Potential transmission chains of variant B.1.1.7 and co-mutations of SARS-CoV-2

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25 Abstract

The presence of SARS-CoV-2 mutants, including the emerging variant B.1.1.7, has raised 26 great concerns in terms of pathogenesis, transmission, and immune escape. Characterizing 27 SARS-CoV-2 mutations, evolution, and effects on infectivity and pathogenicity is crucial to 28 the design of antibody therapies and surveillance strategies. Here we analyzed 454,443 29 SARS-CoV-2 spike genes/proteins and 14,427 whole-genome sequences. We 30 demonstrated that the early variant B.1.1.7 may not have evolved spontaneously in the 31 United Kingdom or within human populations. Our extensive analyses suggested that 32 Canidae, Mustelidae or Felidae, especially the Canidae family (for example, dog) could be a 33 possible host of the direct progenitor of variant B.1.1.7. An alternative hypothesis is that the 34 variant was simply yet to be sampled. Notably, the SARS-CoV-2 whole genome represents a 35 large number of potential co-mutations with very strong statistical significances (p value<E-36 44). In addition, we used an experimental SARS-CoV-2 reporter replicon system to introduce 37 the dominant co-mutations NSP12 c14408t, 5'UTR c241t, and NSP3 c3037t into the viral 38 genome, and to monitor the effect of the mutations on viral replication. Our experimental 39 results demonstrated that the co-mutations significantly attenuated the viral replication. The 40 study provides valuable clues for discovering the transmission chains of variant B.1.1.7 and 41 understanding the evolutionary process of SARS-CoV-2. 42

43 **Key words:** SARS-CoV-2, variant B.1.1.7, transmission chains, co-mutations, viral 44 replication.

45 Introduction

Since the outbreak in December 2019, COVID-19 has been pandemic in over 200 46 countries. Cases of infection and mortalities have been surging and are an ongoing threat to 47 public health^{1,2}. COVID-19 is caused by infection with the novel coronavirus SARS-CoV-2³⁻⁵. 48 Although as a coronavirus, SARS-CoV-2 has genetic proofreading mechanisms⁶⁻⁸, the 49 persistent natural selection pressure in the population drives the virus to gradually 50 accumulate favorable mutations^{6,9,10}. Much attention has been paid to the mutations and 51 evolution of SARS-CoV-2¹¹⁻¹⁵, since mutations are related to the infectivity and pathogenicity 52 of viruses¹⁶⁻²¹. Beneficial mutants of the virus can better evolve and adapt to the host⁹, either 53 strengthening or weakening the infectivity and pathogenicity. In addition, certain variants 54 may generate drug resistance and reduce the efficacy of vaccines and therapeutics²²⁻²⁶. In 55 short, studying mutations and evolution in detail is vital to understand the transformations of 56 viral properties and to control the pandemic. 57

A new variant of SARS-CoV-2 named VOC-202012/01 (Variant of Concern 202012/01) or lineage B.1.1.7 was first detected in the United Kingdom last December²⁷. It appears to be substantially more transmissible than other variants²⁸. The variant has been growing exponentially in the United Kingdom and rapidly spreading to other countries^{29,30}. However, it is not yet clear if it evolved spontaneously in the United Kingdom or was imported from other countries. Studying how the variant B.1.1.7 mutates can enable researchers to track its spread over time and to understand the evolution of SARS-CoV-2.

In this study, large-scale SARS-CoV-2 sequences, consisting of more than 454,000 spike

