

Recombinant BA.1/BA.2 SARS-CoV-2 Virus in Arriving Travelers, Hong Kong, February 2022

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We studied SARS-CoV-2 genomes from travelers arriving in Hong Kong during November 2021–February 2022. In addition to Omicron and Delta variants, we detected a BA.1/BA.2 recombinant with a breakpoint near the 5' end of the spike gene in 2 epidemiologically linked case-patients. Continued surveillance for SARS-CoV-2 recombinants is needed.

The SARS-CoV-2 Omicron variant (Pango lineage B.1.1.529) emerged in November 2021. Within a few weeks, subvariants BA.1, BA.1.1, and BA.2 were detected in varying proportions on different continents, but BA.1 initially was dominant (1). By March 2022, these 3 subvariants accounted for >95% of sequences submitted to GISAID (<https://www.gisaid.org>). We previously demonstrated the feasibility of testing incoming travelers for SARS-CoV-2 genomic surveillance (2). We report detecting a BA.1/BA.2 recombinant SARS-CoV-2 subvariant in travelers arriving in Hong Kong, China.

Using our previously described next-generation sequencing method (2), we analyzed 198 (25%) of 793 SARS-CoV-2 reverse transcription PCR (RT-PCR)-positive samples collected from travelers arriving in Hong Kong during November 15, 2021–February 4, 2022 (Appendix Table 1, <https://wwwnc.cdc.gov/EID/article/28/6/22-0523-App1.pdf>). We randomly selected samples with cycle thresholds <30 and successfully deduced near-full genome sequences from 180 samples (mean coverage 97.6%; depth ≥ 100). Deduced genomes predominantly were Delta ($n = 58$) and Omicron (BA.1 = 66, BA.1.1 = 28, and BA.2 = 26) variants (Appendix Figures 1, 2). Time distribution of these variants agrees with global surveillance data

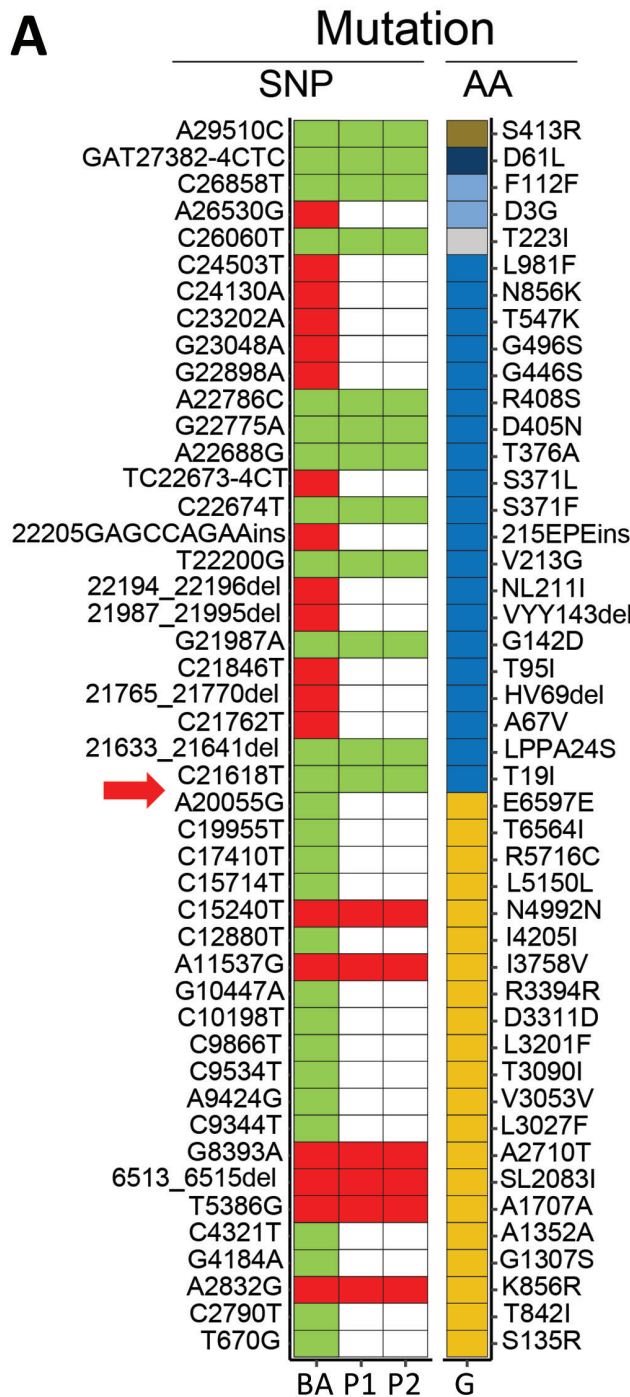
submitted to GISAID, confirming that travel hubs are useful sentinel sites to monitor SARS-CoV-2 circulation (2). Of note, the BA.2 cases we detected predominantly were imported from the Philippines and Nepal, indicating this subvariant might have become established in these countries before detection in Hong Kong.

