 2020; Bai *et al.* 2020; Yang *et al.*
124 2020). Crucially, without generating mutation spectra expected under neutral, background se-
125 lection, and adaptive evolution models, these studies have been unable to test competing evo-
126 lutionary models or to explore adaptive dynamics at the level of individual mutations. In LTEEs,
127 the neutrally evolving populations were created by bottleneck events during serial transfer of
128 cultures from one generation to another (Tenaillon *et al.* 2016). Here, we used *in silico* simula-
129 tion of SARS-CoV-2 genomes evolving under neutral and selective models for understanding
130 and predicting SARS-CoV-2 evolution during the COVID-19 pandemic.

131

Material & Methods

132 [CoV genome simulator and the associated software system](#)

133 *In silico* simulations are a powerful approach to test evolutionary hypotheses by providing
134 fully specified evolutionary processes and parameters as models of species evolution in
135 nature (Yuan *et al.* 2012). However, software tools for simulating the evolution of the gene-rich,
136 finite-size microbial genomes such as those of SARS-CoV-2 are lacking. Simulations based on
137 coalescent (backward-evolution) are highly efficient but are more suitable for modeling the
138 evolution of neutral loci and relatively simple forms of selective and demographic mechanisms
139 (Hudson 2002; Liang *et al.* 2007; Kelleher *et al.* 2016). Software tools based on forward-
140 evolution simulations are less efficient but more flexible in modeling arbitrary selective and
141 demographic forces (Carvajal-Rodríguez 2008; Hernandez 2008; Haller and Messer 2019).
142 For simulating microbial genome evolution, two coalescent-based software tools implemented
143 the realistic form of homologous recombination in bacterial genomes, but were not designed to
144 simulate protein-coding sequences or the strong purifying and positive selective forces com-
145 monly operating on the gene-rich microbial genomes (Didelot *et al.* 2009; Brown *et al.* 2016).
146 Furthermore, to our knowledge all existing simulation software implements infinite-site models
147 of nucleotide substitutions. Consequently, these software tools do not allow for estimation of
148 the chances of recurrent mutations at the same sites, an aspect that cannot be ignored in a
149 rapidly expanding viral population with a small genome, such as SARS-CoV-2 populations dur-
150 ing the COVID-19 pandemic.

151 Previously, we used forward-simulation to validate the origin and maintenance of high
152 sequence diversity at a major surface antigen locus in the Lyme disease bacterium (*Borrelia*
153 *burgdorferi*) by negative frequency-dependent selection (Haven *et al.* 2011). Here we devel-
154 oped a CoV genome evolution simulator (*CovSimulator*) and used it to test whether patterns of
155 CoV genome variability fit better with expectations from neutral (NEU), background-selection
156 (BKG), adaptive (ADPT), or mixed (MIX) evolution models. The software system associated
157 with the *CovSimulator* is diagramed in Supplemental Material Fig S1.

158 Briefly, *CovSimulator* first read the annotated genome of a viral progenitor provided in
159 GenBank format (e.g., Wuhan-Hu-1, GenBank accession NC_045512). It captured the reading
160 frames of the 25 protein-coding loci (Table 1) in the SARS-CoV-2 genome such that coding (or
161 non-coding) information associated with each base of the genome was stored. At a protein-
162 coding nucleotide site, the stored genomic information included the gene locus, codon, amino
163 acid, and codon position. Simulation was initialized with a population of N identical ancestral

164 genomes, each of which was assigned the unit fitness value (see evolution parameters in Ta-
165 ble 2). During each of the total number of g generations, each individual encountered a Pois-
166 son-distributed number of point mutations with the mean genome mutation rate m . If a muta-
167 tion occurred, a uniformly distributed genome position was chosen and an alternative nucleo-
168 tide was selected as the substitute according to the base-substitution frequencies gathered
169 from viral genomes (see section below). Similarly, homologous recombination during each
170 generation occurred with a Poisson distributed mean rate of r per genome. If a recombination
171 event occurred, two individuals from the population were randomly chosen and a uniformly dis-
172 tributed genome position was selected as the break point. Two new individual genomes were
173 created by exchanging the sequences right and left of the break point. Fitness values of the
174 new genomes were re-computed according to a new set of mutated sites.

175 Crucially, we defined the fitness of a simulated viral genome as its adaptiveness to the
176 human host relative to the ancestral viral genome. That is, the fitness of the ancestral viral ge-
177 nome to the human host was defined as one. A simulated viral population displaying an aver-
178 age fitness > 1 could thus be interpreted as being better adapted than the ancestral genome to
179 the human host. A simulated viral population with an average fitness $= 1$ was considered
180 equally fit as the ancestral viral genome to reproduce in the human host. Otherwise, a simula-
181 ted viral population with an average fitness < 1 was considered less fit than the ancestral ge-
182 nome to use the human host. To implement this fitness scheme, we determined synonymous
183 or missense mutations and computed the fitness value of a simulated genome according to
184 comparison with the ancestral viral genome rather than its parental genome. This fitness defi-
185 nition is equivalent to measuring fitness gains in an LTEE study through competing the evolved
186 strains with the original, pre-evolved strain (Lenski 2017).

187 The fitness of an individual genome was the multiplicative product of its composite co-
188 dons. Thus, the fitness of the individual was set to zero if the mutation introduced a stop codon
189 (nonsense) or changed a stop codon into a sense codon (reading-frame extension). Other-
190 wise, the mutation introduced an amino-acid change (missense mutation). In the neutral mod-
191 el, the fitness of an individual remained unchanged by missense mutations. In the background
192 selection model, a missense mutation had a probability of u (e.g., $u=0.8$) of decreasing the fit-
193 ness of its carrier genome by a factor of, e.g., $w=0.95$. In the adaptive evolution model, in con-
194 trast, a missense mutation had a small probability of v (e.g., $v=0.1$) of increasing the fitness of
195 its carrier genome by a factor of, e.g., $w=1.05$. The fitness of the individual was unchanged if
196 the mutation occurred at a non-coding (intergenic or untranslated) site or introduced a synon-

197 ymous amino acid.

198 The probability of an individual to produce an offspring in the next generation was de-
199 termined by its fitness. Specifically, a threshold value between 0 and 1 was computed as an
200 increasing function of the fitness of an individual $c = 1 - e^{-w}$. A random number p between 0
201 and 1 was chosen. If $p < c$, then the individual was able to contribute one offspring. Otherwise,
202 the individual did not have a chance to reproduce. The parental population was repeatedly
203 sampled with replacement for reproduction until the constant population size of N was
204 reached. To validate the genome simulator, we compared the sample statistics with neutral
205 expectations including the level of sequence polymorphism at mutation-drift balance ($\theta =$
206 $2N_e\mu_0$), the rate of sequence divergence with respect to the ancestor ($k=mt$), and the length
207 and shape of genome genealogies under neutral and selective evolution.

208 [Viral genome database and the associated software system](#)

209 Viral genomes and associated information on the geographic origins and collection
210 dates were obtained from GISAID monthly according to submission dates (Shu and McCauley
211 2017). SNVs and indels in each genome with respect to the reference genome (Wuhan-Hu-1,
212 GenBank accession NC_045512) were identified by using the program DNADIST in the Nu-
213 cmer4 package (Marçais *et al.* 2018). To minimize sequencing errors, SNVs at genome ends
214 where missing bases were common were excluded, as were any genomes with more than
215 10% missing bases at SNV sites. Unique haplotypes were obtained with custom Perl scripts
216 based on the BioPerl package (Stajich *et al.* 2002). Isolate information, variants, and haplo-
217 types were deposited into a custom relational database (“cov-db”) to facilitate downstream
218 computational analysis. A custom Python script sampled viral genomes (e.g., $n=100$) by month
219 and at three spatial scales (continent, country, and state). The script also filtered variants and
220 output only the most frequently occurring (e.g., >0.5%) variants. A secondary Python script
221 produced a variant call format (VCF) file based on the sampled isolates and high-frequency
222 variants. Evolutionary statistics, including variant frequencies, linkage disequilibrium (r^2), hap-
223 lotypes, and base substitution frequencies were generated with programs BCFTools and
224 VCFTools (Danecek *et al.* 2011). We used Haploview (version 4.2) to calculate LD scores (D'
225 and r^2) as well as their statistical significance between pairs of SNVs (Barrett 2009). The most
226 parsimonious haplotype networks were estimated with the program TCS ver1.21 (Clement *et*
227 *al.* 2000) and visualized with tcsBU (Múrias dos Santos *et al.* 2016). To visualize genome vari-
228 ants and haplotype networks, we developed a custom web interface (<http://genometracker.org>)
229 using a similar software system supporting *BorreliaBase*, a comparative genomics browser of

230 Lyme disease pathogens (Di *et al.* 2014). The software system associated with the “cov-db”
231 database is diagramed in Supplemental Material Fig S2.

232 Evolution rates, linkage disequilibrium, and homoplasy

233 We estimated the SARS-CoV-2 genome divergence rate from the ancestor by performing
234 a linear regression of sequence differences to the reference genome (NC_045512) with
235 respect to the genome collection dates. The expected variance of the evolutionary rate was
236 estimated according to a Poisson model, which specified, at each time point of t days, an ex-
237 pected number of sequence differences in $\lambda = \mu t L$, where μ being the rate of base substitution
238 per site per day obtained from the regression line and L being the length of the reference ge-
239 nome (NC_045512, $L=29903$). The variance of the Poisson expected difference was expected
240 to be equal to the difference itself ($\sigma^2 = \lambda$).

241 To compare cross-species rates of amino-acid substitutions at protein-coding loci, we
242 downloaded the genomes of 24 viral isolates belonging to the family Coronaviridae. The viral
243 isolates included coronaviruses closely related to SARS-CoV and SARS-CoV-2 and consisted
244 of Wuhan-Hu-1, RaTG13, P1E, P5L, ZC45, ZXC21, SC2018, HuB2013, Shaanxi2011, HKU3-
245 1, Rm1, CoV273, GX2013, Rf4092, YN2013, GD01, SZ3, WIV16, SHC014, YN2018B,
246 As6526, Rs4247, Rs672, and Yunnan2011. Homologous protein sequences were aligned and
247 individual alignments were concatenated using the sequence utility *bioaln* from the BpWrapper
248 software suite (Hernández *et al.* 2018). Per-site substitution rates, normalized to a mean rate
249 of zero, were obtained with *rate4site* (Pupko *et al.* 2002).

250 We used Haploview (version 4.2) to calculate linkage disequilibrium (LD) scores (D' and
251 r^2) as well as their statistical significance (LOD, log odds) between pairs of SNVs (Barrett
252 2009). We used the DNAPARS program of the PHYLIP (version 3.696) package to search for
253 a maximum parsimony tree of unique haplotypes, obtaining the homoplasy index (HI) and the
254 number of base substitutions at each SNV site (Felsenstein 1989). The HI is defined as
255
$$HI = 1 - \frac{1}{num.sub.}$$
 at each SNV site and is zero when the alleles are consistent with the tree
256 (i.e., the number of substitution for a bi-allelic SNV is one).

257 Analysis of synonymous and missense evolutionary rates

258 Genome-wide numbers of synonymous (D_s) and nonsynonymous (D_n) nucleotide diver-
259 gence were obtained through comparison of the reference genome (Wuhan-Hu-1, GenBank
260 accession NC_045512) to its closest known relative (RaTG13, GenBank accession
261 MN996532) (Zhou *et al.* 2020) with the program DNADIST (Marçais *et al.* 2018). Genome-wide

262 synonymous (P_s) and nonsynonymous (P_n) nucleotide polymorphisms in viral populations were
263 estimated with the use of viral samples. In computing the per site synonymous and nonsynon-
264 ymous substitution rates ($d_s=D_s/S$, $d_n=D_n/N$, $p_s=P_s/S$, $p_n=P_n/N$), the effective numbers of avail-
265 able synonymous (S) and nonsynonymous (N) sites at each gene locus must be estimated
266 (Yang 2007). We estimated S and N empirically by using the *CovSimulator*, which accounts for
267 both the genome base composition bias and the strong mutation biases (Supplemental Materi-
268 al Fig S3). Specifically, we ran *CovSimulator* with a high genome mutation rate $m=10$ and a
269 population size $p=200$ for $n=10$ generations, generating an expected total number of 20,000
270 mutation events or $\lambda=0.67$ mutations per genomic site, on average. Assuming a Poisson distri-
271 bution, the proportions of genomic sites encountering 0, 1, and >1 point mutations were ex-
272 pected to be 51.2%, 34.3%, and 14.5%, respectively. Thus, the probability of a site not being
273 mutated was $p=0.512$. To ensure that all genomic sites were mutated at least once, we ran
274 *CovSimulator* ten times such that the chance of a site not undergoing any mutation was small
275 $p = 0.512^{10} = 1.25e-3$. The average numbers of synonymous and missense mutations from ten
276 repeated runs, normalized to gene lengths, were used as estimates of S and N (Table 1; Sup-
277 plemental Material Tables S1 and S2).

