1	Discovery of a 382-nt deletion during the early evolution of SARS-CoV-2
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3	Running title: A SARS-CoV-2 deletion variant
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20 To date, the SARS-CoV-2 genome has been considered genetically more stable than 21 SARS-CoV or MERS-CoV. Here we report a 382-nt deletion covering almost the entire 22 open reading frame 8 (ORF8) of SARS-CoV-2 obtained from eight hospitalized patients 23 in Singapore. The deletion also removes the ORF8 transcription-regulatory sequence 24 (TRS), which in turn enhances the downstream transcription of the N gene. We also 25 found that viruses with the deletion have been circulating for at least four weeks. 26 During the SARS-CoV outbreak in 2003, a number of genetic variants were observed in 27 the human population [1], and similar variation has since been observed across SARS-28 related CoVs in humans and bats. Overwhelmingly these viruses had mutations or deletions in ORF8, that have been associated with reduced replicative fitness of the 29 30 virus [2]. This is also consistent with the observation that towards the end of the 31 outbreak sequences obtained from human SARS cases possessed an ORF8 deletion that 32 may be associated with host adaptation [1]. We therefore hypothesise that the major 33 deletion revealed in this study may lead to an attenuated phenotype of SARS-CoV-2. 34

35 On 1 December 2019, a novel coronavirus emerged from Hubei province in China and 36 infected people visiting the Huanan seafood market in Wuhan [3]. The virus demonstrated efficient human-to-human transmission within mainland China and subsequently spread 37 38 across many countries. The virus was soon identified as 2019-nCoV, more recently 39 designated SARS-CoV-2, while the disease is referred to as COVID-19. On 30 January 2020, the World Health Organization declared a Public Health Emergency of International 40 Concern. As of 10 March 2020, the COVID-19 outbreak has led to 113,702 confirmed cases 41 42 and 4,012 deaths globally [4]. Zoonotic viruses have crossed the species barrier from animals to humans and the success of an interspecies transmission is the ability of a novel virus to 43

44 adapt, via genetic mutation events, to a new host and cause sustained transmission that can45 lead to a significant outbreak.

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47 Nasopharyngeal swabs collected from hospitalized patients positive for SARS-CoV-2 in Singapore were subjected to next generation sequencing (NGS) analysis, with and without 48 49 passaging in Vero-E6 cells. The NGS data revealed a 382-nt deletion towards the 3' end of 50 the viral genomes obtained from multiple patients (Suppl. Table 1). To confirm this 51 observation, specific PCR primers were designed flanking the deleted region and Sanger 52 sequencing performed (Suppl. Fig. 1). This verified the deletion at positions 27,848 to 28,229 53 of the SARS-CoV-2 genome. Interrogation of the NGS assemblies of these 382-nt deletion 54 variants (referred to hereafter as $\Delta 382$) indicated that the virus populations were 55 homogenous.

56

57 Apart from the 382-nt deletion, the genome organization of the Δ 382 viruses is identical to 58 that of other SARS-CoV-2 (Fig. 1A). Closer examination of the deletion indicated that it spans an area of the ORF7b and ORF8 that includes the ORF8 transcriptional regulator 59 60 sequence (TRS), eliminating ORF8 transcription (Fig. 1B). The ORF8 region has been identified as an evolutionary hotspot of SARSr-CoVs. For instance, sequences of human 61 62 SARS-CoV TOR2 and LC2 exhibit 29-nt and 415-nt deletions, whereas those of bat SARS-63 CoV JTMC15 have discontinuous deletions in ORF7/8 (Fig. 1B) [5-8]. The ORF8 region of SARS-CoVs has been shown to play a significant role in adaptation to the human host 64 following interspecies transmission [7] and virus replicative efficiency [2]. Similarly the 65 66 disruption of ORF8 region in $\triangle 382$ viruses may be a result of human adaptation after the emergence of SARS-CoV-2. 67

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69 To investigate the possible effects of the TRS deletion, we calculated and compared the total 70 number of transcripts per million (TPM) of each gene between wild-type (WT) and $\Delta 382$ viruses. For each gene, we counted unambiguous reads that uniquely mapped to the gene-71 72 specific transcripts that include the joint leader and TRS sequences [9]. We analysed three 73 WT and two $\triangle 382$ sequences for which two independent NGS library preparations were 74 available. Our results indicate differential patterns of TPM level between WT and $\Delta 382$ 75 viruses (Fig. 2A). The $\triangle 382$ sequences displayed a greater level of TPMs in the ORF6, 76 ORF7a and N genes compared to WT. More specifically, the $\Delta 382$ N gene, downstream of 77 ORF8, showed a significant increase in the TPM level. To further determine the impact of the 382-nt deletion on the transcription of the N gene, we conducted qPCR to measure the 78 79 relative abundance of the RNAs from 2 genes (E and N) located upstream and downstream of 80 the deleted region, respectively. Results shown in Figure 2B confirm the enhancement of N 81 gene transcription in $\Delta 382$ viruses. This enhancement was consistently observed regardless of 82 the sample type, whether directly from swab samples or passaged viruses.