In our phylogenetic analysis, 2 additional nearly identical sequences formed a distinct branch in the Omicron clade (Appendix Figure 2). We detected these sequences from 2 epidemiologically linked cases, patients 1 and 2, who were work colleagues and traveled together to Hong Kong on February 1, 2022, from Germany via the Netherlands. They tested SARS-CoV-2-positive by RT-PCR at the airport upon arrival (cycle thresholds 27 and 22). Patient 1 reported having a sore throat and cough since January 28, but patient 2 was asymptomatic. Both patients had received 2 doses of Pfizer-BioNTech COVID-19 vaccine (Pfizer Inc., <https://www.pfizer.com>); patient 1 received the second dose on November 1, 2021, and patient 2 received the second dose on June 22, 2021.

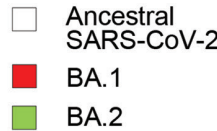
The distinct topology of viral sequences from these patients suggested that they were infected by a recombinant virus. To test that hypothesis, we used previously reported BA.1- and BA.2-defining single-nucleotide polymorphisms (SNPs) to analyze the genomes (Appendix). We found that the 5' end sequences (nucleotide region 1–20055) from the 2 cases only contained BA.1-specific SNPs (Figure, panel A). By contrast, the corresponding 3' end sequences only contained BA.2-specific SNPs. We further conducted a recombination analysis and confirmed that only 1 breakpoint was located within nucleotide positions 20055–21618 (Appendix Figure 3). The nucleotide 5' end of the sequences is phylogenetically similar to authentic BA.1 and the 3' end similar to BA.2 sequences at this breakpoint (Figure, panel B). The breakpoint identified in this recombinant virus is near the 5' end open reading frame of the spike gene. Recombinant viruses, including B.1.1.7/B1.177 and Delta/BA.1, with a breakpoint in this region have been reported (3; P. Colson et al., unpub. data, <https://www.medrxiv.org/content/10.1101/2022.03.03.22271812v2>; T. Peacock, unpub. data, <https://github.com/cov-lineages/pango-designation/issues/441>).

We further examined our sequence data to exclude the possibility of coinfection or contamination (4). We noted that the minor allele frequencies at these BA.1- and BA.2-defining SNP positions were extremely low (mean 0.5%, median 0.06%) (Figure, panel A), indicating these samples contained only 1