genes/proteins and 14,000 whole-genome sequences were analyzed. Our extensive 66 sequence analysis showed that many mutations always co-occur not only in the spike 67 protein of B.1.1.7, but in the whole genome of SARS-CoV-2. The mutation trajectories of 68 the spike protein indicate that the early variant B.1.1.7 did not evolve spontaneously in the 69 United Kingdom or even within human populations. We also investigated possible 70 SARS-CoV-2 transmission chains of the variant B.1.1.7 based on the mutation analysis of 71 large-scale spike proteins and the cluster analysis of spike genes. Over the whole genome, 72 the top 25 high-frequency mutations of SARS-CoV-2 converged into several potential 73 co-mutation patterns, each of which showed a strong correlation with a very strong 74 statistical significance (p value<E-44). The potential co-mutations depicted the 75 evolutionary trajectory of SARS-CoV-2 virus in the population, shaping variable replication 76 77 of SARS-CoV-2. In addition, we further explored the effect of the dominant (co-)mutations 5'UTR c241t, NSP3 c3037t, and NSP12 c14408t on viral replication using a 78 SARS-CoV-2 replicon based on a four plasmid *in-vitro* ligation system. The results 79 suggest that such mutations significantly attenuate the replication of SARS-CoV-2. 80

81 **Results**

82 Evolutionary trajectories of variant B.1.1.7

The variant B.1.1.7 was generally defined by multiple amino acid changes including 3 deletions (69-70del and 145del) and 7 mutations (N501Y, A570D, D614G, P681H, T716I, S982A, and D1118H) in the spike protein³¹. The number of non-adjacent co-occurrent changes indicates that they resulted from accumulated mutations. We therefore explored the

evolutionary trajectories of B.1.1.7 by tracing the incremental mutations (Fig. 1a). All routes 87 along the directions of the arrows are possible evolutionary trajectories of lineage B.1.1.7. 88 Among all the mutation routes, the green one was the most probable mutation trajectory 89 based on the number of variant strains. However, it was unlikely that the earliest variant 90 B.1.1.7 (GISAID: EPI_ISL_601443, 2020-09-20, England) with 9 mutations evolved from the 91 existing variants with 3–8 mutations, because the former arose much earlier than the latter. 92 More than 454,000 SARS-CoV-2 strains have been collected and extensive sequenced from 93 infected humans without finding intermediate variants with 3-9 mutations. It is therefore 94 95 unlikely that the intermediate variants with 3-8 mutations have infected humans. Thus, the early variant B.1.1.7 might not have arisen spontaneously in the UK or within human 96 populations. An alternative hypothesis is that spillover likely occurred from susceptible 97 animals. 98

The co-appearance rates (see Materials and Methods) of all nine mutations are shown in 99 Fig. 1b. We found that at least five mutations (145del, A570D, T716I, S982A, and D1118H) 100 of variant B.1.1.7 significantly co-occurred (rate>95%), which indicates a potential 101 co-mutation pattern in the spike protein, causing us to wonder what selection pressure drove 102 such co-occurrences of mutations and rapid evolution in the population of SARS-CoV-2. 103 Note that coronaviruses generally tend to exhibit rapid evolution when they jump to a 104 different species³². We therefore analyzed the key spike genes and proteins of existing 105 SARS-CoV-2 strains collected from animals to find a possible direct progenitor of variant 106 B.1.1.7. The variant with mutations "56" (labeled by "*" in Fig. 1a, termed star variant) had 107 the minimum phylogenetic distance with EPI_ISL_699508, which was collected from a dog 108

on 2020-07-28 (Fig. 2) using MEGA^{33,34} (see Materials and Methods). The strains collected 109 from tigers, minks, and cats were also close to the star variant. Our extensive analyses 110 including mutations, phylogeny (Fig. 2), collection date/location and the number of 111 sequences (Tables S1-S3) suggested that Canidae, Mustelidae or Felidae, especially the 112 Canidae family (for example, dog) could be a possible host of the direct progenitor of variant 113 B.1.1.7. The possible transmission chains of variant B.1.1.7 are shown in Fig. 1c. This star 114 variant strains in humans could not have evolved into the early variant B.1.1.7, but they 115 might have infected high-density yet susceptible animals (such as dogs) and adapted to 116 these species through rapid mutation. Such progenitor variants comprised most or all of the 117 mutations of the early variant B.1.1.7 within the Canidae family populations, and they may 118 have spilled back to humans after the rapid mutation period. 119