278 [Analysis of mutation trajectories](#)

279 For a simulated population, we followed the trajectories of the most frequent ($>0.5\%$
280 among all samples) missense mutations by first calculating their frequencies in each genera-
281 tion. The trajectory of a mutation X was represented by an n -dimensional vector $X_T =$
282 $(X_{t_0}, \dots, X_{t_n})$, where each X_{t_i} was the frequency of X within in the population at time point t_i .
283 Distance between two trajectories, X_T and Y_T , was defined as $D_{X,Y} = \sum_{i=0}^n |X_{t_i} - Y_{t_i}|$. Trajectories
284 of two or more mutations were merged into a “genotype” if the average distance between them
285 was ≤ 0.05 . A genotype (G_1) was considered as derived from (i.e., nested within) a parental
286 genotype (G_2) if their Jaccard distance $J_{1,2} = \frac{|G_1 \cup G_2| - |G_1 \cap G_2|}{|G_1 \cup G_2|}$ equaled the simplified Jaccard dis-
287 tance when G_1 was nested within G_2 , $J_{1,2} = \frac{|G_2| - |G_1|}{|G_2|}$. In the latter case, the union of the two tra-
288 jectories $|G_1 \cup G_2|$ was the same as the trajectory of the parental genotype G_2 , whereas the in-
289 tersection of the two trajectories $|G_1 \cap G_2|$ was the same as the trajectory of the child genotype
290 G_1 . The merging of mutations into genotypes and the nesting of genotypes were both carried
291 out with the Python package *muller* (version 0.6.0, <https://github.com/cdeitrick/Lollipop>) with de-
292 fault settings. Muller diagrams were subsequently generated using the R package *ggmuller*.
293 Fitness of a “genotype” was defined cumulatively (Desai and Fisher 2007). An adaptive muta-

294 tion within a cluster increases its fitness by $s>0$ and a deleterious mutation decreases its fit-
295 ness by $s<0$. If a parent cluster had fitness ns , and a child of that cluster had fitness ms , the
296 fitness of the child was $(n+m)s$.

297 For viral genome samples, Muller diagrams, which depict mutation frequencies within a
298 single evolving population, are in general not applicable. Since viral genomes were sampled
299 from multiple outbreak locations, it would be misleading to perform analysis including merging
300 of mutations into genotypes and inference of parental and child genotypes. Thus, we used
301 heatmaps as an alternative approach to follow the trajectories of high-frequency mutations in
302 both simulated and viral populations. The R package *pheatmap* was used to generate
303 heatmaps. As in the Muller diagrams, mutations in a heatmap were grouped into hierarchical
304 clusters based on similarities in frequencies over time. Similarity between a pair of mutation
305 trajectories i , and j was defined as $d = 1 - \text{cor}(i, j)$, where $\text{cor}(i, j)$ was the Pearson's correlation
306 coefficient. Unlike in the Muller diagrams, however, mutations with similar frequency trajec-
307 tories were not merged into "genotypes". Nor did the heatmap analysis estimate parent-
308 descendant relationships among mutation clusters.

309 **Data and software availability**

310 SARS-CoV-2 genome sequences and the associated viral isolate information are avail-
311 able from the GISAID EpiCoV™ database (Shu and McCauley 2017). Software tools associat-
312 ed with *CovSimulator*, the forward-evolution simulator, and *cov-db*, the custom database of
313 SARS-CoV-2 genome variability, are available at the Github repository
314 (<https://github.com/weigangq/cov-db>). Programmatic access to the *cov-db* database is availa-
315 ble upon request. Also available in the same Github repository are key datasets including mu-
316 tation trajectories from simulated evolution and VCF files of viral genomes sampled monthly. A
317 web interface to the *cov-db* database is publicly available at <http://cov.genometracker.org>.

318 **Results and Discussion**

319 ***CovSimulator*: a realistic SARS-CoV-2 genome evolution simulator**

320 The SARS-CoV-2 genome is biased in base composition (62.0% AT) and strongly bi-
321 ased in mutation frequency. Approximately ~70% of single-nucleotide mutations occurring dur-
322 ing the pandemic were C>T or G>T substitutions (Supplemental Material Fig S3). To realisti-
323 cally simulate SARS-CoV-2 genome evolution, we used the first known SARS-CoV-2 genome
324 (from the Chinese isolate Wuhan-Hu-1 collected in December 2019) (Zhou *et al.* 2020) as the
325 progenitor and used empirically derived base substitution probabilities (Table 2). *CovSimulator*,

326 currently implemented with the standard Wright-Fisher model with constant population sizes
327 and non-overlapping generations, was validated through comparing simulated outputs with an-
328 alytical expectations including the rates of sequence divergence (Fig 1C and 1D), levels of se-
329 quence polymorphism (Fig 2C), genealogies of genome samples at the end of simulations
330 (Supplemental Material Fig S4A), and fitness values of simulated populations (Supplemental
331 Material Fig S4B).

332 We used the CovSimulator to derive theoretical expectations of synonymous and mis-
333 sense divergence, sequence diversity, and their ratios under neutral, background, adaptive,
334 and mixed evolution models (Figs 1C, 1D, 2C, and 2D). These simulated outcomes provided
335 baseline controls for estimating selective constraints (Figs 1B & 2B) as well as for understand-
336 ing evolutionary dynamics at the level of individual mutations (Fig 3). In addition, the CovSimu-
337 lator provided a simulation-based approach to estimate evolutionary parameters such as the
338 effective numbers of synonymous and nonsynonymous sites at protein-coding loci (Table 1)
339 and frequencies of recurrent mutations (see below). Such parameters are difficult to derive an-
340 alytically because it is necessary to take into account of biases in base composition as well as
341 in mutation frequency.

342 Future versions of the simulator will incorporate more realistic demographic features in-
343 cluding changing population sizes, population admixture, and additional selective mechanisms.
344 In particular, simulating SARS-CoV-2 genome evolution under negative frequency-dependent
345 selection is important to identify mutations contributing to immune escape including escape
346 from vaccines, which are expected to have higher fitness values when they rare (Haven *et al.*
347 2011; Papkou *et al.* 2019). Negative frequency-dependent mutations maintaining coexistence
348 of multiple clonal lineages have been observed in microbial LTEEs (Maddamsetti *et al.* 2015;
349 Good *et al.* 2017). In addition, the CovSimulator paves a way to estimate parameters of SARS-
350 CoV-2 evolution (e.g., population growth rates, selection coefficients, and migration rates)
351 through approximate Bayesian computation (ABC) (Lintusaari *et al.* 2017).

352 Accelerated missense divergence: rise of hyper-mutated variants

353 On the basis of ~1.0 million SARS-CoV-2 genome sequences obtained from the GISAID
354 database (Shu and McCauley 2017) up to March 31, 2021, we generated a custom database
355 of high-quality 815,402 SARS-CoV-2 genomes. We identified 8065 SNVs, 173 deletions, and
356 49 insertions, each of which was represented by 100 or more viral genomes. Genome se-
357 quences were consolidated into 350,094 haplotypes based on the SNVs and indels. We sam-
358 pled ~100 genomes monthly from each of the six continental populations and plotted the syn-

359 onymous, missense, and total mutational differences with respect to the ancestral genome
360 over collection dates (Fig 1A). The total rate of mutation accumulation (gray dots) was well
361 characterized by a linear Poisson model with a highly significant slope and a Poisson-expected
362 variance (Fig 1A). However, significant deviations from the Poisson model were observed in
363 Asian, European, Oceanian, and South American viral populations since October 2020, asso-
364 ciated with the hyper-mutated viral variants discovered first in immuno-compromised patients
365 with COVID-19 (Choi *et al.* 2020; Kemp *et al.* 2021).

366 The Poisson model of linear mutation accumulations over time initially suggested a neu-
367 tral process of viral genome divergence. However, closer examination by measuring the syn-
368 onymous and missense mutation rates separately did not support the neutral divergence mod-
369 el. The ratio of missense to synonymous mutations was expected to be high ($D_n/D_s \sim 3.0$) ac-
370 cording to the simulated neutral evolution (Fig 1C, 1st panel). In reality, the D_n/D_s ratios began
371 at a low level ($D_n/D_s \sim 1.0$) similar to that from the simulated background selection model (Fig
372 1C, 1st panel), thus suggesting considerable selective constraints during the early months (be-
373 fore April 2020) of the pandemic. The D_n/D_s ratio increased across continental populations af-
374 terward and eventually showed a marked increase after October 2020 to the levels expected
375 from the neutral and adaptive evolution models (Fig 1B). The accelerated missense diver-
376 gence, reflected in the steep rise of D_n/D_s ratios, was attributable to the emergence and spread
377 of hyper-mutated viral lineages, which accumulated predominantly missense mutations with
378 little synonymous divergence (Fig 1A). However, the acceleration of missense divergence oc-
379 curred in the North American population before the emergence of hyper-mutated viral lineages
380 therein (Fig 1B, 4th panel).

381 The hyper-mutated viral variants, which first emerged in immune-compromised patients
382 with COVID-19 (Choi *et al.* 2020; Kemp *et al.* 2021), resembled the hyper-mutable microbial
383 lineages with defective DNA repair systems that commonly emerged during LTEE studies
384 (Lenski 2017). In LTEE populations, the “mutator” phenotype was maintained because the
385 consistently higher benefits of new adaptive mutations out-weighing the cost of deleterious mu-
386 tations in a controlled environment (Lenski 2017). Similarly, in an immune-deficient host envi-
387 ronment, mutations that would have been deleterious in a normal host (e.g., those leading to
388 hyper immunogenicity) may become neutral or beneficial to viral reproduction (Choi *et al.*
389 2020; Kemp *et al.* 2021). Freed from host immune constraints, viral evolution essentially fol-
390 lows the adaptive or mixed evolution models in which adaptive lineages dominate the viral
391 population (Fig 1C and 1D, 3rd panel). Nevertheless, all missense mutations observed in the

392 hyper-mutated viral genomes are unlikely to be adaptive because neutral or slightly deleterious
393 mutations are driven to high frequencies through genetic hitchhiking in an asexual population
394 (i.e., genetic draft) (Gillespie 2000; Kim and Stephan 2000; Lang *et al.* 2013).

395 We note that, beyond adaptive mutations, the acceleration of missense divergence as
396 measured by the D_n/D_s ratio could be caused by demographic forces. In the present study, we
397 simulated viral evolution with a constant population size, although neutral and slightly deleteri-
398 ous missense mutations are expected to accumulate in the rapidly expanding viral populations
399 (Messer and Petrov 2013). In addition, our analysis combined viral samples within a continent
400 as representing a single population, whereas numerous local outbreaks and subsequent mi-
401 grations between subpopulations are expected to contribute to increased viral genome diversi-
402 ty including missense divergence (see next section).

403 [Expanding genome diversity: demographic and selective causes](#)

404 SARS-CoV-2 genomic diversity, measured by monthly average genome differences (π),
405 increased in the six continents during the first year of the COVID-19 pandemic (Fig 2A). Ex-
406 panding genomic diversity is expected for a nascent viral population before it reaches muta-
407 tion-drift balance even if the population remains at a constant size (Fig 2C). Clearly, the global
408 viral populations are far from reaching an equilibrium level of genomic diversity as the virus
409 has spread within and across continents, mirroring the failures in local and global outbreak
410 control. Furthermore, the increasing genomic diversity may be a reflection of increasing admix-
411 ture of viral subpopulations distributed across the continents.

412 Beyond demographic forces, the relaxation of selective constraints and adaptive muta-
413 tions may also have contributed to the rising viral genomic diversity. Ratios of missense to
414 synonymous polymorphisms (π_n/π_s) were generally higher in continental populations than ex-
415 pected under strong purifying selection (Fig 2B), suggesting the accumulation of neutral and
416 slightly deleterious missense mutations in the expanding viral population. Adaptive hyper-
417 mutated lineages contributed to the increase in π_n/π_s ratios in later months in most continental
418 populations and caused the elevated D_n/D_s ratios described in the previous section. An addi-
419 tional possible cause of rising viral genomic diversity is the presence of negative frequency-
420 dependent selection by which rare immune-escape variants gain a selectively advantage (Ha-
421 ven *et al.* 2011; Papkou *et al.* 2019).

422 [Asexuality, recurrent mutations, and recombination](#)

423 We estimated the genome-wide levels of LD on the basis of 93 most frequent missense

424 SNVs segregating in 8215 genomes sampled from six continents (Supplemental Material Ta-
425 ble S3). These mutations were present with a frequency of 20% or higher in at least one month
426 in one continent. Complete LD (D' close to 1) dominated the D' values between pairs of SNVs
427 and, furthermore, there was no evidence of LD decay over genomic distances between the
428 SNVs (Supplemental Material Fig S5). LD decay over distance is expected if recombination
429 among viral strains occurs with sufficient frequency. In microbial species, recombination occur-
430 ring at a rate comparable to the rate of point mutation is sufficient to cause LD decay over ge-
431 nomic distances (Fraser *et al.* 2007; Ansari and Didelot 2014). Thus we conclude that SARS-
432 CoV-2 populations during the first year of pandemic were largely asexual with little evidence of
433 recombination. The asexual population structure of SARS-CoV-2 populations mirrors the low
434 recombination rates during previous SARS and Middle East respiratory syndrome (MERS)
435 coronavirus outbreaks (Chinese SARS Molecular Epidemiology Consortium 2004; de Wit *et al.*
436 2016).

437 An analysis of SARS-CoV-2 genomes from early isolates suggested active recombi-
438 nation during human transmission based on a high level of homoplasy – independent mutations
439 occurring at the same sites that cause inconsistencies with the viral phylogeny (Yi 2020). A
440 prominent example of phylogenetically inconsistent mutation is the nonsynonymous SNV
441 TTT[Phe]/TTG[Leu] at the genomic position 11083 of the *Nsp6* locus (Yi 2020). By reconstruct-
442 ing genome phylogeny through haplotype networks, we observed a similarly high level of ho-
443 moplasy caused by either recombination or by mutations that have occurred independently in
444 multiple evolutionary lineages (Supplemental Material Fig S6). Recurrent mutations and se-
445 quencing errors may have contributed to the observed homoplasy (Turakhia *et al.* 2020). Re-
446 curring mutations are inevitable in SARS-CoV-2 with its relatively small genome size. The
447 chance of mutation recurrence increases as the pandemic spreads and persists. Indeed, we
448 were able to estimate the rate of mutation recurrence with the use of *CovSimulator*. In a simu-
449 lated population evolving under neutral conditions, ~2.9% genomic sites (860 out of the total of
450 29903 sites) experienced two or more mutations after 500 generations. This number was sig-
451 nificantly greater than expected from a random Poisson process ($p=2.1e-270$ by a χ^2 test of
452 goodness of fit). For two mutations occurring at the same genomic site, strong mutation biases
453 seen during SARS-CoV-2 genome evolution stipulate a high chance of parallel base substi-
454 tutions. For example, a mutation at a cytosine (C) site will almost certainly (with a ~95% chance)
455 result in a thymine (T) (Table 2; Supplemental Material Fig S3).