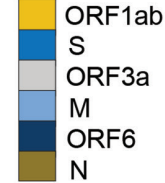
A



SNP



Gene



B

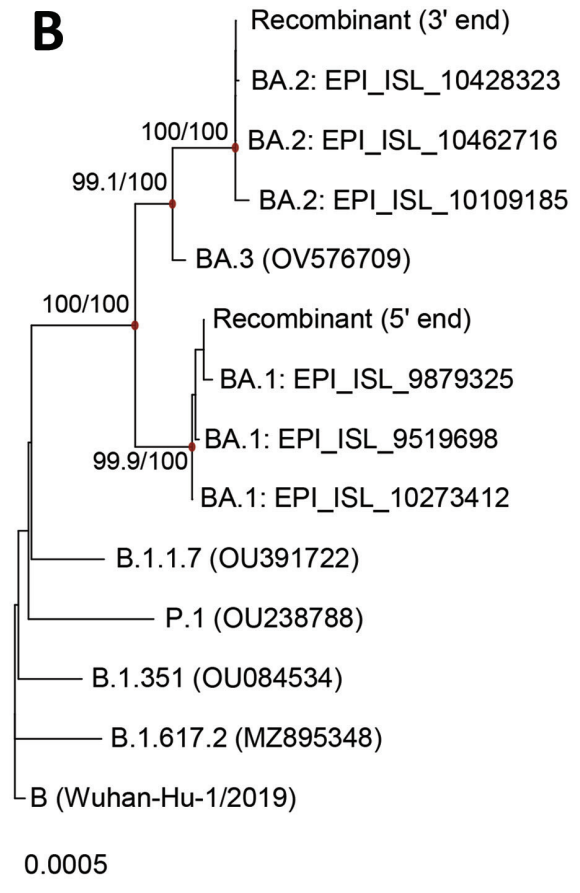


Figure. Detection of recombinant BA.1/BA.2 SARS-CoV-2 virus in arriving travelers, Hong Kong, China, February 2022. A) Mapping of BA.1- and BA.2-specific SNPs against the reference sequence genome (Genbank accession no. MN908947.3). Red boxes indicate BA.1-specific SNPs and green boxes indicate BA.2-specific SNPs found in samples from P1 and P2; the corresponding AA changes of these SNPs also are indicated. Red arrow indicates the putative breaking point. B) Phylogeny of viral RNA sequences at the 5' and 3' ends to the putative breakpoint. The maximum-likelihood tree was generated by using IQ-TREE (<http://www.iqtree.org>) and the transition plus empirical base frequencies plus proportion of invariable site nucleotide substitution model with Wuhan-Hu-1 (GenBank accession no. MN908947.3) as the outgroup. References sequences are shown with GISAID (<https://www.gisaid.org>) or GenBank accession numbers. Red node points show strongly supported branches as detected by SH-aLRT and ultrafast bootstrap values. Scale bar indicates nucleotide substitutions per site. AA, amino acid; BA, BA.1/BA.2 recombinant; G, gene; M, membrane; N, nucleocapsid; ORF, open reading frame; P1, patient 1; P2, patient 2; S, spike; SNP, single-nucleotide polymorphism.

virus population. We used the patient 2 sample to clone an RT-PCR product (≈ 2.2 kbp) spanning the recombination breakpoint. We detected BA.1-specific (19955C/20055A) and BA.2-specific (21618T/21633–21641del/21762C) SNPs in the same plasmid clone, confirming the 2 patients were infected by a BA.1/BA.2 recombinant virus.

We found no similar BA.1/BA.2 recombinant sequences in GISAID or GenBank (as of March 7, 2022), suggesting a novel recombinant. The BA.1 region of this recombinant virus is genetically close to 3 BA.1 sequences detected in Europe and the United States (Figure, panel B), whereas its BA.2 region is identical to 19,555 BA.2 sequences from multiple continents. Because global cocirculation of BA.1 and BA.2 subvariants is high, pinpointing the geographic location where this recombination event occurred would be difficult.

Emerging Omicron subvariants could allow vaccine breakthrough and widespread reinfection. Previous studies reported detection of SARS-CoV-2 interlineage recombinants at the same time as different SARS-CoV-2 lineages were cocirculating (3; D. VanInsberghe et al., unpub. data, <https://doi.org/10.1101/2020.08.05.238386>; P. Colson et al., unpub. data; T. Peacock, unpub. data). The high transmissibility of Omicron (5,6) has led to wide cocirculation of BA.1 and BA.2 subvariants in many regions, which might provide ample opportunities to generate novel recombinants among these or other variants via coinfection events. Although current global surveillance data suggest that our recombinant might only be a sporadic case, the potential effects of novel recombinants should not be underestimated. Of note, homologous recombination is common in animal and other human coronaviruses (7), and some recombination events could generate recombinants with enhanced virulence (8,9). Long-term global SARS-CoV-2 genomic surveillance will be needed to monitor for possible more virulent or transmissible strains.