120 High-frequency mutations converge into potential co-mutations

Based on sequence alignment and mutation analysis, we found that 7,441 nucleotide 121 alterations in the viral 29903-letter RNA code occurred at least once in the samples from 122 COVID-19 patients. These mutations were dispersed in the 14,427 SARS-CoV-2 strains 123 collected from all around the world. As shown in the heatmap of the top 1% high-frequency 124 mutations (Table S4), some sites show very similar mutation rates on most days in samples 125 isolated globally (Fig. S1), including 8,898 and 815 samples isolated from the U.S. (Fig. S2) 126 and Australia (Fig. S3). Therefore, these mutations shown in Fig. S4a were selected and 127 clustered into co-occurrences, which we called potential co-mutation patterns. From the 128 landscape of the mutation rates (Fig. S4a), 25 nucleotide sites were clearly clustered into 129

several potential co-mutation patterns. Among these patterns, there was one consisting of 130 the top 4 high-frequency mutations (i.e., 5'UTR c241t, NSP3 c3037t, NSP12 c14408t, and 131 S_a23403g), which converged into a dominant potential co-mutation pattern. Such 132 co-occurrence lineage has been found in almost all sequenced samples of SARS-CoV-2. 133 Within this co-occurrence pattern, mutation S_ a23403g resulted in the amino acid change 134 (D614G) that apparently enhances viral infectivity^{6,35}, albeit debate exists¹⁶. Notably, there 135 were three successive sites at the 28881st to 28883rd positions of the virus (N g28881a, 136 N g28882a, and N g28883c) that strictly co-occurred. Comparing Fig. S4a-c and Table S4, 137 138 we found that the top 14 high-frequency mutations formed five common co-occurrence patterns. 139

To assess the above co-occurrence patterns, we analyzed the correlations and statistical 140 significance levels of the high-frequency co-occurrence mutations. The heatmap of the 141 paired Pearson-correlation-coefficients (Fig. 3a) shows that the top 25 high-frequency 142 mutations clearly cluster into several potential co-mutation groups/patterns with very strong 143 correlation (≥0.8). By regression analyses, the above co-occurrence patterns have statistical 144 significance levels with p values less than 10^{-44} (Fig. 3b). The detailed mutation transitions 145 (Fig. 3c-k, Figs. S5-7) provide further evidence that the above mutations form co-mutation 146 patterns. 147

Dominant mutations attenuate viral replication

We further explored the effect of the dominant mutations 5'UTR_c241t, NSP3_c3037t, and NSP12_c14408t on viral replication using a SARS-CoV-2 replicon based on a

four-plasmid *in-vitro* ligation system. This replicon is devoid of the viral structural proteins 151 while undergoing viral replication, and the viral replication is sensitive to the antiviral agent 152 remdesivir³⁶. The 5'UTR_c241t mutation resides in a highly conserved region in the 5'UTR 153 (Fig. 4a). The NSP3 c3037t mutation is synonymous. The NSP12 c14408t mutation is 154 nonsynonymous with an amino acid change of a conserved amino acid P323 in the viral 155 RNA-dependent RNA polymerase (Fig. 4b). We introduced the NSP12_c14408t mutation or 156 the NSP12_c14408t mutation with the other two mutations 5'UTR_c241t and NSP3_c3037t 157 into the replicon plasmids. The fragments were released from the plasmids by Bsal digestion, 158 and then assembled by *in-vitro* ligation with T4 ligase (Fig. 4c). Replicon RNA transcribed 159 from the ligation products was co-transfected with N mRNA into Huh7 cells. RNA replication 160 was monitored by measuring the secreted Gaussia luciferase activity in the supernatants. 161 162 Enzymatic dead mutants (759-SAA-761) of the RNA-dependent RNA polymerase NSP12 were introduced, and the mutated replicon served as a non-replication control. As shown in 163 Fig. 4d, transfection of WT replicon RNA resulted in an obvious increase of luciferase activity, 164 and SAA RNA did not replicate as expected. Introduction of NSP12_c14408t mutation 165 resulted in a significant reduction of viral replication. The combination of NSP12 c14408t 166 mutation with the other two mutations further significantly but only marginally reduced viral 167 replication. These results demonstrate that the P323L mutation in the viral RNA-dependent 168 RNA polymerase reduces viral replication, and the synonymous mutations may further 169 attenuate viral replication. 170