83

The global phylogeny of SARS-CoV-2 (Suppl. Fig. 2) revealed a comb-like appearance in 84 85 the phylogenetic tree and a general lack of phylogenetic resolution, reflecting the high similarity of the virus genomes. This tree topology is a typical characteristic of a novel 86 interspecies transmission, wherein the viruses are infecting immunologically naïve 87 88 populations with sustained human-to-human transmission, as previously shown for the early 89 stages of the 2009 H1N1 pandemic [10]. Notably, we observe an emerging virus lineage (tentatively designated as "Lineage 1" in Suppl. Fig. 2) that consists of sequences from 90 91 various newly affected countries with recent virus introductions. All viruses from this recent 92 lineage possess an amino acid substitution at residue 250 (from lysine to serine) on the ORF8

gene [11], which is known to undergo strong positive selection during animal-to-humantransmission [7].

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96 To estimate the divergence times among lineages of SARS-CoV-2, we reconstructed a dated 97 phylogeny based on 137 complete genomes, including data from this study. The estimated 98 rate of nucleotide substitutions among SARS-CoV-2 viruses is approximately at 8.68×10⁻⁴ substitutions per site per year (95% HPD: $5.44 \times 10^{-4} - 1.22 \times 10^{-3}$), which is moderately lower 99 than SARS-CoV ([12]: 0.80–2.38×10⁻³) and MERS-CoV (with a mean rate [13] of: 1.12×10⁻³ 100 and 95% HPD: $58.76 \times 10^{-4} - 1.37 \times 10^{-3}$) as well as human seasonal influenza viruses (ranging 101 1.0-5.5×10⁻³ depending on individual gene segments [10, 14-16]. In contrast, the estimated 102 103 rate of SARS-CoV-2 viruses is greater than estimates for human coronaviruses by a 104 magnitude order of 4×10^{-4} [17].

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106 The mean TMRCA estimates indicate the introduction of SARS-CoV-2 into humans (Fig. 3. 107 node A) occurred in the middle of November 2019 (95% HPD: 2019.77–2019.94) (Table 1), suggesting that the viruses were present in human hosts approximately one month before the 108 109 outbreak was detected. A single amino-acid mutation was fixed on the ORF8 region of all 110 full-genome Lineage 1 viruses (Fig. 3, node B), with the TMRCA estimate of 22 December 111 2019 (95% HPD: 2019.9–2020.0). The Δ 382 viruses form a monophyletic group within 112 Lineage 1, and dating estimates indicate that they may have arisen in the human population in 113 Singapore around 07 February 2020 (95% HPD: 2020.12–2020.06), consistent with the date of collection of the $\triangle 382$ positive samples (17–19 February 2020). The three $\triangle 382$ viruses 114 115 shared a high level of nucleotide similarities (99.9%) with only 5 nucleotide differences 116 between them. Although the dated phylogeny indicates that $\Delta 382$ viruses clustered together 117 and are possibly derived from a single source, there is a general lack of statistical support in

SARS-CoV-2 phylogenies due to its recent emergence, and evolutionary inferences should bemade with caution.

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121 In this report we describe the first major evolutionary event of the SARS-CoV-2 virus 122 following its emergence into the human population. Although the biological consequences of 123 this deletion remain unknown, the alteration of the N gene transcription would suggest this may have an impact on virus phenotype. Recent work has indicated that ORF8 of SARS-CoV 124 125 plays a functional role in virus replicative fitness and may be associated with attenuation 126 during the early stages of human-to-human transmission [2]. Given the prevalence of a variety of deletions in the ORF8 of SARSr-CoVs, it is likely that we will see further deletion 127 128 variants emerge with the sustained transmission of SARS-CoV-2 in humans. Future research 129 should focus on the phenotypic consequences of $\Delta 382$ viruses in the transmission dynamics 130 of the current epidemics and the immediate application of this genomic marker for molecular epidemiological investigation. 131

132	Online content. Any methods, additional references, Nature Research reporting summaries,
133	source data, extended data, supplementary information, acknowledgements, peer review
134	information; details of author contributions and competing interests; and statements of data
135	availability are available at [Article DOI].
136	
137	Data availability. The new sequences generated in this study have been deposited in
138	GISAID database under accession numbers EPI_ISL_407987, EPI_ISL_407988,
139	EPI_ISL_410535 to EPI_ISL_410537 for WT viruses and EPI_ISL_414378 to
140	EPI_ISL_414380 for $\triangle 382$ viruses.
141	
142	Acknowledgements. This study was supported by the Duke-NUS Signature Research
143	Programme funded by the Ministry of Health, Singapore, the National Medical Research
144	Council under its COVID-19 Research Fund (NMRC Project No. COVID19RF-001) and by
145	National Research Foundation Singapore grant NRF2016NRFNSFC002-013 (Combating the
146	Next SARS-or MERS-Like Emerging Infectious Disease Outbreak by Improving Active
147	Surveillance). We thank Viji Vijayan, Benson Ng and Velraj Sivalingam of the Duke-NUS
148	Medical School ABSL3 facility for logistics management and assistance. We thank all
149	scientific staff who assisted with processing clinical samples, especially Velraj Sivalingam,
150	Adrian Kang, Randy Foo, Wan Ni Chia and Akshamal Gamage. We thank Su Ting Tay,
151	Ming Hui Lee and Angie Tan from the Duke-NUS Medical School Genome Biology Facility
152	for expert and rapid technical assistance.
153	
154	Author contributions. YCFS, DEA, LFW, GJDS designed and supervised research. BEY,
155	SK, JGHL, LS, GZY, YSL, DCL collected and provided samples. DEA, ML, YZ conducted

156	experiments. YCFS, ZF, JJ, IHM, GJDS performed analyses. YCFS, ZF, ML, LFW, GJDS
157	wrote the paper. All authors reviewed and approved the manuscript.