Acknowledgments

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About the Author

Dr. Gu is a postdoctoral fellow at The University of Hong Kong, Hong Kong, China. His research interests focus on bioinformatics and virus evolution.

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Appendix

Additional Methods

Sequencing

SARS-CoV-2 RT-PCR–positive samples with a C_t value <30 were randomly selected for next-generation sequencing (NGS) analysis. RNA samples were sent to a World Health Organization (WHO) reference laboratory at the University of Hong Kong (China) for full genome analyses (IRB no. UW 20–168). We deduced near full-length genomes from the samples using an Illumina (<https://www.illumina.com>) sequencing protocol previously described by us (1,2). Briefly, virus genome was reverse transcribed with multiple gene-specific primers targeting different regions of the viral genome. The synthesized cDNA was then subjected to multiple overlapping 2-kb PCRs for full-genome amplification. PCR amplicons obtained from the same specimen were pooled and sequenced using Novaseq or iSeq sequencing platform (Illumina). Specifically, for the sample from case-patient 1 with putative recombinant virus, we additionally performed NGS sequencing with the COVIDSeq kit (Illumina) for cross-validation. Generated sequencing reads were quality-trimmed by fastp (<https://github.com/OpenGene/fastp>) and mapped to a reference virus genome (Genbank accession no. MN908947.3) by BWA-MEM2 v2.1 (3). Potential PCR duplicates were identified and removed by samtools markdup (<https://www.htslib.org/doc/samtools-markdup.html>). The genome consensus was generated by iVar (4) with the PCR primer trimming protocol (minimum sequence depth of 5 for iSeq samples and minimum sequence depth of 10 for Novaseq samples, and minimum Q value of 30). The deduced sequences are available GISAID (<https://www.gisaid.org>; accession nos. are available at https://github.com/Leo-PoonLab/BA1_BA2_recombinant_HK/blob/main/GISAID_accessions.txt).

The average sequencing depths at the breakpoint region were 1,086 in patient 1 samples and 24,604 in patient 2 samples. We also cloned a \approx 2.2 kbp RT-PCR amplicon spanning the putative breakpoint region using patient 2's sample. The 5' and 3' end of this clone was subjected to standard Sanger sequencing.

Identification of Putative Recombinants

We scanned all the sequenced samples from imported cases in Hong Kong after November 15, 2021 for putative BA.1/BA.2 recombinants. The lineage defining mutations for BA.1 and BA.2 were curated from Cov-lineages (<https://github.com/cov-lineages/pango-designation/issues/361>) and CoVariants (<https://covariants.org>). For defining a sequence as a putative BA.1/BA.2 recombinant, it must have ≥ 3 BA.1- and BA.2-defining mutations, each with an allele frequency $>90\%$. The statistics of sample's read depth, allele frequency and minor allele frequency were deduced from aligned reads in bam files by using pysamstats (<https://github.com/alimanfoo/pysamstats>).

Identification of Putative Parental Sequences of the Recombinant

The available public 1,222,642 BA.1 and 767,399 BA.2 sequences from GISAID and GenBank (accessed on March 7, 2022) were downloaded and mapped to the reference sequence (GenBank accession no. MN908947.3) by using minimap2 (<https://github.com/lh3/minimap2>). The aligned sequences were used as the database for searching putative parental sequences. The leading/tailing partial sequences (positions 1–22005 and 21618–29903) of the recombinant genome were extracted by masking the remainder of the genome with “N”. Masking was also performed for the aligned public sequences with letter “?” by using figleaf (https://github.com/Koohoko/figleaf_fasta). After masking, low-quality sequences were dropped (if >300 “N” bases were found within the non-masked regions). The closest matches of the 2 partial recombinant sequences were separately identified from the above aligned public sequences using gofasta (<https://github.com/virus-evolution/gofasta>).