171 Discussion

A well-resolved phylogeny of variant B.1.1.7 spike genes provides an opportunity to 172 understand the evolutionary process and transmission chains of variant B.1.1.7. Our 173 incremental mutation and phylogenetic analyses on large-scale SARS-CoV-2 spike 174 proteins/genes revealed that the early variant B.1.1.7 might not have evolved spontaneously 175 in the United Kingdom or within human populations. In this case the spillover likely occurred 176 from susceptible animals. Current evidence³⁷⁻³⁹ indicates that SARS-CoV-2 can effectively 177 infect both domestic animals (for example, dog, cat, pig and bovine) and wild animals (for 178 example, mink, rabbit and fox) by binding their angiotensin converting enzyme 2 (ACE2). 179 Our further analyses including mutations, phylogeny, collection date/location and the 180 number of sequences suggested that the earliest variant B.1.1.7 possibly originated from 181 182 Canidae, Mustelidae or Felidae, especially the Canidae family (for example, dog). The cases⁴⁰ that the variant B.1.1.7 can easily infect dogs and cats indicated that both are 183 susceptible to B.1.1.7. Still, due to the limited information available to date, an alternative 184 hypothesis is that the direct progenitor of variant B.1.1.7 is yet to be sampled. In addition to 185 variant B.1.1.7, as a future topic we will work on the analysis of other lineages such as P.1, 186 B.1.351, B.1.427, and B.1.42, when sufficient numbers of their sequences are available. 187

By tracing the mutation trajectories, we found that at least five mutations of the spike proteins always co-occurred, and a large number of potential co-mutations appeared in the top 1% high-frequency mutations of SARS-CoV-2 whole genome. It has been documented that the mutation S_ a23403g results in the amino acid change of the spike protein D614G

and enhances viral infectivity^{19,41-44}. Here, by using a SARS-CoV-2 reporter replicon system, 192 we demonstrated that the one of the dominant co-mutations NSP12 c14408t significantly 193 reduced viral replication and combination of NSP12_c14408t mutation with the other two 194 synonymous mutations 5'UTR c241t and NSP3 c3037t although significantly but only 195 marginally reduced viral replication further. As the 5'UTR play an important role in regulating 196 viral replication, the synonymous mutations 5'UTR c241t may attenuate viral replication by 197 change RNA secondary structure⁴⁵. These findings imply that SARS-CoV-2 undergoes an 198 evolution toward enhancing viral infectivity while attenuating viral replication. SARS-CoV-2 199 200 has exhibited significant mutations and co-mutations. We evaluated the replication of a co-mutation pattern including three dominant mutations. If other mutations act similarly on 201 the viral replication needs to be verified. These results can be further explored for efficient 202 203 vaccine design in our future work. In summary, this study provides insights into the transmission chains of variant B.1.1.7 and the effect of viral dominant mutations on viral 204 evolution. 205

206 Materials and Methods

207 Data selection and pre-processing

The 454,443 spike gene/protein sequences of SARS-CoV-2 were obtained at https://www.gisaid.org/. The NCBI website at https://www.ncbi.nlm.nih.gov/sars-cov-2/ has released more than 1.7 thousand sequences of SARS-CoV-2 viruses before July 31, 2020. We selected 14,427 sequences that satisfied two criteria: (1) having specific collection dates; (2) sequence-lengths being no less than 29,305 nt (29903*0.98). It is inevitable that some

sites of sequences are equivocal owing to the limitation of sequencing depth. For instance, many sites were labeled as letter N in genome sequences. The noise of indeterminate nucleic-acids was taken into consideration in our experiments so as to boost accuracy. The co-mutation rate of multi-site co-mutations was calculated by co – mutation rate = $\frac{number of sequences containing co-mutations}{number of all sequences}$. Moreover, the co-appearance rate of a mutation in B.1.1.7 variant was defined by co – appearance rate = $\frac{number of B.1.1.7 sequences}{number of sequences containing a mutation}$.