- 158
- 159 Methods

Ethics statement. This study was undertaken as part of the national disease outbreak and the
response and the protocols were approved by the ethics committee of the National Healthcare
Group. Patient samples were collected under PROTECT (2012/00917), a multi-centred
Prospective Study to Detect Novel Pathogens and Characterize Emerging Infections. Work
undertaken at the Duke-NUS Medical School ABSL3 laboratory was approved by the DukeNUS ABSL3 Biosafety Committee, National University of Singapore and Ministry of Health
Singapore.

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Virus culture, RNA extraction and sequencing. Clinical samples from infected patients 168 169 were collected from public hospitals in Singapore from January-February 2020. Material 170 from clinical samples was used to inoculate Vero-E6 cells (ATCC[®]CRL-1586TM). Total RNA 171 was extracted using E.Z.N.A. Total RNA Kit I (Omega Bio-tek) according to manufacturer's instructions and samples analysed by real-time quantitative reverse transcription-PCR (RT-172 qPCR) for the detection of SARS-CoV-2 as previously described [18]. Whole genome 173 174 sequencing was performed using next-generation sequencing (NGS) methodology. The 175 cDNA libraries were constructed using TruSeq RNA Library Prep Kit (Illumina) according to 176 the manufacturer's instructions and sequenced on an Illumina MiSeg System. Raw NGS reads were trimmed by Trimmomatic v0.39 [19] to remove adaptors and low-quality bases. 177 178 Genome sequences were assembled and consensus sequences obtained using the BWA algorithm in UGENE v.33. To verify the presence of the deletion in the SARS-CoV-2 179 180 genome, we designed two specific PCR primers (F primer: 5'-

181 TGTTAGAGGTACAACAGTACTTT-3' and R primer: 5'-

- 182 GGTAGTAGAAATACCATCTTGGA-3') targeting the ORF7–8 regions. For samples with
- 183 low Ct values, a hemi-nested PCR was then performed with primers (5'-
- 184 TGTTTATAACACTTTGCTTCACA-3') and (5'-GGTAGTAGAAATACCATCTTGGA-
- 185 3'). The PCR mixture contained the cDNA, primers (10μ M each), 10x Pfu reaction buffer
- 186 (Promega), Pfu DNA polymerase (Promega) and dNTP mix (10mM, Thermo Scientific). The
- 187 PCR reaction was carried out with the following conditions: 95°C for 2 min, 35 cycles at
- 188 95°C for 1 min, 52°C for 30 sec, 72°C for 1 min and a final extension at 72°C for 10 min in a
- 189 thermal cycler (Applied Biosystems Veriti). Deletions in the PCR products were visualized
- 190 by gel electrophoresis and confirmed by Sanger sequencing. Three full $\Delta 382$ genomes were
- 191 generated and designated BetaCoV/Singapore/12/2020, BetaCoV/Singapore/13/2020 and
- 192 BetaCoV/Singapore/14/2020 along with five WT SARS-CoV-2 genomes previously
- 193 generated in our laboratories and deposited in GISAID (see Suppl. Table 1).
- 194
- **Genomic characterization.** To characterize and map the deletion regions of SARS-CoV-2
- 196 viruses, we compared with available SARS-CoV-2 and SARs-CoV related genomes from bat
- 197 SARs-CoV-JTMC15 (accession number: KY182964) and humans SARs-CoV-Tor 2
- 198 (accession number: AY274119) and LC2 (accession number: AY394999). The viral genome
- 199 organizations of Wuhan-Hu-1 (accession number: MN908947) and Singapore SARS-CoV-2
- 200 (Singapore/2 /2020: EPI_ISL_407987) comprise the following gene order and lengths:
- 201 ORF1ab (open-reading frame) replicase (21291 nt), spike (S: 3822 nt), ORF3 (828 nt),
- 202 envelope (E: 228 nt), membrane (M: 669 nt), ORF6 (186 nt), ORF7ab (498 nt), ORF8 (366
- 203 nt) and nucleocapsid (N: 1260 nt).
- 204

205 To understand possible effects of the TRS deletion on the ORF8 region, we analysed the total 206 number of transcripts per million (TPM) of each gene between wildtype and $\Delta 382$ variants. 207 For each gene, unambiguous reads which uniquely mapped to the specific joint leader and 208 TRS sequences are counted. We analysed three patients of SARS-CoV-2 wild type and two 209 patients of SARS-CoV-2 variants and independent NGS library preparations were performed 210 twice for each sample. The Wuhan-Hu-1 was used as the reference sequence for genomic 211 coordinates. Leader RNA sequence is situated at the 5' UTR region of the genome and the 212 core sequence consists of 26 nt: UCUCUAAACGAACUUUAAAAUCUGUG. Each of the 213 coding genes is preceded by a core transcription regulatory sequence (TRS) which is highly 214 conserved (i.e. ACGAAC), resembling the leader core sequence. The leader RNA is joined to 215 the TRS regions of different genes and the joining is required for driving subgenomic 216 transcription. Coding sequence (CDS), leader and TRS sequences annotations were generated 217 in Geneious and followed published SARS-CoV studies [9]. To characterize the differential 218 levels of TRS for each gene, 70 nt leader sequence and 230 nt downstream of each TRS 219 sequence were annotated individually to form a 300bp leader-TRS transcript for the splicing-220 aware aligners in R package rtracklayer (v 1.44). NGS raw fastq reads were then mapped to 221 the reference genome by Geneious RNA-Seq mapper with the annotation of splice-junctions for the leader-TRS. For each gene, the TPM of leader-TRS was then calculated in Geneious 222 223 by excluding ambiguous reads which may come from other TRS. The resulting TPM data 224 was generated in Geneious and graphs plotted using ggplot 2 (v3.2.0) in R v3.6.1. Wilcoxon 225 one-sided tests were performed in R to test the significant differences between WT and $\Delta 382$ 226 samples. We also compared individual Ct values of E and N genes between WT and $\Delta 382$ 227 viruses using qRT-PCR assays as previously described [18]. Samples used in this analysis included nasopharyngeal swabs from three WT patient samples and both nasopharyngeal 228