456 It should be cautioned that a clonal population structure in SARS-CoV-2 does not imply

457 an absence of or an inability of homologous recombination. In fact, coronaviruses are known
458 for their high potential for homologous recombination in natural reservoirs as well as in the la-
459 boratory conditions (Masters 2006; Denison *et al.* 2011; Cui *et al.* 2019). SARS-CoV-2 ge-
460 nomes showed a mixed ancestry containing parts of the genome from coronaviruses associat-
461 ed with the pangolin (*Manis javanica*) and other parts from related viruses associated with the
462 bat (*Rhinolophus affinis*) (Andersen *et al.* 2020; Lam *et al.* 2020). Consequently, the high clon-
463 ality of SARS-CoV-2 populations is likely to be due to the explosive population growth world-
464 wide (“epidemic structure”) rather than to an inability of recombination (Smith *et al.* 1993). By
465 reducing clonal interference among competing adaptive mutations as well as by removing de-
466 letierous mutations without decreasing the frequencies of beneficial alleles, recombination is a
467 powerful mechanism accelerating adaptation across species including microorganisms (Smith
468 *et al.* 1993; Barton and Charlesworth 1998). As such, it is important to be vigilant about the ris-
469 ing chance of recombination among SARS-CoV-2 variants as the COVID-19 pandemic be-
470 comes entrenched. Previously, we have quantified recombination rates in natural populations
471 of Lyme disease bacterium based on genome comparisons and computer simulations (Qiu *et*
472 *al.* 2004; Haven *et al.* 2011). Similarly, *CovSimulator* can be used to detect the presence of re-
473 combination and to estimate recombination rates in SARS-CoV-2 populations through a com-
474 parison of homoplasy levels in populations simulated with and without recombination.

475 [Adaptation despite background selection: a model of SARS-CoV genome evolution](#)

476 The highly clonal population structure of SARS-CoV-2 and the microbial species in
477 LTee studies implies that genetic variations across the entire genome are highly linked. As
478 such, various selective forces interfere with one another including, for example, purifying selec-
479 tion at housekeeping loci, diversifying selection at antigenic loci, and adaptive evolution at
480 host-binding sites (Hill and Robertson 1966; Gillespie 2000; Lang *et al.* 2013; da Silva and
481 Galbraith 2017; Lenski 2017; Campos and Charlesworth 2019). In addition, neutral or even
482 deleterious mutations may “ride along” with a newly emerged adaptive mutation and to reach
483 high frequency in an asexual population.

484 On the basis of the conclusions on adaptive mutations contributing to genome variability
485 and on the strong LD across the SARS-CoV-2 genome, we propose the mixed evolution as a
486 model to understand the dynamics of SARS-CoV-2 genome evolution. In the mixed model
487 (Figs 1C, 1D, 2C, and 2D), the majority of missense mutations were slightly deleterious (~80%
488 probability with a multiplicative fitness cost of 0.95) and a small proportion of missense muta-
489 tions were slightly adaptive (~10% probability with a multiplicative fitness benefit of 1.05).

490 First, we traced the genealogy of the final 20 sampled genomes, which showed a substantially
491 shortened coalescence time since the most recent common ancestor (Fig 3A). Next, we
492 tracked the frequencies of the top most frequent ($\geq 5\%$) missense mutations for 500 genera-
493 tions (Fig 3B). Adaptive mutations (11 of a total of 19 mutations) dominated the final popula-
494 tion. Nevertheless, not all fixed mutations were adaptive. Three neutral (#1, #2, and #4, in
495 gray) and three deleterious missense (#2, #3, and #4 in blue) mutations became fixed through
496 linkage with adaptive driver mutations, exemplifying genetic draft (Gillespie 2000). Conversely,
497 not all adaptive mutations were destined to be fixed, indeed, one adaptive mutation (#5, in red)
498 was lost because of competition with other adaptive mutations, exemplifying clonal interfer-
499 ence (Lang *et al.* 2013; Maddamsetti *et al.* 2015). Thirdly, we generated the Muller diagrams,
500 which grouped mutations sharing similar frequency trajectories (i.e., temporal linkage) into a
501 single “genotype” (Fig 3C). The diagrams highlighted regular selective sweeps driven by adap-
502 tive mutations. Critically, it is clear from the Muller diagrams that within each “genotype” (e.g.,
503 G1, G2, G3, G8, and G9), at least one genetic change was the driver adaptive mutation. To
504 facilitate comparison of evolutionary dynamics among evolutionary models, we provided the
505 genome genealogies of the last-generation samples and the Muller diagrams of the topmost
506 frequent missense mutations from all four models of simulated evolution as Supplemental Ma-
507 terial Fig S4.

508 In summary, the mixed evolution model illustrates that, first, adaptive mutations and vi-
509 ral lineages quickly dominate the viral population despite that most of the missense mutations
510 are deleterious. Second, neutral and deleterious mutations can become fixed through genetic
511 hitchhiking with adaptive mutations. Third, adaptive mutations can be lost because of strong
512 clonal interference. Fourth, recurring mutations become increasingly common because of a
513 small viral genome, strong mutation biases, longer evolution time, and prolonged maintenance
514 of adaptive lineages in the viral population. Fifth, temporal linkage among missense mutations
515 provides a way to identify adaptive driver mutations. These conclusions are anticipated by ob-
516 servations from microbial LTEE studies as well as by results of theoretical analysis, both of
517 which showed dominance of adaptive mutations in asexually evolving populations despite
518 presence of strong purifying selection (Kim and Stephan 2000; Desai and Fisher 2007; Lenski
519 2017). We note that only one set of evolutionary parameters was used in the present simula-
520 tion of the mixed model, the outcome of which would vary quantitatively with selection parame-
521 ters including the proportions and strengths of deleterious and adaptive mutations.

522 **Spatiotemporal characteristics of adaptive mutations**

523 The mixed model of SARS-CoV-2 genome evolution revealed a number of characteristics of adaptive mutations that are informative for their identification. First, adaptive mutations were over-represented in high-frequency SNVs (Fig 3). In one population simulated with mixed model, 258 adaptive mutations (10.9% out of a total of 2376 missense mutations) were present in the combined sampled genomes. However, 14 (45%) of the adaptive mutations were among the 31 missense mutations that have reached a frequency of 0.5% or higher. Second, the proportion of adaptive mutations among missense mutations increased over time (Fig 3). In the same simulated population, among the 20 genomes sampled from the last generation, the fixed missense mutations included 7 (70%) adaptive, 2 (20%) deleterious, and 1 (10%) neutral mutations. Third, in clusters of mutations that shared similar temporal trajectories, at least one of the consortium was the adaptive driver (e.g., G1, G2, G3, G8, and G9 in Fig 3). These characteristics of adaptive mutations suggest ways to identify adaptive mutations driving SARS-CoV-2 adaptation to humans through spatiotemporal tracking of mutation frequencies.

536 Guided by the above insights from the simulated evolution and LTEEs, we identified a genome-wide set of 101 missense mutations with a presence of 20% or higher frequency in at 537 least one month within a continent (Supplemental Material Table S4). The high-frequency mutations were most often found in genes encoding the spike (S, $n=21$, 20.8%), nucleocapsid (N, 539 $n=19$, 18.8%), Nsp3 ($n=13$, 12.9%), and ORF8 ($n=9$, 8.9%) proteins. Similarly, we identified a 540 set of 52 missense mutations on the spike protein with a presence of 5% or higher frequency 541 in at least one month within a continent (Supplemental Material Table S5). Mutations on the 542 spike protein are of particular interest because of its use as vaccinogen. Half ($n=26$) of these 543 spike protein mutations were located within the N-terminus domain (NTD) and receptor-binding 544 domain (RBD), suggesting an oversized role the NTD and RBD mutations play in driving 545 SARS-CoV-2 adaptation to humans. Sequences in NTD and RBD evolve faster relative to the 546 genome average during coronavirus divergence, further supporting the role of mutations within 547 these domains in driving viral adaptation to humans (Luk *et al.* 2019; Phan 2020; Cagliani *et al.* 548 2020) (Supplemental Material Fig S6).

550 We subsequently clustered these high-frequency mutations on the basis of their temporal 551 trajectories within each continent. A heatmap of frequency trajectories of 52 missense 552 mutations (with >5% frequency in at least one month) on the spike protein revealed clusters of 553 mutations that were distributed either across the globe or more limitedly within continents (Fig 554 4). The globally distributed mutations included the D614G substitution that arose during the

555 first month (January 2020) of the SARS-CoV-2 outbreaks in Asia and quickly reached global
556 fixation. Clinical and experimental studies suggested enhanced human transmissibility but not
557 increased disease severity associated with D614G viral variants (Korber *et al.* 2020; Volz *et al.*
558 2021; Plante *et al.* 2021). It is possible that missense mutations strongly linked with the D614G
559 mutation, including P323L in Nsp12 (RNA polymerase) and R203K and G204R in the N (nu-
560 cleocapsid) protein, may have also played a role in increased viral fitness in humans (Yang *et*
561 *al.* 2020).

562 A temporally linked group of six spike protein mutations – N501Y, P681H, T716I,
563 D1118H, S982A, and A570D – associated with the hyper-mutated B.1.1.7 lineage (Fig 4) that
564 emerged in September 2020 in England and quickly spread worldwide represent another set of
565 mutations that have enhanced viral transmission in humans (Galloway 2021; Kemp *et al.*
566 2021). These spike protein mutations pose the additional risk of viral escape from protective
567 immunity elicited with vaccines designed on the basis of the bat-adapted progenitor genome
568 (Wang *et al.* 2021; Collier *et al.* 2021). Other mutations have so far been confined within one or
569 more continents and have not reached global presence. These spike protein mutations includ-
570 ed those associated with the B.1.351 lineage in Africa, the P.1 lineage in South America, the
571 B.1.427/B.1.429 lineages in North America, and the B.1.617 lineage in Asia (Fig 4).

572 The strong candidates of human-adaptive mutations shown in the above have risen rel-
573 atively early during the pandemic. The latest emergent human-adaptive mutations, however,
574 would first reach high frequencies only in local outbreak populations. Thus, it is necessary to
575 track allele frequencies at regional levels for early detection of human-adaptive mutations. As
576 an example, we tracked the spatiotemporal frequencies of 56 spike missense mutations with
577 ≥5% allele frequencies in at least one month in the United States and and its five states includ-
578 ing Washington, California, New York, Texas and Michigan (Fig 5). Except for the globally
579 fixed D614G mutations and mutations associated with the B.1.1.7, B.1.427 and B.1.429 line-
580 ages, mutations associated with the latest emergent viral lineages only reached the threshold
581 5% level within the states and not at the national level. For example, the B.1.526 and B.1.243
582 lineages emerged during December 2020 in New York and have not yet spread to the other
583 four states. The B.1.2 lineage in Washington and the B.1.234 lineage in Michigan have thus far
584 been observed only within the states.

585 In summary, whereas missense mutations in the SARS-CoV-2 genome that have
586 reached high frequency at local or global levels are not necessarily human-adaptive mutations
587 because of the possibilities of genetic drift and hitchhiking, clusters of missense mutations that

588 display temporal linkage and have reached high frequencies are indicative of adaption to hu-
589 mans. Within each of the cluster of temporally linked high-frequency missense mutations, we
590 expect at least one to be a human-adaptive driver mutation. As such, the high-frequency clus-
591 ters of missense mutations are top-priority candidates for clinical development of therapeutics
592 and vaccines that target human-adapted viral variants.

593 **Concluding remarks**

594 In the present work, we used realistic simulations of genome evolution and insights from
595 microbial long-term evolution experiments (LTEEs) (Tenailleon *et al.* 2016; Cvijović *et al.* 2018)
596 to understand the evolutionary transition of the SARS-CoV-2 virus from a bat-adapted to a
597 human-adapted pathogen. The two evolving systems share salient evolutionary characteristics
598 including strong purifying selection associated with a compact genome and large population
599 sizes, forced adaptation to a new environment, and an asexual population structure. Not sur-
600 prisingly, the variety of adaptive dynamics occurred in LTEEs were all discernable during
601 SARS-CoV-2 evolution including the early rise and rapid fixation of adaptive mutations, clonal
602 interference with competing adaptive mutations, fixation of neutral and deleterious mutations
603 due to genetic hitchhiking. Specifically, both LTEEs and our analysis suggest that temporal
604 linkage among mutations is a sensitive means for identifying emerging human-adaptive muta-
605 tions and vaccine-escape mutations, particularly when mutation frequencies are tracked at the
606 local and regional levels.