219 **Possible animal host analyses**

220 In addition to the phylogenetic analysis, we further explored the possible animal hosts of the direct progenitor of variant B.1.1.7 by mutations, collection time/space of strains, the 221 number of sequences and the edit distance^{46,47} of mutations (Table S1-2). Due the late 222 223 lockdown policies of some governmental agencies, the spread of SARS-CoV-2 has not been prevented well in Europe, America, and Australia. We could ignore the impact of policies for 224 studying the origin of variant B.1.1.7. We quantified the multiple impact factors of viral 225 226 transmission as shown in Table S3 based on the criterion that the smaller the value, the more similar. The results still supported that the Canidae family is a possible host of the 227 direct progenitor of variant B.1.1.7. 228

229 MEGA version and parameter settings

- 230 Version: MEGA-X
- 231 Statistical Method: Maximum Likelihood
- 232 Test of Phylogeny: None
- 233 Model/Method: Jones-Taylor-Thornton (JTT) model

- 234 Rates among Sites: Uniform Rates
- 235 Gaps/Missing Data Treatment: Use all sites
- 236 ML Heuristic Method: Nearest-Neighbor-Interchange (NNT)
- Initial Tree for ML: Make initial tree automatically (Default NJ/BioNJ)
- 238 Branch Swap Filter: None
- Number of Threads: 7

240 Statistical analysis

The Pearson-correlation-coefficient (PCC) is a classic statistic that measures linear correlation between two variables. Its value ranges from -1.0 to 1.0. Normally, the two variables meet a strong correlation or a very strong correlation when the absolutes value of PCC is between 0.6 and 0.8 or between 0.8 and 1.0. Linear regression is a linear approach to model the relationship between a scalar response and one or more variables. We used PCC and significance level (p value) of regression analysis to evaluate the relationships of the co-occurrence mutations in large-scale SARS-CoV-2 examples.

248 Plasmids

Four plasmids encompassing the viral genome (pLC-nCoV-A-Bsal, pLC-nCoV-B-Bsal, 249 previously³⁶. pnCoV-D-sGluc-Bsal) pLC-nCoV-C-Bsal, and were described The 250 5'UTR c-241-t and NSP3 c-3037-t mutations were introduced into the pLC-nCoV-A-Bsal by 251 fusion PCR. The NSP12_c-14408-t mutation was introduced into the pLC-nCoV-B-Bsal by 252 fusion PCR. 253

254 Cell lines

The human hepatoma cells Huh 7 were purchased from the Cell Bank of the Chinese Academy of Sciences (www.cellbank.org.cn) and routinely maintained in Dulbecco's modified medium supplemented with 10% FBS (Gibco) and 25 mM HEPES (Gibco).

258 *In-vitro* ligation

Bsal digested fragments were gel purified using Gel Extraction Kit (OMEGA) and ligated with T4 ligase (New England Biolabs) at room temperature for 1 h. The ligation products were phenol/chloroform extracted, precipitated by absolute ethanol, and resuspended in nuclease-free water, quantified by determining the A260 absorbance.

263 *In-vitro* transcription

Purified *in-vitro* ligated product was used as template for the *in-vitro* transcription by 264 mMESSAGE mMACHINE T7 Transcription Kit (Ambion) according to the manufacturer's 265 protocol. For N mRNA production, we amplified the N coding region by PCR (sense: GGC 266 267 TTT TTT TCT AGG CCT GAG TTG AGT CAG CAC) with phCMV-N as template. Then the 268 purified PCR product was used as a template for *in-vitro* transcription by mMESSAGE 269 mMACHINE T7 Transcription Kit as described above. RNA was purified by RNeasy mini 270 Elute (Qiagen), eluted in nuclease-free water, and quantified by UV absorbance (260 nm). 271

272 Transfection

273 Cells were seeded onto 48-well plates at a density of 7.5×10⁴ per well and then

transfected with 0.3 µg *in-vitro* transcribed RNA using a TransIT-mRNA transfection kit
 (Mirus) according to the manufacturer's protocol.