swabs and passaged samples from two Δ382 patients (n=7). Significance between groups was
assessed using a one-way analysis of variance (ANOVA) in PRISM v8.3.1.

231

232 Phylogenetic analyses. All available genomes of SARS-CoV-2 with associated virus 233 sampling dates were downloaded from the GISAID database. Genome sequence alignment 234 was performed and preliminary maximum likelihood phylogenies of complete genome were 235 reconstructed using RAxML with 1,000 bootstrap replicates in Geneious R9.0.3 software 236 (Biomatters Ltd). Any sequence outliers were removed from subsequent analyses. To 237 reconstruct a time-scaled phylogeny, an uncorrelated lognormal relaxed-clock model with an 238 exponential growth coalescent prior and the HKY85+ Γ substitution model was used in the 239 program BEAST v1.10.4 [20] to simultaneously estimate phylogeny, divergence times and 240 rates of nucleotide substitution. Four independent Markov Chain Monte Carlo (MCMC) runs 241 of 100 million generations were performed and sampled every 10,000 generations. The runs 242 were checked for convergence in Tracer v1.7 [21] and that effective sampling size (ESS) 243 values of all parameters was \geq 200. The resulting log and tree files were combined after 244 removing appropriate burn-in values using LogCombiner [20], and the maximum clade 245 credibility (MCC) tree was subsequently generated using TreeAnnontator [20].

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298

299 Table 1. TMRCA estimates of major SARS-CoV-2 nodes.

Node	Mean TMRCA	Lower 95% HPD	Upper 95% HPD
А	2019.87 (15 November 2019)	2019.77 (09 October 2019)	2019.94 (11 December 2019)
В	2019.97 (22 Dec 2019)	2019.90 (26 November 2019)	2020.00 (01 January 2020)
С	2020.09 (07 Feb 2020)	2020.06 (24 January 2020)	2020.12 (14 Feb 2020)

300 Abbreviations: TMRCA, time to most recent common ancestor, HPD: highest posterior

301 density.

Figure 1.

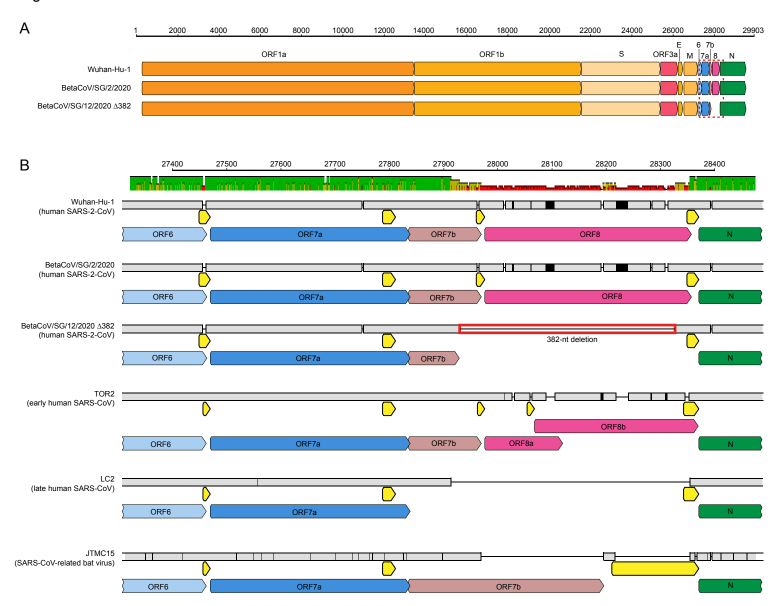


Figure 1. Schematic comparison of SARS-2-CoV, SARS-CoV and SARSr-CoV genomes.

A. Full genome sequences of SARS-2-CoV isolates Wuhan-Hu-1 (GenBank: MN908947), BetaCoV/Singapore/2/2020 (GISAID: 407987), and BetaCoV/Singapore/12/2020 \triangle 382 (GISAID: EPI_ISL_414378). **B.** Magnification of genomic region (dashed box in Figure 1A), numbers on horizontal axes indicate the nucleotide position relative to Wuhan-Hu-1, grey boxes indicate sequence coverage at a certain position, black horizontal lines represent a deletion at a certain position, open reading frames (ORFs) are indicated by colored arrows. A red-lined box indicates the 382-nt deletion. Transcription-regulatory sequences (TRSs) are indicated by yellow arrows.