607 Epidemiological models based on human coronaviruses and influenza viruses predict
608 the COVID-19 to be a recurrent seasonal disease in the next 2 ~ 5 years (Kissler *et al.* 2020;
609 Cobey 2020). We expect to see continued expansion of viral genome diversity as the pandem-
610 ic persists, entailing increasing risks for viral adaptation to humans and viral escape from natu-
611 ral and vaccine-induced protective immunity. Prolonged pandemic incurs the additional risks of
612 recurrent mutations and recombination among viral variants, which would accelerate viral ad-
613 aptation to humans. The software systems we developed facilitate real-time tracking of SARS-
614 CoV-2 outbreaks. The *CovSimulator* software system is capable of modeling the trajectories of
615 SARS-CoV-2 genome evolution and could be further improved by including more realistic pa-
616 rameters such as population expansion, migration and admixture between subpopulations, and
617 frequency-dependent fitness imitating vaccine-escape mutations. The second software system
618 associated with the *cov-db* genome database is capable of rapid tracking of emergent adaptive
619 mutations through temporal sampling of genomes in a continent, country, or region therein.
620 Thirdly, the cov.genometracker.org website provides a public-friendly user interface to search,

621 browse, and visualize SARS-CoV-2 genome evolution and mutation trajectories (Supplemental
622 Material Fig S7). Mutations appear in genes encoding proteins that down-regulate host im-
623 mune responses (e.g., ORF3a and ORF8) and bind host cells (e.g., Spike) are high priority
624 targets for the development of therapeutics and vaccines against human-adapted SARS-CoV-
625 2 variants.

626

627 **List of Supplemental Material**

628 • Supplemental Tables

629 ○ Table S1. Estimated total number of synonymous and nonsynonymous sites
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635 ○ Fig S1. The CovSimulator software system
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642

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653 The authors declare no conflicts of interest.

654 Author Contributions

655 S. Akther initiated the project, performed all evolutionary analysis, and participated in
656 manuscript drafting. E. Bezrucenkovas downloaded viral genomes, prepared associated
657 metadata, and developed the prototype evolution simulator. L. Li participated in the development
658 of the simulator and performed simulation, linkage, and evolutionary rate analyses. B.
659 Sulkow contributed to conceptual development and performed Muller diagram analysis. L. Di
660 developed the genome database and web browser. D. Pante contributed to haplotype network
661 and mutation analyses, C.L. Martin contributed to mutation analysis. B.J. Luft contributed to
662 conceptual development and manuscript preparation. W. Qiu designed the software systems,
663 implemented the simulator, and drafted the manuscript.

664 References Cited

665 Andersen K. G., A. Rambaut, W. I. Lipkin, E. C. Holmes, and R. F. Garry, 2020 The proximal origin of SARS-
666 CoV-2. *Nat. Med.* 1–3. <https://doi.org/10.1038/s41591-020-0820-9>

667 Ansari M. A., and X. Didelot, 2014 Inference of the properties of the recombination process from whole bacterial
668 genomes. *Genetics* 196: 253–265. <https://doi.org/10.1534/genetics.113.157172>

669 Bai Y., D. Jiang, J. R. Lon, X. Chen, M. Hu, *et al.*, 2020 Comprehensive evolution and molecular characteristics
670 of a large number of SARS-CoV-2 genomes reveal its epidemic trends. *Int. J. Infect. Dis. IJID Off. Publ. Int. Soc. Infect. Dis.* 100: 164–173. <https://doi.org/10.1016/j.ijid.2020.08.066>

672 Barrett J. C., 2009 Haploview: Visualization and analysis of SNP genotype data. *Cold Spring Harb. Protoc.* 2009:
673 *pdb.ip71*. <https://doi.org/10.1101/pdb.ip71>

674 Barton N. H., and B. Charlesworth, 1998 Why sex and recombination? *Science* 281: 1986–1990.

675 Benvenuto D., M. Giovanetti, M. Salemi, M. Prosperi, C. De Flora, *et al.*, 2020 The global spread of 2019-nCoV:
676 a molecular evolutionary analysis. *Pathog. Glob. Health* 114: 64–67.
677 <https://doi.org/10.1080/20477724.2020.1725339>

678 Bergh B. V. den, T. Swings, M. Fauvert, and J. Michiels, 2018 Experimental Design, Population Dynamics, and
679 Diversity in Microbial Experimental Evolution. *Microbiol. Mol. Biol. Rev.* 82.
680 <https://doi.org/10.1128/MMBR.00008-18>

681 Bobay L.-M., C. C. Traverse, and H. Ochman, 2015 Impermanence of bacterial clones. *Proc. Natl. Acad. Sci.*
682 112: 8893–8900. <https://doi.org/10.1073/pnas.1501724112>

683 Brown T., X. Didelot, D. J. Wilson, and N. D. Maio, 2016 SimBac: simulation of whole bacterial genomes with
684 homologous recombination. *Microb. Genomics* 2. <https://doi.org/10.1099/mgen.0.000044>

685 Burton D. R., and E. J. Topol, 2021 Variant-proof vaccines — invest now for the next pandemic. *Nature* 590:
686 386–388. <https://doi.org/10.1038/d41586-021-00340-4>

687 Cagliani R., D. Forni, M. Clerici, and M. Sironi, 2020 Computational inference of selection underlying the evolution
688 of the novel coronavirus, SARS-CoV-2. *J. Virol.* <https://doi.org/10.1128/JVI.00411-20>

689 Campos J. L., and B. Charlesworth, 2019 The Effects on Neutral Variability of Recurrent Selective Sweeps and
690 Background Selection. *Genetics* 212: 287–303. <https://doi.org/10.1534/genetics.119.301951>

691 Carvajal-Rodríguez A., 2008 GENOMEPOP: a program to simulate genomes in populations. *BMC Bioinformatics*
692 9: 223. <https://doi.org/10.1186/1471-2105-9-223>

693 Charlesworth J., and A. Eyre-Walker, 2006 The Rate of Adaptive Evolution in Enteric Bacteria. *Mol. Biol. Evol.*
694 23: 1348–1356. <https://doi.org/10.1093/molbev/msk025>

695 Charlesworth J., and A. Eyre-Walker, 2008 The McDonald-Kreitman test and slightly deleterious mutations. *Mol.*
696 *Biol. Evol.* 25: 1007–1015. <https://doi.org/10.1093/molbev/msn005>

697 Charlesworth B., 2013 Background selection 20 years on: the Wilhelmine E. Key 2012 invitational lecture. *J.*
698 *Hered.* 104: 161–171. <https://doi.org/10.1093/jhered/ess136>

699 Chinese SARS Molecular Epidemiology Consortium, 2004 Molecular evolution of the SARS coronavirus during
700 the course of the SARS epidemic in China. *Science* 303: 1666–1669.
701 <https://doi.org/10.1126/science.1092002>

702 Choi B., M. C. Choudhary, J. Regan, J. A. Sparks, R. F. Padera, *et al.*, 2020 Persistence and Evolution of SARS-
703 CoV-2 in an Immunocompromised Host. *N. Engl. J. Med.* 383: 2291–2293.
704 <https://doi.org/10.1056/NEJMc2031364>

705 Clement M., D. Posada, and K. A. Crandall, 2000 TCS: a computer program to estimate gene genealogies. *Mol.*
706 *Ecol.* 9: 1657–1659. <https://doi.org/10.1046/j.1365-294x.2000.01020.x>

707 Cobey S., 2020 Modeling infectious disease dynamics. *Science* 368: 713–714.
708 <https://doi.org/10.1126/science.abb5659>

709 Cohen A. A., P. N. P. Gnanapragasam, Y. E. Lee, P. R. Hoffman, S. Ou, *et al.*, 2021 Mosaic nanoparticles elicit
710 cross-reactive immune responses to zoonotic coronaviruses in mice. *Science* 371: 735–741.
711 <https://doi.org/10.1126/science.abf6840>

712 Collier D. A., A. De Marco, I. A. T. M. Ferreira, B. Meng, R. P. Datir, *et al.*, 2021 Sensitivity of SARS-CoV-2
713 B.1.1.7 to mRNA vaccine-elicited antibodies. *Nature*. <https://doi.org/10.1038/s41586-021-03412-7>

714 Cui J., F. Li, and Z.-L. Shi, 2019 Origin and evolution of pathogenic coronaviruses. *Nat. Rev. Microbiol.* 17: 181–
715 192. <https://doi.org/10.1038/s41579-018-0118-9>

716 Cvijović I., A. N. Nguyen Ba, and M. M. Desai, 2018 Experimental Studies of Evolutionary Dynamics in Mi-
717 crobes. *Trends Genet. TIG* 34: 693–703. <https://doi.org/10.1016/j.tig.2018.06.004>

718 Danecek P., A. Auton, G. Abecasis, C. A. Albers, E. Banks, *et al.*, 2011 The variant call format and VCFtools.
719 *Bioinforma. Oxf. Engl.* 27: 2156–2158. <https://doi.org/10.1093/bioinformatics/btr330>

720 DeLong E. F., 2004 Microbial population genomics and ecology: the road ahead. *Environ. Microbiol.* 6: 875–878.
721 <https://doi.org/10.1111/j.1462-2920.2004.00668.x>

722 Denison M. R., R. L. Graham, E. F. Donaldson, L. D. Eckerle, and R. S. Baric, 2011 Coronaviruses. *RNA Biol.* 8:
723 270–279. <https://doi.org/10.4161/rna.8.2.15013>

724 Desai M. M., and D. S. Fisher, 2007 Beneficial mutation selection balance and the effect of linkage on positive
725 selection. *Genetics* 176: 1759–1798. <https://doi.org/10.1534/genetics.106.067678>

726 Di L., P. E. Pagan, D. Packer, C. L. Martin, S. Akther, *et al.*, 2014 BorreliaBase: a phylogeny-centered browser of
727 *Borrelia* genomes. *BMC Bioinformatics* 15: 233. <https://doi.org/10.1186/1471-2105-15-233>

728 Didelot X., D. Lawson, and D. Falush, 2009 SimMLST: simulation of multi-locus sequence typing data under a
729 neutral model. *Bioinforma. Oxf. Engl.* 25: 1442–1444. <https://doi.org/10.1093/bioinformatics/btp145>

730 Felsenstein J., 1989 PHYLIP - Phylogeny Inference Package. *Cladistics* 5: 164–166.

731 Fraser C., W. P. Hanage, and B. G. Spratt, 2007 Recombination and the nature of bacterial speciation. *Science*
732 315: 476–480. <https://doi.org/10.1126/science.1127573>

733 Galloway S. E., 2021 Emergence of SARS-CoV-2 B.1.1.7 Lineage — United States, December 29, 2020–January
734 12, 2021. *MMWR Morb. Mortal. Wkly. Rep.* 70. <https://doi.org/10.15585/mmwr.mm7003e2>

735 Garud N. R., P. W. Messer, E. O. Buzbas, and D. A. Petrov, 2015 Recent selective sweeps in North American
736 *Drosophila melanogaster* show signatures of soft sweeps. *PLoS Genet.* 11: e1005004.
737 <https://doi.org/10.1371/journal.pgen.1005004>

738 Gillespie J. H., 2000 Genetic Drift in an Infinite Population: The Pseudohitchhiking Model. *Genetics* 155: 909–
739 919.

740 Good B. H., M. J. McDonald, J. E. Barrick, R. E. Lenski, and M. M. Desai, 2017 The dynamics of molecular evo-
741 lution over 60,000 generations. *Nature* 551: 45–50. <https://doi.org/10.1038/nature24287>

742 Hadfield J., C. Megill, S. M. Bell, J. Huddleston, B. Potter, *et al.*, 2018 Nextstrain: real-time tracking of pathogen
743 evolution. *Bioinformatics* 34: 4121–4123. <https://doi.org/10.1093/bioinformatics/bty407>

744 Haller B. C., and P. W. Messer, 2019 SLiM 3: Forward Genetic Simulations Beyond the Wright–Fisher Model.
745 *Mol. Biol. Evol.* 36: 632–637. <https://doi.org/10.1093/molbev/msy228>

746 Haven J., L. C. Vargas, E. F. Mongodin, V. Xue, Y. Hernandez, *et al.*, 2011 Pervasive Recombination and Sym-
747 patric Genome Diversification Driven by Frequency-Dependent Selection in *Borrelia burgdorferi*, the
748 Lyme Disease Bacterium. *Genetics* 189: 951–966. <https://doi.org/10.1534/genetics.111.130773>

749 Hernandez R. D., 2008 A flexible forward simulator for populations subject to selection and demography. *Bioin-
750 forma. Oxf. Engl.* 24: 2786–2787. <https://doi.org/10.1093/bioinformatics/btn522>

751 Hernández Y., R. Bernstein, P. Pagan, L. Vargas, W. McCaig, *et al.*, 2018 BpWrapper: BioPerl-based sequence

752 and tree utilities for rapid prototyping of bioinformatics pipelines. *BMC Bioinformatics* 19: 76.
753 <https://doi.org/10.1186/s12859-018-2074-9>

754 Hill W. G., and A. Robertson, 1966 The effect of linkage on limits to artificial selection. *Genet. Res.* 8: 269–294.

755 Huang Y., C. Yang, X.-F. Xu, W. Xu, and S.-W. Liu, 2020 Structural and functional properties of SARS-CoV-2
756 spike protein: potential antivirus drug development for COVID-19. *Acta Pharmacol. Sin.* 41: 1141–1149.
757 <https://doi.org/10.1038/s41401-020-0485-4>

758 Hudson R. R., M. Kreitman, and M. Aguadé, 1987 A test of neutral molecular evolution based on nucleotide data.
759 *Genetics* 116: 153–159.

760 Hudson R. R., and N. L. Kaplan, 1995 Deleterious background selection with recombination. *Genetics* 141:
761 1605–1617.

762 Hudson R. R., 2002 Generating samples under a Wright-Fisher neutral model of genetic variation. *Bioinforma.*
763 Oxf. Engl. 18: 337–338.