276 Luciferase activity

277 Supernatants were taken from cell medium and mixed with equal volumes of 2×lysis 278 buffer (Promega). Luciferase activity was measured with Renilla luciferase substrate 279 (Promega) according to the manufacturer's protocol.

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291 Author contributions

L.N.C. and J.S.Z. designed the study. Z.G.Y. and J.S.Z. designed the experiments. J.S.Z.

analyzed data. Y. Z. performed the experiments of viral replication. J.S.Z., Z.G.Y., and J.Y.K.
designed the figures. S.C. repeated and checked the experiments of viral replication. H.B.H.
checked the computational analyses. J.S.Z. and Z.G.Y. wrote the manuscript. Y.Q.H, M.F.L.,
L.N.L, and L.L. polished the manuscript. All authors participated in result interpretation and
discussion.

298 **Data availability**

- 299 The raw sequence data reported in this paper have been deposited in the GISAID and NCBI
- 300 websites at https://www.gisaid.org/ and https://www.ncbi.nlm.nih.gov/sars-cov-2/,
- respectively. Code is available from the corresponding author on reasonable request.

302 **Conflict of interest**

The authors declare that they have no conflict of interest.

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401 Figures and Figure legends

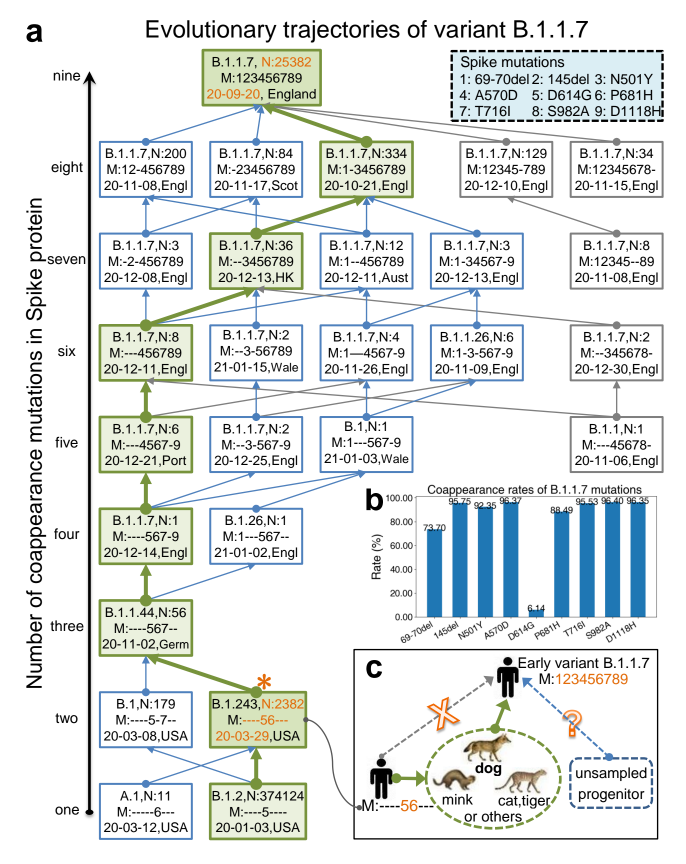
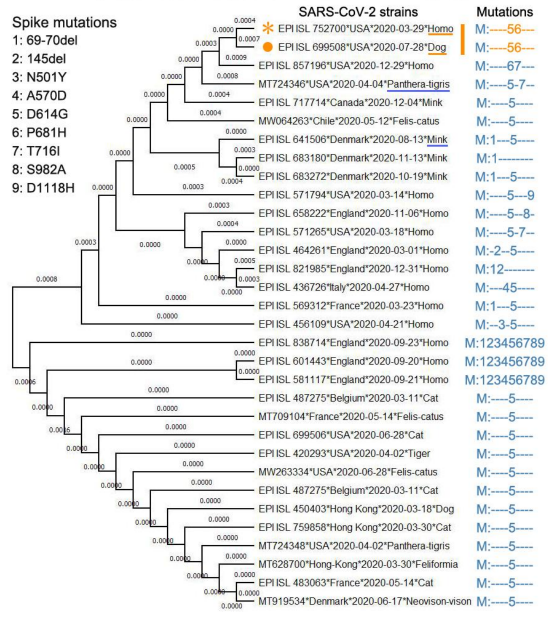
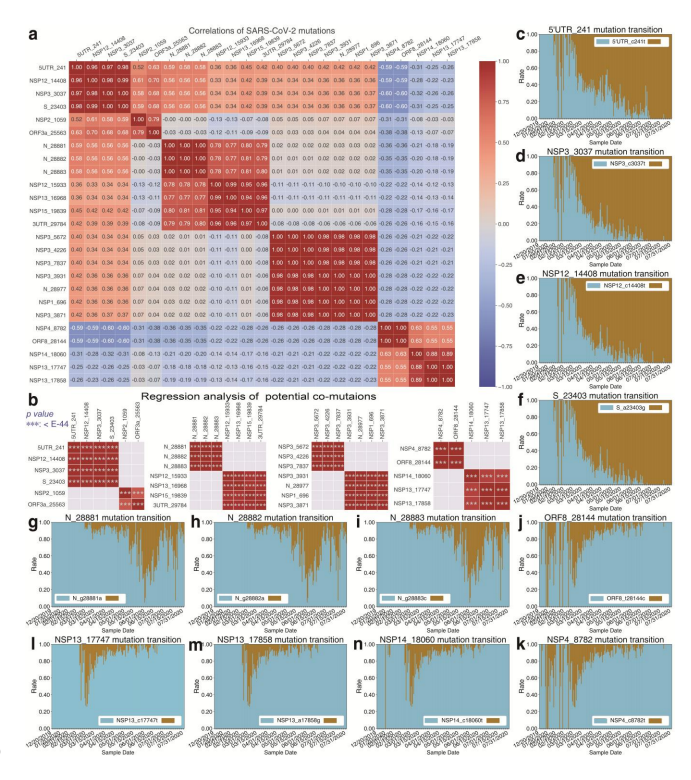