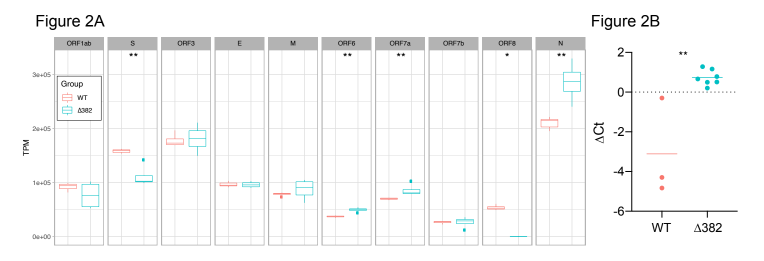


Figure 2. Comparison of transcription of the transcription of the N gene in wild-type (WT) versus \triangle 382 viruses. A. Abundance of mapped reads relative to transcriptional regulatory sequence (TRS) positions across the genome. 70 bp of leader sequence and 230 of each TRS-downstream sequence were merged in a splice-junction in the annotation. Transcripts per million reads (TPM) was calculated from reads mapped specifically to each leader-TRS region and a whisker and scatter plot was drawn for each gene. A Wilcoxon test was applied to the TPM for each gene of \triangle 382 to WT (*: p ≤ 0.05 **: p ≤ 0.01). B. Relative abundance of the RNAs of the E gene in comparison to the N gene of WT versus \triangle 382 SARS-CoV-2 (p = 0.002) using qPCR. Horizontal bars indicate the mean of each group.

Figure 3.

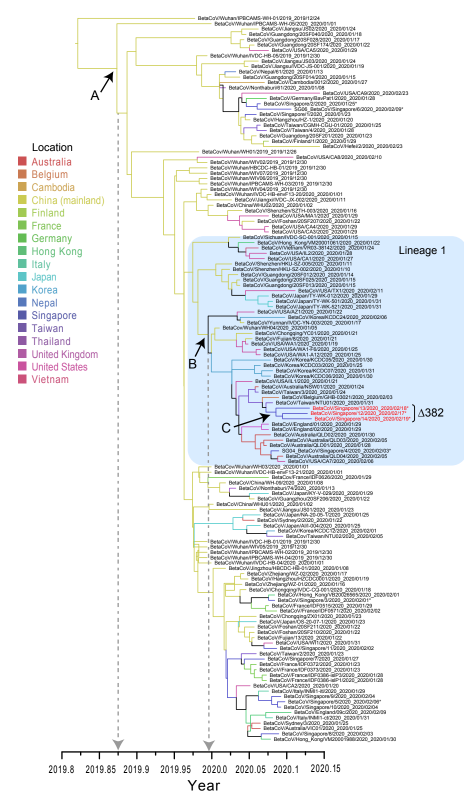
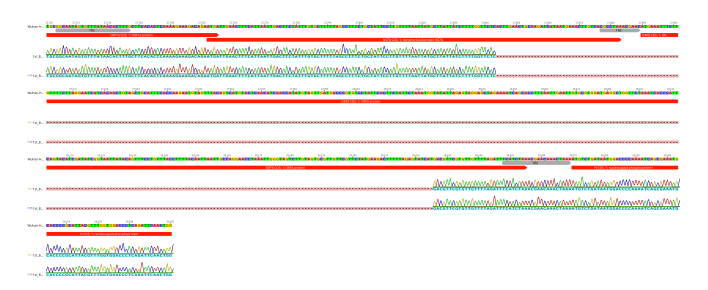
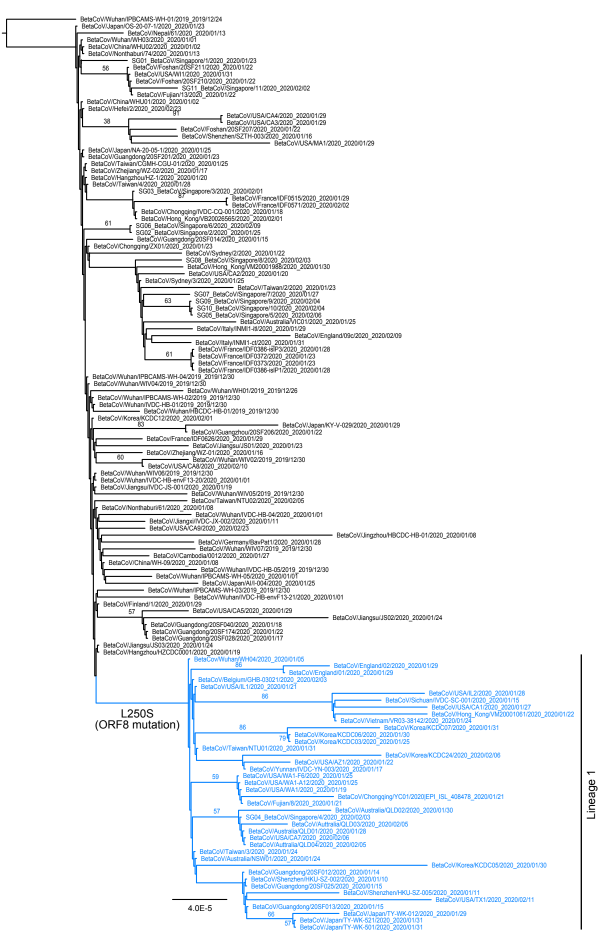


Figure 3. Temporal phylogeny of the complete genomes of SARS-CoV-2 viruses reconstructed using an uncorrelated lognormal relaxed clock model in BEAST. Isolate names in red indicate Δ 382 viruses with a large deletion in the ORF7/8 region as described in this study. *Indicates virus genomes generated in this study. Colored branches denote different geographical locations.