764 Kelleher J., A. M. Etheridge, and G. McVean, 2016 Efficient Coalescent Simulation and Genealogical Analysis
765 for Large Sample Sizes. *PLoS Comput. Biol.* 12: e1004842. <https://doi.org/10.1371/journal.pcbi.1004842>

766 Kemp S. A., D. A. Collier, R. P. Datir, I. A. T. M. Ferreira, S. Gayed, *et al.*, 2021 SARS-CoV-2 evolution during
767 treatment of chronic infection. *Nature* 592: 277–282. <https://doi.org/10.1038/s41586-021-03291-y>

768 Kim Y., and W. Stephan, 2000 Joint effects of genetic hitchhiking and background selection on neutral variation.
769 *Genetics* 155: 1415–1427.

770 Kissler S. M., C. Tedijanto, E. Goldstein, Y. H. Grad, and M. Lipsitch, 2020 Projecting the transmission dynamics
771 of SARS-CoV-2 through the postpandemic period. *Science*. <https://doi.org/10.1126/science.abb5793>

772 Korber B., W. M. Fischer, S. Gnanakaran, H. Yoon, J. Theiler, *et al.*, 2020 Tracking Changes in SARS-CoV-2
773 Spike: Evidence that D614G Increases Infectivity of the COVID-19 Virus. *Cell* 182: 812-827.e19.
774 <https://doi.org/10.1016/j.cell.2020.06.043>

775 Koropoulis A., N. Alachiotis, and P. Pavlidis, 2020 Detecting Positive Selection in Populations Using Genetic
776 Data. *Methods Mol. Biol.* Clifton NJ 2090: 87–123. https://doi.org/10.1007/978-1-0716-0199-0_5

777 Koyama T., D. Weeraratne, J. L. Snowdon, and L. Parida, 2020 Emergence of Drift Variants That May Affect
778 COVID-19 Vaccine Development and Antibody Treatment. *Pathog. Basel Switz.* 9.
779 <https://doi.org/10.3390/pathogens9050324>

780 Lam T. T.-Y., M. H.-H. Shum, H.-C. Zhu, Y.-G. Tong, X.-B. Ni, *et al.*, 2020 Identifying SARS-CoV-2 related
781 coronaviruses in Malayan pangolins. *Nature* 1–6. <https://doi.org/10.1038/s41586-020-2169-0>

782 Lang G. I., D. P. Rice, M. J. Hickman, E. Sodergren, G. M. Weinstock, *et al.*, 2013 Pervasive genetic hitchhiking
783 and clonal interference in forty evolving yeast populations. *Nature* 500: 571–574.
784 <https://doi.org/10.1038/nature12344>

785 Lenski R. E., 2017 Experimental evolution and the dynamics of adaptation and genome evolution in microbial
786 populations. *ISME J.* 11: 2181–2194. <https://doi.org/10.1038/ismej.2017.69>

787 Liang L., S. Zöllner, and G. R. Abecasis, 2007 GENOME: a rapid coalescent-based whole genome simulator. *Bi-
788 oinforma.* Oxf. Engl. 23: 1565–1567. <https://doi.org/10.1093/bioinformatics/btm138>

789 Lintusaari J., M. U. Gutmann, R. Dutta, S. Kaski, and J. Corander, 2017 Fundamentals and Recent Developments
790 in Approximate Bayesian Computation. *Syst. Biol.* 66: e66–e82. <https://doi.org/10.1093/sysbio/syw077>

791 Lu W., B.-J. Zheng, K. Xu, W. Schwarz, L. Du, *et al.*, 2006 Severe acute respiratory syndrome-associated coro-
792 navirus 3a protein forms an ion channel and modulates virus release. *Proc. Natl. Acad. Sci. U. S. A.* 103: 12540–12545. <https://doi.org/10.1073/pnas.0605402103>

794 Lu R., X. Zhao, J. Li, P. Niu, B. Yang, *et al.*, 2020 Genomic characterisation and epidemiology of 2019 novel
795 coronavirus: implications for virus origins and receptor binding. *Lancet Lond. Engl.* 395: 565–574.
796 [https://doi.org/10.1016/S0140-6736\(20\)30251-8](https://doi.org/10.1016/S0140-6736(20)30251-8)

797 Luk H. K. H., X. Li, J. Fung, S. K. P. Lau, and P. C. Y. Woo, 2019 Molecular epidemiology, evolution and phy-
798 logeny of SARS coronavirus. *Infect. Genet. Evol. J. Mol. Epidemiol. Evol. Genet. Infect. Dis.* 71: 21–30.
799 <https://doi.org/10.1016/j.meegid.2019.03.001>

800 Maddamsetti R., R. E. Lenski, and J. E. Barrick, 2015 Adaptation, Clonal Interference, and Frequency-Dependent
801 Interactions in a Long-Term Evolution Experiment with *Escherichia coli*. *Genetics* 200: 619–631.
802 <https://doi.org/10.1534/genetics.115.176677>

803 Marçais G., A. L. Delcher, A. M. Phillippy, R. Coston, S. L. Salzberg, *et al.*, 2018 MUMmer4: A fast and versa-
804 tile genome alignment system. *PLoS Comput. Biol.* 14: e1005944.
805 <https://doi.org/10.1371/journal.pcbi.1005944>

806 Masters P. S., 2006 The molecular biology of coronaviruses. *Adv. Virus Res.* 66: 193–292.
807 [https://doi.org/10.1016/S0065-3527\(06\)66005-3](https://doi.org/10.1016/S0065-3527(06)66005-3)

808 McDonald J. H., and M. Kreitman, 1991 Adaptive protein evolution at the *Adh* locus in *Drosophila*. *Nature* 351:
809 652–654. <https://doi.org/10.1038/351652a0>

810 Messer P. W., and D. A. Petrov, 2013 Frequent adaptation and the McDonald-Kreitman test. *Proc. Natl. Acad. Sci. U. S. A.* 110: 8615–8620. <https://doi.org/10.1073/pnas.1220835110>

812 Milkman R., and M. M. Bridges, 1990 Molecular evolution of the *Escherichia coli* chromosome. III. Clonal
813 frames. *Genetics* 126: 505–517.

814 Muller H. J., 1964 The relation of recombination to muational advance. *Mutat. Res.* 106: 2–9.

815 Múrias dos Santos A., M. P. Cabezas, A. I. Tavares, R. Xavier, and M. Branco, 2016 tcsBU: a tool to extend TCS
816 network layout and visualization. *Bioinformatics* 32: 627–628.
817 <https://doi.org/10.1093/bioinformatics/btv636>

818 Papkou A., T. Guzella, W. Yang, S. Koepper, B. Pees, *et al.*, 2019 The genomic basis of Red Queen dynamics
819 during rapid reciprocal host-pathogen coevolution. *Proc. Natl. Acad. Sci. U. S. A.* 116: 923–928.
820 <https://doi.org/10.1073/pnas.1810402116>

821 Peeri N. C., N. Shrestha, M. S. Rahman, R. Zaki, Z. Tan, *et al.*, 2020 The SARS, MERS and novel coronavirus
822 (COVID-19) epidemics, the newest and biggest global health threats: what lessons have we learned? *Int. J. Epidemiol.* <https://doi.org/10.1093/ije/dyaa033>

824 Phan T., 2020 Genetic diversity and evolution of SARS-CoV-2. *Infect. Genet. Evol. J. Mol. Epidemiol. Evol. Genet. Infect. Dis.* 81: 104260. <https://doi.org/10.1016/j.meegid.2020.104260>

826 Plante J. A., Y. Liu, J. Liu, H. Xia, B. A. Johnson, *et al.*, 2021 Spike mutation D614G alters SARS-CoV-2 fitness.
827 *Nature* 592: 116–121. <https://doi.org/10.1038/s41586-020-2895-3>

828 Pupko T., R. E. Bell, I. Mayrose, F. Glaser, and N. Ben-Tal, 2002 Rate4Site: an algorithmic tool for the identifi-
829 cation of functional regions in proteins by surface mapping of evolutionary determinants within their
830 homologues. *Bioinformatics* 18: S71–S77. https://doi.org/10.1093/bioinformatics/18.suppl_1.S71

831 Qiu W.-G., S. E. Schutzer, J. F. Bruno, O. Attie, Y. Xu, *et al.*, 2004 Genetic exchange and plasmid transfers in
832 *Borrelia burgdorferi* sensu stricto revealed by three-way genome comparisons and multilocus sequence
833 typing. *Proc. Natl. Acad. Sci. U. S. A.* 101: 14150–14155. <https://doi.org/10.1073/pnas.0402745101>

834 Rambaut A., E. C. Holmes, Á. O'Toole, V. Hill, J. T. McCrone, *et al.*, 2020 A dynamic nomenclature proposal
835 for SARS-CoV-2 lineages to assist genomic epidemiology. *Nat. Microbiol.* 5: 1403–1407.
836 <https://doi.org/10.1038/s41564-020-0770-5>

837 Rocha E. P. C., 2018 Neutral Theory, Microbial Practice: Challenges in Bacterial Population Genetics. *Mol. Biol.*
838 *Evol.* 35: 1338–1347. <https://doi.org/10.1093/molbev/msy078>

839 Sabeti P. C., D. E. Reich, J. M. Higgins, H. Z. P. Levine, D. J. Richter, *et al.*, 2002 Detecting recent positive se-
840 lection in the human genome from haplotype structure. *Nature* 419: 832–837.
841 <https://doi.org/10.1038/nature01140>

842 Shu Y., and J. McCauley, 2017 GISAID: Global initiative on sharing all influenza data - from vision to reality.
843 *Euro Surveill. Bull. Eur. Sur Mal. Transm. Eur. Commun. Dis. Bull.* 22. <https://doi.org/10.2807/1560-7917.ES.2017.22.13.30494>

845 Silva J. da, and J. D. Galbraith, 2017 Hill-Robertson interference maintained by Red Queen dynamics favours the
846 evolution of sex. *J. Evol. Biol.* 30: 994–1010. <https://doi.org/10.1111/jeb.13068>

847 Siu K.-L., K.-S. Yuen, C. Castaño-Rodriguez, Z.-W. Ye, M.-L. Yeung, *et al.*, 2019 Severe acute respiratory syn-
848 drome coronavirus ORF3a protein activates the NLRP3 inflammasome by promoting TRAF3-dependent
849 ubiquitination of ASC. *FASEB J.* 33: 8865–8877. <https://doi.org/10.1096/fj.201802418R>

850 Smith J. M., N. H. Smith, M. O'Rourke, and B. G. Spratt, 1993 How clonal are bacteria? *Proc. Natl. Acad. Sci. U.*
851 *S. A.* 90: 4384–4388.

852 Stajich J. E., D. Block, K. Boulez, S. E. Brenner, S. A. Chervitz, *et al.*, 2002 The Bioperl toolkit: Perl modules for
853 the life sciences. *Genome Res.* 12: 1611–1618. <https://doi.org/10.1101/gr.361602>

854 Tenaillon O., J. E. Barrick, N. Ribeck, D. E. Deatherage, J. L. Blanchard, *et al.*, 2016 Tempo and mode of ge-
855 nome evolution in a 50,000-generation experiment. *Nature* 536: 165–170.
856 <https://doi.org/10.1038/nature18959>

857 To K. K.-W., S. Sridhar, K. H.-Y. Chiu, D. L.-L. Hung, X. Li, *et al.*, 2021 Lessons learned 1 year after SARS-
858 CoV-2 emergence leading to COVID-19 pandemic. *Emerg. Microbes Infect.* 10: 507–535.
859 <https://doi.org/10.1080/22221751.2021.1898291>

860 Turakhia Y., N. D. Maio, B. Thornlow, L. Gozashti, R. Lanfear, *et al.*, 2020 Stability of SARS-CoV-2 phyloge-
861 nies. *PLOS Genet.* 16: e1009175. <https://doi.org/10.1371/journal.pgen.1009175>

862 Volz E., V. Hill, J. T. McCrone, A. Price, D. Jorgensen, *et al.*, 2021 Evaluating the Effects of SARS-CoV-2 Spike
863 Mutation D614G on Transmissibility and Pathogenicity. *Cell* 184: 64–75.e11.
864 <https://doi.org/10.1016/j.cell.2020.11.020>

865 Wang Q., Y. Zhang, L. Wu, S. Niu, C. Song, *et al.*, 2020 Structural and Functional Basis of SARS-CoV-2 Entry
866 by Using Human ACE2. *Cell* 181: 894–904.e9. <https://doi.org/10.1016/j.cell.2020.03.045>

867 Wang P., M. S. Nair, L. Liu, S. Iketani, Y. Luo, *et al.*, 2021 Antibody resistance of SARS-CoV-2 variants B.1.351
868 and B.1.1.7. *Nature*. <https://doi.org/10.1038/s41586-021-03398-2>

869 Wit E. de, N. van Doremalen, D. Falzarano, and V. J. Munster, 2016 SARS and MERS: recent insights into
870 emerging coronaviruses. *Nat. Rev. Microbiol.* 14: 523–534. <https://doi.org/10.1038/nrmicro.2016.81>

871 Yang Z., 2007 PAML 4: phylogenetic analysis by maximum likelihood. Mol. Biol. Evol. 24: 1586–1591.
872 <https://doi.org/10.1093/molbev/msm088>

873 Yang H.-C., C. Chen, J.-H. Wang, H.-C. Liao, C.-T. Yang, *et al.*, 2020 Analysis of genomic distributions of
874 SARS-CoV-2 reveals a dominant strain type with strong allelic associations. Proc. Natl. Acad. Sci. 117:
875 30679–30686. <https://doi.org/10.1073/pnas.2007840117>