Fig. 1 Evolutionary trajectories of variant B.1.1.7. a Incremental mutations of variant B.1.1.7. The 403 404 digits in the upper-right-corner rectangle with dotted line indicate the labels of mutations. For simplicity, the 69–70 deletions were labeled as "1", and the other mutations "2"-"9" respectively. The 405 bottom nodes (rectangles) represent the variants with one mutation and the top one was the early 406 variant B.1.1.7. Each rectangle with solid line consists of lineage (e.g., B.1.243), number of strains 407 (e.g., N:2382), mutation sites (e.g., M:---56---), the earliest collection date (e.g., 20-03-29, i.e., 408 2020-03-29), and collection location (e.g., USA). In the labels of the mutation sites, sign "-" indicated 409 410 the corresponding site did not mutate. All routes along the directions of the arrows are possible evolutionary trajectories of lineage B.1.1.7, where the green one was the most probable mutation 411 trajectory. Large-scale SARS-CoV-2 analysis demonstrates that the early variant B.1.1.7 might not 412 have arisen spontaneously in the UK or within human hosts. b Coappearances of variant B.1.1.7 413 mutations. At least five mutations form a potential co-mutation pattern (coappearance rate > 95%). c 414 Possible transmission chains of variant B.1.1.7. Canidae, Mustelidae or Felidae, especially the 415 Canidae family (for example, dog) could be a possible host of the direct progenitor of variant B.1.1.7. 416