Supplementary Figure 1. Sanger sequences of BetaCoV/Singapore/12/2020 \triangle 382 mapped to Wuhan-Hu-1 showing the position of the 382-nt deletion in the SARS-CoV-2 genome.



Supplementary Figure 2. Maximum-likelihood tree of SARS-CoV-2 genomes reconstructed using RAxML with 1,000 bootstrap replicates. \triangle 382 viruses were excluded from this analysis as the L250S mutation is located in the ORF8 deletion.

Supplementary Table 1.

Accession number	Virus isolato	Collection date	Noto
Accession number EPI_ISL_414378	Virus isolate BetaCoV/Singapore/12/2020	Collection date 17/02/2020	Note 382-nt deletion (this study)
EPI_ISL_414378	BetaCoV/Singapore/13/2020	18/02/2020	382-nt deletion (this study)
EPI_ISL_414379	BetaCoV/Singapore/14/2020	19/02/2020	382-nt deletion (this study)
EPI_ISL_407987	BetaCoV/Singapore/2/2020	25/01/2020	Wild type (this study)
EPI ISL 407988	BetaCoV/Singapore/3/2020	01/02/2020	Wild type (this study) Wild type (this study)
EPI_ISL_410535	BetaCoV/Singapore/4/2020	03/02/2020	Wild type (this study) Wild type (this study)
EPI_ISL_410536	BetaCoV/Singapore/5/2020	06/02/2020	Wild type (this study) Wild type (this study)
EPI_ISL_410537	BetaCoV/Singapore/6/2020	09/02/2020	Wild type (this study)
EPI_ISL_402119	BetaCoV/Wuhan/IVDC-HB-01/2019	30/12/2019	Wild type
EPI_ISL_402120	BetaCoV/Wuhan/IVDC-HB-04/2020	01/01/2020	Wild type
EPI_ISL_402121	BetaCoV/Wuhan/IVDC-HB-05/2019	30/12/2019	Wild type
EPI_ISL_402123	BetaCoV/Wuhan/IPBCAMS-WH-01/2019	24/12/2019	Wild type
EPI_ISL_402124	BetaCoV/Wuhan/WIV04/2019	30/12/2019	Wild type
EPI_ISL_402127	BetaCoV/Wuhan/WIV02/2019	30/12/2019	Wild type
EPI_ISL_402128	BetaCoV/Wuhan/WIV05/2019	30/12/2019	Wild type
EPI_ISL_402129	BetaCoV/Wuhan/WIV06/2019	30/12/2019	Wild type
EPI_ISL_402130	BetaCoV/Wuhan/WIV07/2019	30/12/2019	Wild type
EPI ISL 402132	BetaCoV/Wuhan/HBCDC-HB-01/2019	30/12/2019	Wild type
EPI_ISL_403928	BetaCoV/Wuhan/IPBCAMS-WH-05/2020	01/01/2020	Wild type
EPI_ISL_403929	BetaCoV/Wuhan/IPBCAMS-WH-04/2019	30/12/2019	Wild type
EPI ISL 403930	BetaCoV/Wuhan/IPBCAMS-WH-03/2019	30/12/2019	Wild type
EPI_ISL_403931	BetaCoV/Wuhan/IPBCAMS-WH-02/2019	30/12/2019	Wild type
EPI_ISL_403932	BetaCoV/Guangdong/20SF012/2020	14/01/2020	Wild type
EPI_ISL_403933	BetaCoV/Guangdong/20SF013/2020	15/01/2020	Wild type
EPI_ISL_403934	BetaCoV/Guangdong/20SF014/2020	15/01/2020	Wild type
EPI_ISL_403935	BetaCoV/Guangdong/20SF025/2020	15/01/2020	Wild type
EPI_ISL_403936	BetaCoV/Guangdong/20SF028/2020	17/01/2020	Wild type
EPI_ISL_403937	BetaCoV/Guangdong/20SF040/2020	18/01/2020	Wild type
EPI_ISL_403962	BetaCoV/Nonthaburi/61/2020	08/01/2020	Wild type
EPI_ISL_403963	BetaCoV/Nonthaburi/74/2020	13/01/2020	Wild type
EPI_ISL_404227	BetaCoV/Zhejiang/WZ-01/2020	16/01/2020	Wild type
EPI_ISL_404228	BetaCoV/Zhejiang/WZ-02/2020	17/01/2020	Wild type
EPI_ISL_404253	BetaCoV/USA/IL1/2020	21/01/2020	Wild type
EPI_ISL_404895	BetaCoV/USA/WA1/2020	19/01/2020	Wild type
EPI_ISL_405839	BetaCoV/Shenzhen/HKU-SZ-005/2020	11/01/2020	Wild type
EPI_ISL_406030	BetaCoV/Shenzhen/HKU-SZ-002/2020	10/01/2020	Wild type
EPI_ISL_406031	BetaCoV/Taiwan/2/2020	23/01/2020	Wild type
EPI_ISL_406034	BetaCoV/USA/CA1/2020	27/01/2020	Wild type
EPI_ISL_406036	BetaCoV/USA/CA2/2020	20/01/2020	Wild type
EPI_ISL_406223	BetaCoV/USA/AZ1/2020	22/01/2020	Wild type
EPI_ISL_406531	BetaCoV/Guangdong/20SF174/2020	22/01/2020	Wild type
EPI_ISL_406533	BetaCoV/Guangzhou/20SF206/2020	22/01/2020	Wild type
EPI_ISL_406534	BetaCoV/Foshan/20SF207/2020	22/01/2020	Wild type
EPI_ISL_406535	BetaCoV/Foshan/20SF210/2020	22/01/2020	Wild type
EPI_ISL_406536	BetaCoV/Foshan/20SF211/2020	22/01/2020	Wild type
EPI_ISL_406538	BetaCoV/Guangdong/20SF201/2020	23/01/2020	Wild type
EPI_ISL_406594	BetaCoV/Shenzhen/SZTH-003/2020	16/01/2020	Wild type
EPI_ISL_406597	BetaCoV/France/IDF0373/2020	23/01/2020	Wild type
EPI_ISL_406716	BetaCoV/China/WHU01/2020	02/01/2020	Wild type
EPI_ISL_406717	BetaCoV/China/WHU02/2020	02/01/2020	Wild type
EPI_ISL_406798	BetaCov/Wuhan/WH01/2019	26/12/2019	Wild type
EPI_ISL_406800	BetaCov/Wuhan/WH03/2020	01/01/2020	Wild type
EPI_ISL_406801	BetaCov/Wuhan/WH04/2020	05/01/2020	Wild type
EPI_ISL_406844	BetaCoV/Australia/VIC01/2020	25/01/2020	Wild type
EPI_ISL_406862	BetaCoV/Germany/BavPat1/2020	28/01/2020	Wild type
EPI_ISL_406970	BetaCoV/Hangzhou/HZ-1/2020	20/01/2020	Wild type
EPI_ISL_406973	BetaCoV/Singapore/1/2020	23/01/2020	Wild type
EPI_ISL_407071	BetaCoV/England/01/2020	29/01/2020	Wild type