876 Yi H., 2020 2019 Novel Coronavirus Is Undergoing Active Recombination. Clin. Infect. Dis.
877 <https://doi.org/10.1093/cid/ciaa219>

878 Yuan X., D. J. Miller, J. Zhang, D. Herrington, and Y. Wang, 2012 An overview of population genetic data simu-
879 lation. J. Comput. Biol. J. Comput. Mol. Cell Biol. 19: 42–54. <https://doi.org/10.1089/cmb.2010.0188>

880 Zhou P., X.-L. Yang, X.-G. Wang, B. Hu, L. Zhang, *et al.*, 2020 A pneumonia outbreak associated with a new
881 coronavirus of probable bat origin. Nature 579: 270–273. <https://doi.org/10.1038/s41586-020-2012-7>

882 Zinzula L., 2021 Lost in deletion: The enigmatic ORF8 protein of SARS-CoV-2. Biochem. Biophys. Res. Com-
883 mun. 538: 116–124. <https://doi.org/10.1016/j.bbrc.2020.10.045>

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Tables and Figures

887 **Table 1. Protein-coding loci (n=25) included in simulated SARS-CoV-2 genome evolution**

| Protein ID ^a | Locus symbol | Protein function ^b | Location ^a | Locus length | Syn sites (S) ^c | Missense sites (N) ^c |
|-------------------------|--------------|---|----------------------------|--------------|----------------------------|---------------------------------|
| YP_009725297.1 | <i>nsp1</i> | Leader protein; inhibits host translation | 266, 805 | 540 | 164.48 | 375.52 |
| YP_009725298.1 | <i>nsp2</i> | Unknown | 806, 2719 | 1914 | 583.94 | 1330.06 |
| YP_009725299.1 | <i>nsp3</i> | Polyprotein processing | 2720, 8554 | 5835 | 1738.36 | 4096.64 |
| YP_009725300.1 | <i>nsp4</i> | Formation of double membrane vesicles associated with replication complexes | 8555, 10054 | 1500 | 449.50 | 1050.50 |
| YP_009725301.1 | <i>nsp5</i> | 3C-like proteinase; polyprotein processing | 10055, 10972 | 918 | 278.19 | 639.81 |
| YP_009725302.1 | <i>nsp6</i> | Formation of double membrane vesicles associated with replication complexes | 10973, 11842 | 870 | 253.34 | 616.66 |
| YP_009725303.1 | <i>nsp7</i> | Accessory subunit of RNA-dependent RNA polymerase | 11843, 12091 | 249 | 75.97 | 173.03 |
| YP_009725304.1 | <i>nsp8</i> | Accessory subunit of RNA-dependent RNA polymerase; primase | 12092, 12685 | 594 | 164.77 | 429.23 |
| YP_009725305.1 | <i>nsp9</i> | RNA-binding protein | 12686, 13024 | 339 | 105.61 | 233.39 |
| YP_009725306.1 | <i>nsp10</i> | Co-factor of Nsp14 and Nsp16 for methyltransferase activity | 13025, 13441 | 417 | 123.76 | 293.24 |
| YP_009725307.1 | <i>nsp12</i> | RNA-dependent RNA polymerase | 13442, 13468; 13468, 16236 | 2796 | 843.60 | 1952.40 |
| YP_009725308.1 | <i>nsp13</i> | Helicase | 16237, 18039 | 1803 | 541.37 | 1261.63 |
| YP_009725309.1 | <i>nsp14</i> | Proof-reading 3'-to-5' exonuclease | 18040, 19620 | 1581 | 469.80 | 1111.20 |
| YP_009725310.1 | <i>nsp15</i> | Endonuclease | 19621, 20658 | 1038 | 319.98 | 718.02 |
| YP_009725311.1 | <i>nsp16</i> | Ribose 2'-O-methyltransferase; RNA cap formation | 20659, 21552 | 894 | 268.86 | 625.14 |
| YP_009724390.1 | <i>S</i> | Surface glycoprotein; spike protein; binding of host cell receptor | 21563, 25384 | 3822 | 1150.89 | 2671.11 |
| | <i>S1</i> | S1 subunit containing the angiotensin- | 21563, 23185 | 1623 | 482.32 | 1140.68 |

| | | | | | | |
|----------------|--------------|--|--------------|------|--------|---------|
| | | converting enzyme 2 (ACE2) receptor-binding domain (RBD) (Wang <i>et al.</i> 2020; Huang <i>et al.</i> 2020) | | | | |
| YP_009724391.1 | <i>S2</i> | S2 subunit responsible for membrane fusion | 23186, 25381 | 2296 | 667.38 | 1528.62 |
| | <i>orf3a</i> | Ion channel promoting virus release (Lu <i>et al.</i> 2006; Siu <i>et al.</i> 2019) | 25393, 26220 | 828 | 252.54 | 575.46 |
| YP_009724392.1 | <i>E</i> | Envelope protein forming homopentameric cation channel | 26245, 26472 | 228 | 80.43 | 147.57 |
| YP_009724393.1 | <i>M</i> | Membrane glycoprotein | 26523, 27191 | 669 | 202.40 | 466.60 |
| YP_009724394.1 | <i>orf6</i> | Unknown | 27202, 27387 | 186 | 53.74 | 132.26 |
| YP_009724395.1 | <i>orf7a</i> | Unknown | 27394, 27759 | 366 | 115.71 | 250.29 |
| YP_009725318.1 | <i>orf7b</i> | Unknown | 27756, 27887 | 132 | 46.24 | 85.76 |
| YP_009724396.1 | <i>orf8</i> | Ion channel contributing to suppression of host immunity (Zinzula 2021) | 27894, 28259 | 366 | 121.59 | 244.41 |
| YP_009724397.2 | <i>N</i> | Nucleocapsid phosphoprotein; viral RNA genome protection and packaging | 28274, 29533 | 1260 | 396.51 | 863.49 |
| YP_009725255.1 | <i>orf10</i> | Unknown; suspected membrane protein forming viroporin | 29558, 29674 | 117 | 32.88 | 84.12 |

888 ^a Start and stop positions (stop codon included) based on the genome of the strain Wuhan-Hu-1 (GenBank Accession NC_045512)
889 (Zhou *et al.* 2020). The *nsp11* locus (39 bases; 13442-13480), which is small and enclosed within the *nsp12* locus, was excluded. The
890 4-base overlap between *orf7a* and *orf7b* was counted towards the former locus.

891 ^b (To *et al.* 2021)

892 ^c Effective numbers of synonymous (S) and nonsynonymous (N) substitution sites of a protein-coding locus, derived with the *CovSim-*
893 *ulator* (see Material and Methods)

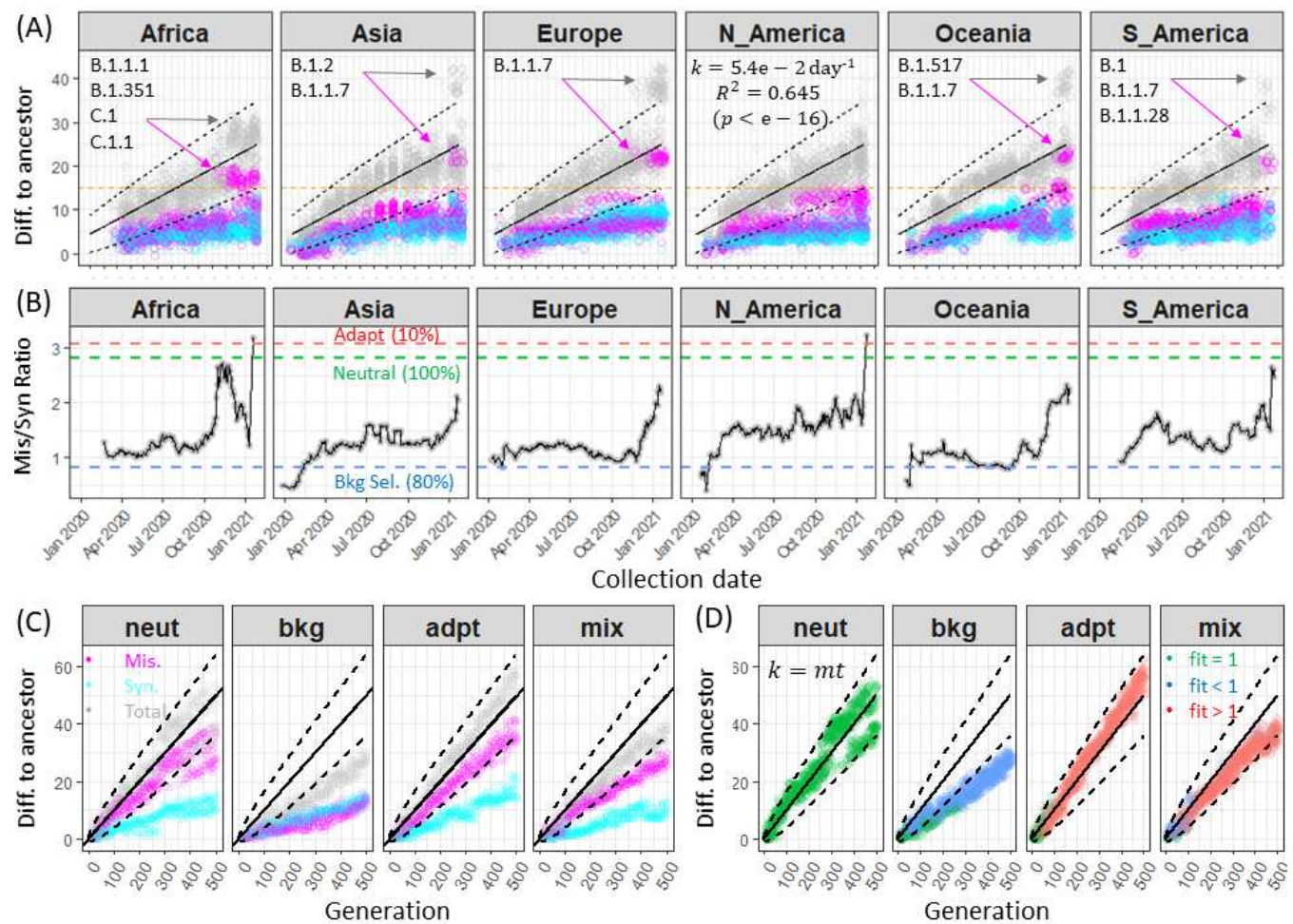
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896 **Table 2. Evolutionary parameters and neutral expectations**

| Symbol | Description and Settings |
|----------------------|--|
| N_e | Effective population size ($N_e = 200$); held constant |
| g | Total number of generations ($g = 500$) |
| L | Genome length ($L = 29,903$ bases, GenBank Accession NC_045512) |
| μ, μ_0 | Mutation rate per site; neutral mutation rate |
| $\theta = 2N_e\mu_0$ | Expected level of neutral sequence diversity at mutation-drift balance |
| $TMRC$ | Time to the most recent common ancestor; $\sim 2N_e$ for a haplotype population |
| π, π_s, π_n | Average pair-wise total, synonymous, nonsynonymous sequence differences |
| m | Mutation rate per genome ($m = 0.1$); Poisson distributed and uniform across the genome |
| $k=mt$ | Expected sequence difference with respect to the ancestral sequence over generation time (t); Poisson distributed with variance equal to mean, or standard deviation $sd(k) = \sqrt{mt}$ |
| r | Recombination rate per genome ($r = 0$ for clonal evolution without recombination); Poisson distributed with uniform breakpoint probabilities across the genome |
| s | Sample size per generation ($s = 20$, 10% of population size) |
| w_1, w_2 | Multiplicative fitness loss for a deleterious ($w_1 = 0.95$) missense mutation or fitness gain for an adaptive ($w_2 = 1.05$) missense mutation; relative to the progenitor genome |
| u, v | Probabilities of a missense mutation being deleterious ($u = 0.5$) or adaptive ($v = 0.05$); relative to the progenitor genome |
| Q | Empirically derived base substitution probabilities: $Q = \begin{pmatrix} 0 & 0.1083 & 0.7000 & 0.1917 \\ 0.0475 & 0 & 0.0033 & 0.9492 \\ 0.2102 & 0.0931 & 0 & 0.6967 \\ 0.1025 & 0.795 & 0.1025 & 0 \end{pmatrix}$. Both rows (source bases) and columns (destination) are in the order of A, C, G, and T, probabilities for each source base (in a row) summing to 1. |
| S | Total number of synonymous sites in a protein-coding locus |
| N | Total number of nonsynonymous sites in a protein-coding locus |
| d_s, d_n | Levels of synonymous and nonsynonymous divergence between viral species: $d_s = D_s/S$; $d_n = D_n/N$, where D_s and D_n are the numbers of synonymous and nonsynonymous base differences |
| p_s, p_n | Levels of synonymous and nonsynonymous polymorphism in a viral population: $p_s = P_s/S$; $p_n = P_n/N$, where P_s and P_n are the numbers of synonymous and nonsynonymous polymorphic sites |

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899 **Fig 1. Rates of synonymous and nonsynonymous divergence of SARS-CoV-2 genomes**

900 (A) In each panel, points represent differences with respect to the reference genome (y-axis) of
901 viral genomes originating in a continent with collection dates (x-axis) ranging from Dec 2019
902 through March 2021. A random sample of 100 genomes was chosen for each month, resulting
903 in ~1400 genomes for each continent with evenly distributed monthly representations. Each
904 genome was represented three times, including the number of synonymous mutations (cyan),
905 the number of missense mutations (magenta), and the total number of genetic changes (in-
906 cluding indels; gray). A linear regression line (solid, with statistics shown within the
907 "N_America" panel) was derived by using genomes from all continents. Dashed lines show two
908 standard deviations above and below the regression line on the basis of the Poisson expected
909 variance $\sigma^2(k) = k$. Hyper-mutated genomes (marked by the lineage designations) that
910 emerged in late 2020 showed accelerated accumulation of missense (but not synonymous)
911 mutations (Choi *et al.* 2020; Kemp *et al.* 2021). The orange lines indicate a cutoff value of 15
912 missense mutations to determine outliers.