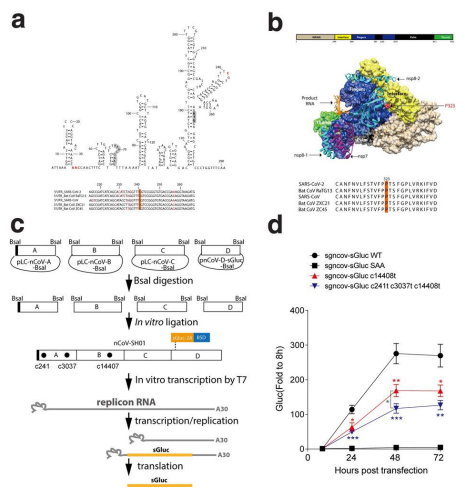
Phylogenetic analysis of variant B.1.1.7 by Spike

Fig. 2 The Canidae family could be a possible host of the direct progenitor of variant B.1.1.7. 419 The digits on the left of the figure indicate the labels of mutations, which correspond with the 420 mutation labels in Fig. 1a. The strains shown in the center of the figure contain at least one spike 421 mutation of variant B.1.1.7. And these strain examples cover all existing SARS-CoV-2 viruses that 422 collected from animal hosts. The strain labeled by orange star corresponds with the star variant in Fig. 423 1a. The strain with orange solid-round label was collected from a dog on 2020-07-28. Such two 424 strains share the same mutations "56" and have the minimum phylogenetic distance by MEGA tool. 425 Canidae, Mustelidae or Felidae, especially the Canidae family (for example, dog) could be a possible 426 host of the direct progenitor of variant B.1.1.7 based on existing stains collected before the end of 427 428 Jan. 2021.



429

Fig. 3 The strong correlations suggest that the top 25 mutations form eight potential co-mutation patterns. a The correlation heatmap of the top 25 mutations. These mutations could be grouped into several clusters with high Pearson-correlation-coefficient (PCC). b Regression analysis of mutations shows that eight clusters all denote the statistical significance level: ***p value < E-44. c to k show the transitions of the high-frequency mutations. The sky-blue represents the rate per day of initial residue in population and the golden the rate per day of substitution/mutant. These mutation transitions provide further evidence that the above mutations potentially form co-mutation patterns.



437

Fig. 4 Dominant co-mutation attenuates viral replication. a Predicted RNA structure of the 438 SARS-CoV-2 5'UTR. RNA structure of the 400-nt 5'UTR was predicted by "RNAstructure" 439 (http://rna.urmc.rochester.edu/RNAstructureWeb). The start codon for nsp1 is grey, the TRS-L is 440 orange, and the mutated nucleotides are red. The bottom panel shows the alignment of the 5'UTR of 441 SARS-CoV-2 with 5'UTRs of related viruses, with c241 highlighted. b Structure of SARS-CoV-2 442 RdRp/RNA complex. The structure of SARS-CoV-2 RdRp/RNA complex (PDB, 6X2G) was visualized 443 by Chimera (UCSF). The P323 mutation is highlighted in red, with the alignment of the amino acid 444 sequences of SARS-CoV-2 and related viruses near the P323 position. c Schematic of the in-vitro 445 ligation system for SARS-CoV-2 replicon. Four plasmids encompassing the viral genome were 446 digested by Bsal to release the four fragments. After gel purification, the fragments were ligated by 447 T4 ligase. The ligation products were purified and used as template for RNA *in-vitro* transcription. 448 sGluc, secreted Gaussia luciferase; 2A, foot-and-mouth disease virus (FMDV) 2A peptide; BSD, 449 blasticidin. d Huh7 cells were co-transfected with in-vitro transcribed replicon RNA (WT or the 450 451 indicated mutants) and an mRNA encoding the SARS-CoV-2 N protein. The luciferase activity in the supernatants was measured at the time points indicated. Medium was changed at 8 hours 452 453 post-transfection. Data are shown as mean±SEM (n=8). SAA, the NSP12 polymerase active-site mutant. Unpaired Student's t-test was performed between the mutants and wild type (WT) and 454 between the mutants as indicated (statistical significance level: *p value<0.05, **p value<0.01, ***p 455 value<0.001). 456