EPI_ISL_407073	BetaCoV/England/02/2020	29/01/2020	Wild type
EPI_ISL_407079	BetaCoV/Finland/1/2020	29/01/2020	Wild type
EPI_ISL_407084	BetaCoV/Japan/AI/I-004/2020	25/01/2020	Wild type
EPI_ISL_407193	BetaCoV/Korea/KCDC03/2020	25/01/2020	Wild type
EPI_ISL_407214	BetaCoV/USA/WA1-A12/2020	25/01/2020	Wild type
 EPI_ISL_407215	BetaCoV/USA/WA1-F6/2020	25/01/2020	Wild type
EPI_ISL_407313	BetaCoV/Hangzhou/HZCDC0001/2020	19/01/2020	Wild type
EPI_ISL_407893	BetaCoV/Australia/NSW01/2020	24/01/2020	Wild type
EPI ISL 407894	BetaCoV/Australia/QLD01/2020	28/01/2020	Wild type
EPI ISL 407896	BetaCoV/Australia/QLD02/2020	30/01/2020	Wild type
EPI ISL 407976	BetaCoV/Belgium/GHB-03021/2020	03/02/2020	Wild type
EPI_ISL_408008	BetaCoV/USA/CA3/2020	29/01/2020	Wild type
EPI_ISL_408008	BetaCoV/USA/CA4/2020	29/01/2020	Wild type
		29/01/2020	
EPI_ISL_408010	BetaCoV/USA/CA5/2020		Wild type
EPI_ISL_408430	BetaCoV/France/IDF0515/2020	29/01/2020	Wild type
EPI_ISL_408431	BetaCov/France/IDF0626/2020	29/01/2020	Wild type
EPI_ISL_408478	BetaCoV/Chongqing/YC01/2020	21/01/2020	Wild type
EPI_ISL_408479	BetaCoV/Chongqing/ZX01/2020	23/01/2020	Wild type
EPI_ISL_408480	BetaCoV/Yunnan/IVDC-YN-003/2020	17/01/2020	Wild type
EPI_ISL_408481	BetaCoV/Chongqing/IVDC-CQ-001/2020	18/01/2020	Wild type
EPI_ISL_408484	BetaCoV/Sichuan/IVDC-SC-001/2020	15/01/2020	Wild type
EPI_ISL_408486	BetaCoV/Jiangxi/IVDC-JX-002/2020	11/01/2020	Wild type
EPI_ISL_408488	BetaCoV/Jiangsu/IVDC-JS-001/2020	19/01/2020	Wild type
EPI_ISL_408489	BetaCoV/Taiwan/NTU01/2020	31/01/2020	Wild type
EPI_ISL_408514	BetaCoV/Wuhan/IVDC-HB-envF13-20/2020	01/01/2020	Wild type
EPI_ISL_408515	BetaCoV/Wuhan/IVDC-HB-envF13-21/2020	01/01/2020	Wild type
EPI_ISL_408665	BetaCoV/Japan/TY-WK-012/2020	29/01/2020	Wild type
EPI_ISL_408666	BetaCoV/Japan/TY-WK-501/2020	31/01/2020	Wild type
EPI_ISL_408667	BetaCoV/Japan/TY-WK-521/2020	31/01/2020	Wild type
EPI_ISL_408668	BetaCoV/Vietnam/VR03-38142/2020	24/01/2020	Wild type
EPI_ISL_408669	BetaCoV/Japan/KY-V-029/2020	29/01/2020	Wild type
EPI ISL 408670	BetaCoV/USA/WI1/2020	31/01/2020	Wild type
EPI_ISL_408976	BetaCoV/Sydney/2/2020	22/01/2020	Wild type
EPI ISL 408977	BetaCoV/Sydney/3/2020	25/01/2020	Wild type
EPI_ISL_409067	BetaCoV/USA/MA1/2020	29/01/2020	Wild type
EPI_ISL_410045	BetaCoV/USA/IL2/2020	28/01/2020	Wild type
EPI_ISL_410218	BetaCov/Taiwan/NTU02/2020	05/02/2020	Wild type
EPI_ISL_410301	BetaCoV/Nepal/61/2020	13/01/2020	Wild type
EPI_ISL_410531	BetaCoV/Japan/NA-20-05-1/2020	25/01/2020	Wild type
EPI_ISL_410532	BetaCoV/Japan/OS-20-07-1/2020	23/01/2020	Wild type
EPI_ISL_410545	BetaCoV/Italy/INMI1-itl/2020	29/01/2020	Wild type
EPI ISL 410546	BetaCoV/Italy/INMI1-ct/2020	31/01/2020	Wild type
EPI ISL 410713	BetaCoV/Singapore/7/2020	27/01/2020	Wild type
EPI ISL 410714	BetaCoV/Singapore/8/2020	03/02/2020	Wild type
EPI_ISL_410715	BetaCoV/Singapore/9/2020	04/02/2020	Wild type
EPI_ISL_410716	BetaCoV/Singapore/10/2020	04/02/2020	Wild type
EPI_ISL_410717	BetaCoV/Australia/QLD03/2020	05/02/2020	Wild type
EPI_ISL_410718	BetaCoV/Australia/QLD03/2020 BetaCoV/Australia/QLD04/2020	05/02/2020	Wild type
		02/02/2020	
EPI_ISL_410719	BetaCoV/Singapore/11/2020		Wild type
EPI_ISL_410720	BetaCoV/France/IDF0372/2020	23/01/2020	Wild type
EPI_ISL_411060	BetaCoV/Fujian/8/2020	21/01/2020	Wild type
EPI_ISL_411066	BetaCoV/Fujian/13/2020	22/01/2020	Wild type
EPI_ISL_411218	BetaCoV/France/IDF0571/2020	02/02/2020	Wild type
EPI_ISL_411219	BetaCoV/France/IDF0386-isIP1/2020	28/01/2020	Wild type
EPI_ISL_411220	BetaCoV/France/IDF0386-isIP3/2020	28/01/2020	Wild type
EPI_ISL_411902	BetaCoV/Cambodia/0012/2020	27/01/2020	Wild type
EPI_ISL_411915	BetaCoV/Taiwan/CGMH-CGU-01/2020	25/01/2020	Wild type
EPI_ISL_411926	BetaCoV/Taiwan/3/2020	24/01/2020	Wild type
EPI_ISL_411927	BetaCoV/Taiwan/4/2020	28/01/2020	Wild type
EPI_ISL_411950	BetaCoV/Jiangsu/JS01/2020	23/01/2020	Wild type
EPI_ISL_411952	BetaCoV/Jiangsu/JS02/2020	24/01/2020	Wild type