913 (B) Ratios of the numbers of missense (D_n) to synonymous (D_s) mutations relative to the refer-
914 ence genome (y-axis), a measure of selective constraint, were plotted against the viral sample
915 collection dates (x-axis). Each point was the ratio of the average number of missense to syn-
916 onymous mutations within a moving window of 14 days. Horizontal dashed lines mark the rati-
917 os obtained from simulated evolution and percentages represent proportions of missense mu-
918 tations that were deleterious (blue), neutral (green) and adaptive (red) (see below). D_n/D_s rati-
919 os in all populations started at low levels, indicating strong purifying selection during the early
920 months (before April 2020) of the pandemic. The D_n/D_s ratio increased greatly in later months
921 of 2020 worldwide, suggesting rapid population expansion and the emergence of human-
922 adaptive viral variants.

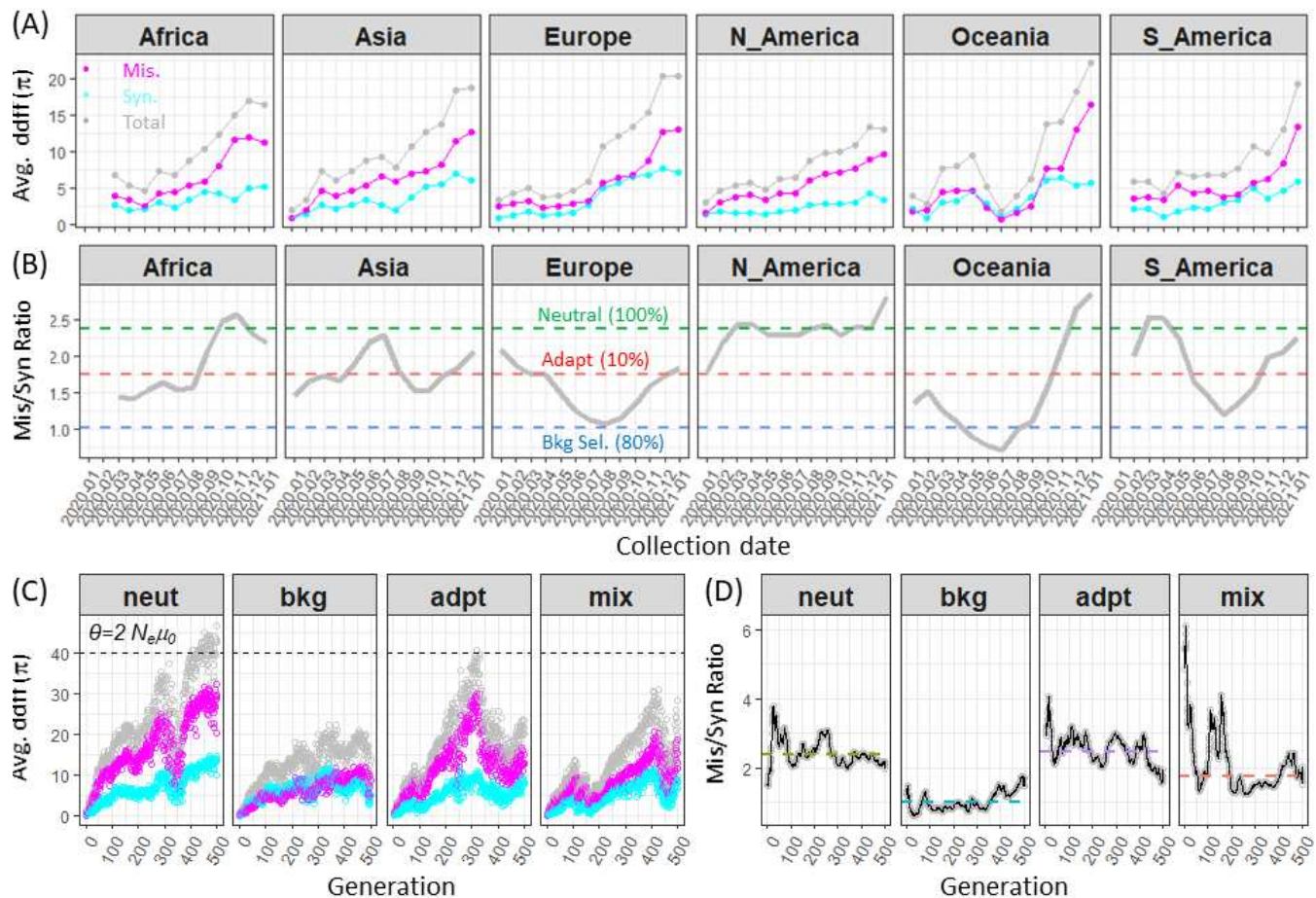
923 (C) Mutational divergence (y-axis) over the generation (x-axis) of genomes simulated with four
924 evolution models. For each model, a sample of $s=20$ genomes was chosen for each genera-
925 tion, resulting in $\sim 10,000$ genomes for each model. Solid lines indicate the expected mutation
926 rate in the neutral model. Dashed lines show two standard deviations above and below the ex-
927 pected total mutation rate on the basis of the Poisson expected variance $\sigma^2(k) = mt$. Genome
928 evolution was simulated with a population size $N=200$, genome mutation rate $m=0.1$ and no
929 recombination ($r=0$). In the neutral evolution model (“NEUT”), all missense mutations carried a
930 fitness of 1. In the background selection model (“BKG”), a missense mutation had an 80%
931 chance of incurring a fitness cost of 0.95 and was otherwise neutral. In the adaptive evolution
932 model (“ADPT”), a missense mutation had a 10% chance of incurring a fitness benefit of 1.05
933 and was otherwise neutral. In the mixed evolution model (“MIX”), a missense mutation had an
934 80% chance of incurring a fitness cost of 0.95, a 10% chance of incurring a fit benefit of 1.05,
935 and 10% chance of being neutral. As expected, the ratio of missense to synonymous muta-
936 tions (~ 1.0) in the BKG model was markedly lower than that in neutral evolution and was used
937 as a control (blue dashed line in *panel B*) for measuring viral evolution during the pandemic.
938 The ratios of missense to synonymous mutations from the neutral and mixed evolution models
939 were much higher (~ 3.0) and were used as another set of controls (green and red dashed lines
940 in *panel B*) to understand SARS-CoV genome evolution.

941 (D) Simulated genomes colored by fitness values. In the neutral evolution model, the total mu-
942 tuation rate and its variability accurately followed the expectations, thereby validating the
943 *CovSimulator*. In the background selection model, the overall mutation rate decreased and the
944 population was increasingly dominated by low-fitness genomes, showing a gradual loss of fit-
945 ness in a clonal population known as Muller’s ratchet (Muller 1964). In the adaptive evolution

946 model, mutation accumulation was accelerated and the population was dominated by adaptive
947 lineages except in the first 100 generations. In the mixed evolution model, adaptive lineages
948 dominated the population despite the presence of strong purifying selection.

949

950



951

952 Fig 2. Synonymous and nonsynonymous polymorphisms of SARS-CoV-2 populations

953 The average number of pairwise sequence differences (π) is a measure of viral genetic diversity,
 954 which reflects the viral effective population size and viral reproduction rate.

955 (A) Each panel represents a continental population. Nonsynonymous (π_n), nonsynonymous (π_n),
 956 and total pairwise sequence differences (y-axis) were calculated from monthly samples from
 957 December 2020 through March 2021 (x-axis). Sequence diversity increased in all populations.
 958 An initial increase in genetic diversity was expected for a nascent viral population before
 959 reaching mutation-drift balance. However, the acceleration of viral diversity in later months was
 960 unexpected and reflected the accumulation of neutral, deleterious, and adaptive genetic diversity
 961 in rapidly expanding viral populations.

962 (B) The ratio of nonsynonymous (π_n) to synonymous (π_s) diversity, similarly to D_n/D_s (Fig 1), is
 963 a measure of selective constraints. The π_n/π_s ratios were elevated and fluctuated substantially,
 964 in agreement with a lack of selective constraints in rapidly expanding viral populations.

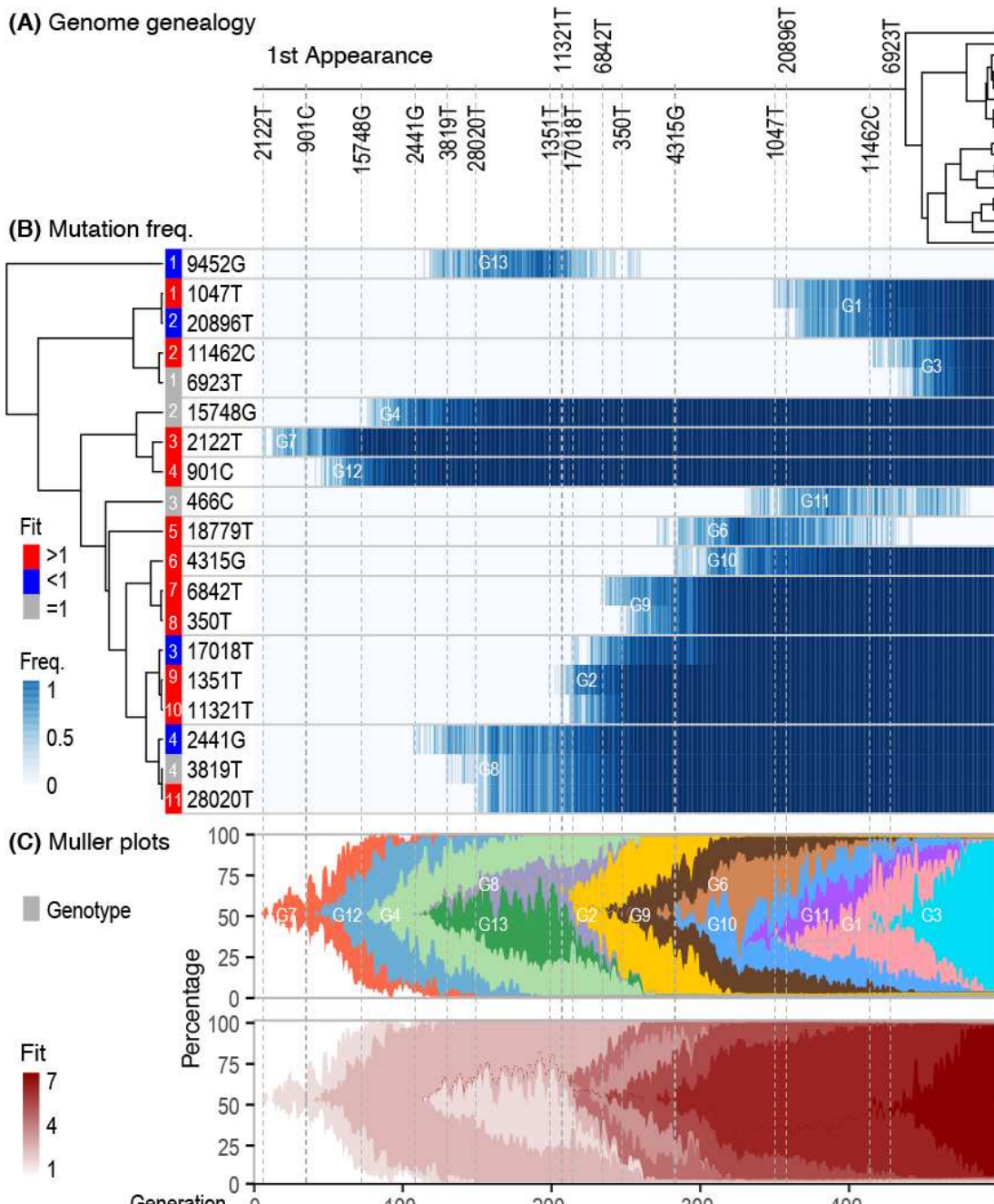
965 (C) The π values of simulated genomes under four evolution models. In the neutral evolution

966 model the total π value stabilized at the expected value of $\theta = 40$, further validating the
967 *CovSimulator*. The π values were relatively lower in the background and adaptive selection
968 models, in agreement with smaller effective population sizes due to natural selection and
969 shorter coalescent times (Supplemental Fig S4). Genome evolution was simulated with a
970 population size $N=200$, genome mutation rate $m=0.1$ and without recombination ($r=0$). In the
971 neutral evolution model (“NEUT”), all missense mutations carried a fitness of one. In the back-
972 ground selection model (“BKG”), a missense mutation had an 80% chance of carrying a fitness
973 cost of 0.95. In the adaptive evolution model (“ADPT”), a missense mutation had a 10%
974 chance of carrying a fitness benefit of 1.05.

975 (D) The π_n/π_s ratios for the four evolution models, showing a low value for a population under
976 purifying selection, a high value during neutral evolution, and intermediate values for a popula-
977 tion under adaptive evolution. These average ratios are shown in panel (B) as references for
978 comparison with values based on viral samples.