EPI_ISL_411953	BetaCoV/Jiangsu/JS03/2020	24/01/2020	Wild type
EPI_ISL_411954	BetaCoV/USA/CA7/2020	06/02/2020	Wild type
EPI_ISL_411955	BetaCoV/USA/CA8/2020	10/02/2020	Wild type
EPI_ISL_411956	BetaCoV/USA/TX1/2020	11/02/2020	Wild type
EPI_ISL_411957	BetaCoV/China/WH-09/2020	08/01/2020	Wild type
EPI_ISL_412026	BetaCoV/Hefei/2/2020	23/02/2020	Wild type
EPI_ISL_412028	BetaCoV/Hong_Kong/VM20001061/2020	22/01/2020	Wild type
EPI_ISL_412029	BetaCoV/Hong_Kong/VM20001988/2020	30/01/2020	Wild type
EPI_ISL_412030	BetaCoV/Hong_Kong/VB20026565/2020	01/02/2020	Wild type
EPI_ISL_412116	BetaCoV/England/09c/2020	09/02/2020	Wild type
EPI_ISL_412459	BetaCoV/Jingzhou/HBCDC-HB-01/2020	08/01/2020	Wild type
EPI_ISL_412862	BetaCoV/USA/CA9/2020	23/02/2020	Wild type
EPI_ISL_412869	BetaCoV/Korea/KCDC05/2020	30/01/2020	Wild type
EPI_ISL_412870	BetaCoV/Korea/KCDC06/2020	30/01/2020	Wild type
EPI_ISL_412871	BetaCoV/Korea/KCDC07/2020	31/01/2020	Wild type
EPI_ISL_412872	BetaCoV/Korea/KCDC12/2020	01/02/2020	Wild type
EPI_ISL_412873	BetaCoV/Korea/KCDC24/2020	06/02/2020	Wild type