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982 **Fig 3. Mixed evolution as a model of SARS-CoV-2 genome evolution**

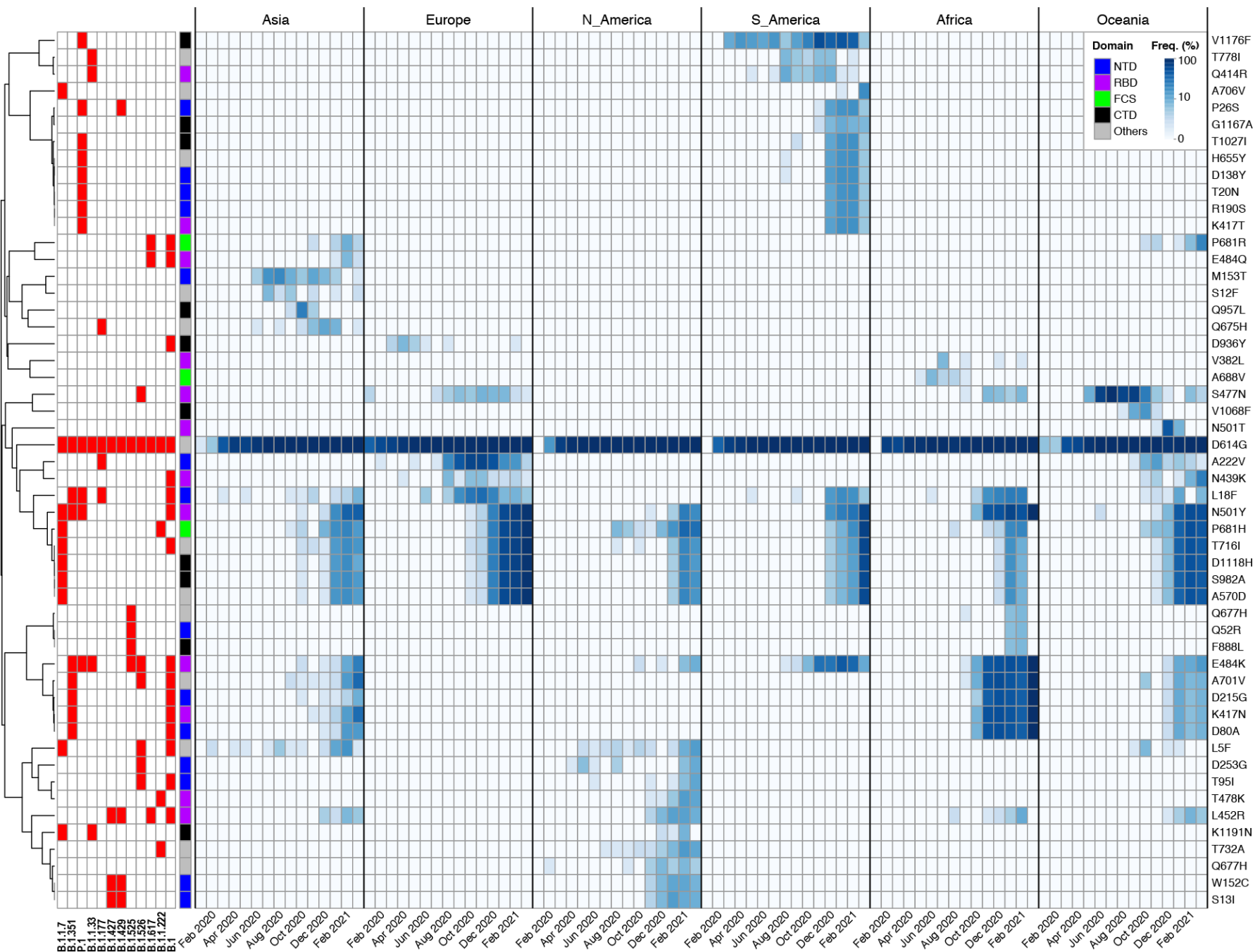
983 Simulated evolution of a viral population subject to both purifying and adaptive selection (pa-
984 rameters defined in Fig 1) captured viral adaptation at the level of individual mutations.

985 (A) Genealogy of the $n=20$ simulated genomes sampled from the last generation. The 16 mu-
986 tations were fixed in the final population and their times of first appearance in the population
987 (tick marks on the root edge) corresponded to the timing of selective sweeps shown in the Mul-
988 ler diagram (panel C). The genealogy shows that the final viral population descended from the
989 latest selective sweep, which occurred only \sim 50 generations ago.

990 (B) In the heatmap, rows show $n=19$ high frequency ($\geq 5\%$ in 10,000 genome samples) non-

991 synonymous SNVs, labeled by the genome position and destination base, from the mixed evo-
992 lution model. Frequencies were represented by color intensity (*middle panel*). Unlike those of
993 natural viral variants, the fitness values of simulated variants were precisely known (*colored*
994 *side bar*). The variants were grouped according to the correlated evolutionary trajectories
995 (*dendrogram*) such that mutations with similar trajectories (“genotypes”) – indicating temporal
996 linkage – were adjacent. Adaptive mutations (red, numbered from #1 through #11) dominated
997 the population. Adaptive mutations (#3 and #4) arose early. One adaptive mutation (#5) was
998 lost, probably because of clonal interference but also possibly because of genetic drift. Among
999 the four deleterious mutations, one (#1) was lost whereas the other three hitchhiked with linked
1000 adaptive mutations to fixation. Similarly, one neutral mutation (#3) was lost and three others
1001 hitchhiked to fixation. Furthermore, the simulation suggests a high rate of multiple mutations
1002 occurring at the same genomic site associated with adaptive mutations (#1, #10, and #11).
1003 Three of the four multiple-hit mutations had T as the destination base, reflecting the strong mu-
1004 tation bias in which ~70% SNVs during viral evolution were due to C>T or G>T substitutions.
1005 (C) Muller diagrams of mutation trajectories in the simulated population. In the top diagram, at
1006 each vertical cross section, the heights of colored blocks represent the relative frequencies of
1007 the “genotypes”. A genotype represented one or more mutations displaying a similar evolu-
1008 tionary trajectory reflecting temporal linkage. The Muller diagram reveals selective sweeps oc-
1009 ccurring regularly and strong competition among adaptive mutations. For example, during gen-
1010 erations 100 to 250, the top Muller plot shows competition between two genotypes (G8 and
1011 G13), both of which carried a deleterious mutation. They had similar fitness values (bottom
1012 Muller plot). At the ~250 generation, however, G13 was outcompeted by G8 and went extinct.
1013 G8 itself subsequently gave rise to G2. Similarly, at the ~300th generation, two adaptive muta-
1014 tions (#5 in G6 and #6 in G10) began to compete against each other and coexisted until the
1015 ~450th generation when G6 was eventually displaced by G1 and G3, both descendants of G10.
1016 The latter case represents a loss of an adaptive allele (#5) due to clonal interference.

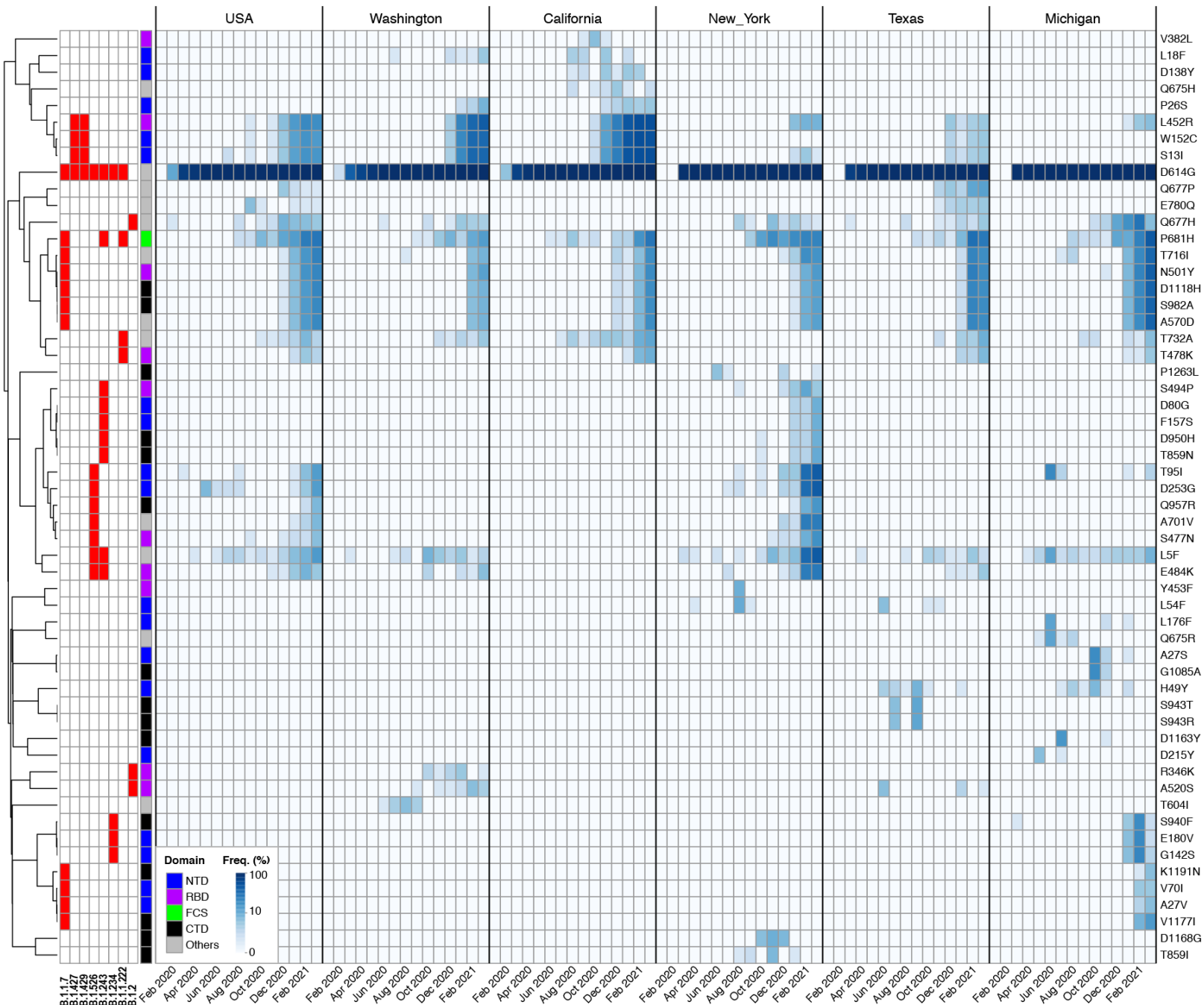
1017



019 **Fig 4. Frequency trajectories of high-frequency missense SNVs in six continental SARS-CoV-2 populations**

020 The heatmap depicts allele frequencies (*colored cells*, in percentage, scaled with 10-based logarithm) by month (*columns*) of 51 mis-
021 sense mutations (*rows*) on the spike protein in viral populations from six continents (*vertical blocks*). Each mutation frequency was cal-
022 culated on the basis of ~100 genomes randomly sampled from a month within a continent. Each mutation was present with a ≥5% allele
023 frequency in at least one monthly sample. Mutations were grouped according to similarities in frequency trajectories (*rowside dendro-
024 gram*). The rowside table shows mutations associated with major viral lineages (Rambaut *et al.* 2020) (columns #1 through #12). Col-
025 umn #13 of the rowside table shows the spike protein domains associated with the mutations, including the N-terminus domain (NTD),
026 receptor-binding domain (RBD), Furin cleavage site (FCS); and the C-terminus domain (CTD). The heatmap reveals the early rise and
027 rapid fixation of the D614G mutation across the globe (dark blue stripe in the middle). Also discernable is rapid global spread of six
028 spike protein mutations (N501Y, P681H, T716I, D1118H, S982A, and A570D) associated with the hyper-mutated B.1.1.7 lineage
029 (rowside column #1) after its first emergence during October 2020 in Europe (Choi *et al.* 2020). Other mutations were associated with
030 lineages that have so far shown limited geographic ranges, including the B.1.351 (originated in Africa, rowside column #2), P.1 (South
031 America, rowside column #3), B.1.427 and B.1.429 (North American, rowside columns #6 and #7), and B.1.617 (Asia, rowside column
032 #10) lineages. For early detection of human-adaptive mutations, it is necessary to track mutation frequencies at country and regional
033 levels before they become more widespread (Fig 5).

034



036 [Fig 5. Tracking emergent adaptive mutations at regional levels](#)

037 The heatmap depicts monthly (*columns*) frequencies (*colored cells*) of 56 missense mutations (*rows*) on the spike protein that were
038 present with $\geq 5\%$ allele frequencies in at least one monthly SARS-CoV-2 sample from the United States and its five states (*vertical*
039 *blocks*). As in the global heatmap (Fig 4), the D614G mutation reached fixation across the country since March 2020. The European
040 lineage B.1.1.7 (*rowside column #1*) first arrived the US in December 2020 and quickly spread to all five states. The B.1.427 and
041 B.1.429 lineages (*rowside columns #2 and #3*) were first identified in fall 2020 in California and have spread to the four other states by
042 the end of March 2021. Similarly, the B.1.1.222 (*rowside column #7*) lineage was first identified in summer in California and has since
043 spread to Washington, Texas, and Michigan. The B.1.526 and B.1.243 (*rowside columns #4 and #5*) lineages emerged during Decem-
044 ber 2020 in New York and have not yet spread to the other four states. Similarly, two other lineages have thus far not yet spread out-
045 side of the state of origination, including the B.1.2 lineage (*rowside column #8*) in Washington and the B.1.234 lineage (*rowside col-*
046 *umns #6*) in Michigan. None of the latter four regional lineages has reached a $\geq 5\%$ frequency at the national level (1st *vertical block*),
047 highlighting the importance of identifying human-adaptive mutations by tracking mutation frequencies at the level of local populations.