Phylogenetic Analysis

The consensus sequences deduced from NGS data were aligned to the reference genome (GenBank accession no. MN908947.3) by using MAFFT-add (<https://mafft.cbrc.jp/alignment/server/add.html>). The representative sequences from SARS-CoV-2 variants of concern Alpha, Beta, Gamma, Delta, and Omicron BA.3 were also included.

The 5' and 3' untranslated regions were masked before tree building. The maximum-likelihood phylogenies were estimated by using IQ-TREE version 2.1.3 (5), and the best-fit nucleotide substitution models searched by the software by using Wuhan-Hu-1 (GenBank accession no. MN908947.3) as the outgroup. Branch supports are assessed by SH-aLRT and the ultrafast bootstrap, a node is considered supported if SH-aLRT $\geq 80\%$ and UFboot $\geq 95\%$ (<http://www.iqtree.org/doc/Frequently-Asked-Questions>).

Simplot Analysis

The putative BA.1/BA.2 recombinant virus sequence was analyzed in Simplot v3.5.1 (6) for the recombination signals. Its similarity was plotted against a smaller group of representative variants of concern (VOCs) including Alpha, Beta, Gamma, Delta, Omicron BA.1, Omicron BA.2, and the prototype Wuhan/WH01/2019 (GenBank accession no. MN908947.3). Due to the relatively large proportion of strictly conserved sites, these sites were excluded from the alignment before subjecting to the similarity plot analysis. The BA.1/BA.2 recombinant, its putative parents from Omicron BA.1 and BA.2, and prototype Wuhan/WH01/2019 (as outgroup) were analyzed for their informative sites of recombination.

The GenBank/GISAID accession numbers of viral sequences used in the Simplot analysis are as follows:

Alpha: OL807059, OU272361, OU052790, OU179605, OL315388, OU208527, OU174622, OU208088, MW933836, MZ280980, MZ296197, MZ077208, OU022681

Beta: OU202380, OU516338, OU136527, OK433425, OM765676, OL779105, OU114765, MW963525, OU233168

Gamma: MW913237, OL803729, MZ414874, MZ217960, MZ536412, OV921949, MZ211976, OM485550, MZ037589

Delta: OK208965, OK258803, MZ988451, OK101403, OK243904, MZ764878, OK160402, OK054978, MZ888548, MZ888540, MZ888535, MZ888534, OU338538, EPI_ISL_8880068

BA.1: EPI_ISL_10273412

BA.1: EPI_ISL_10462716

Code Availability

Detailed analyzing scripts used in the study can be accessed in a GitHub repository (https://github.com/Leo-Poon-Lab/BA1_BA2_recombinant_HK).

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Appendix Table 1. Number and country of origin of studied imported COVID-19 cases (n = 198), Hong Kong, China, November 15, 2021–February 4, 2022

Country of importation	No. cases
United Kingdom	23
United States of America	22
Philippines	13
Nepal	12
Canada	10
Pakistan	9
Australia	7
Japan	7
Finland	6
France	6
India	6
Germany	5
Italy	5
Spain	5
Thailand	5
Ghana	4
Russia	4
Denmark	3
Ireland	3
Kenya	3
Switzerland	3
Vietnam	3
Brazil	2
Ethiopia	2
Kazakhstan	2
Korea	2
Nigeria	2
Poland	2
Singapore	2
South Africa	2
Sweden	2
Argentina	1
The Bahamas	1
Belgium	1
Chile	1
Cyprus	1
Czech Republic	1
Estonia	1
Lithuania	1
Morocco	1
The Netherlands	1
Papua New Guinea	1
Qatar	1
Republic of Moldova	1
Saudi Arabia	1
Ukraine	1
United Republic of Tanzania	1

Appendix Table 2. GISAID sequences used in a study of SARS-CoV-2 BA.1/BA.2 recombinant variant in arriving travelers, Hong Kong, China, February 2022*

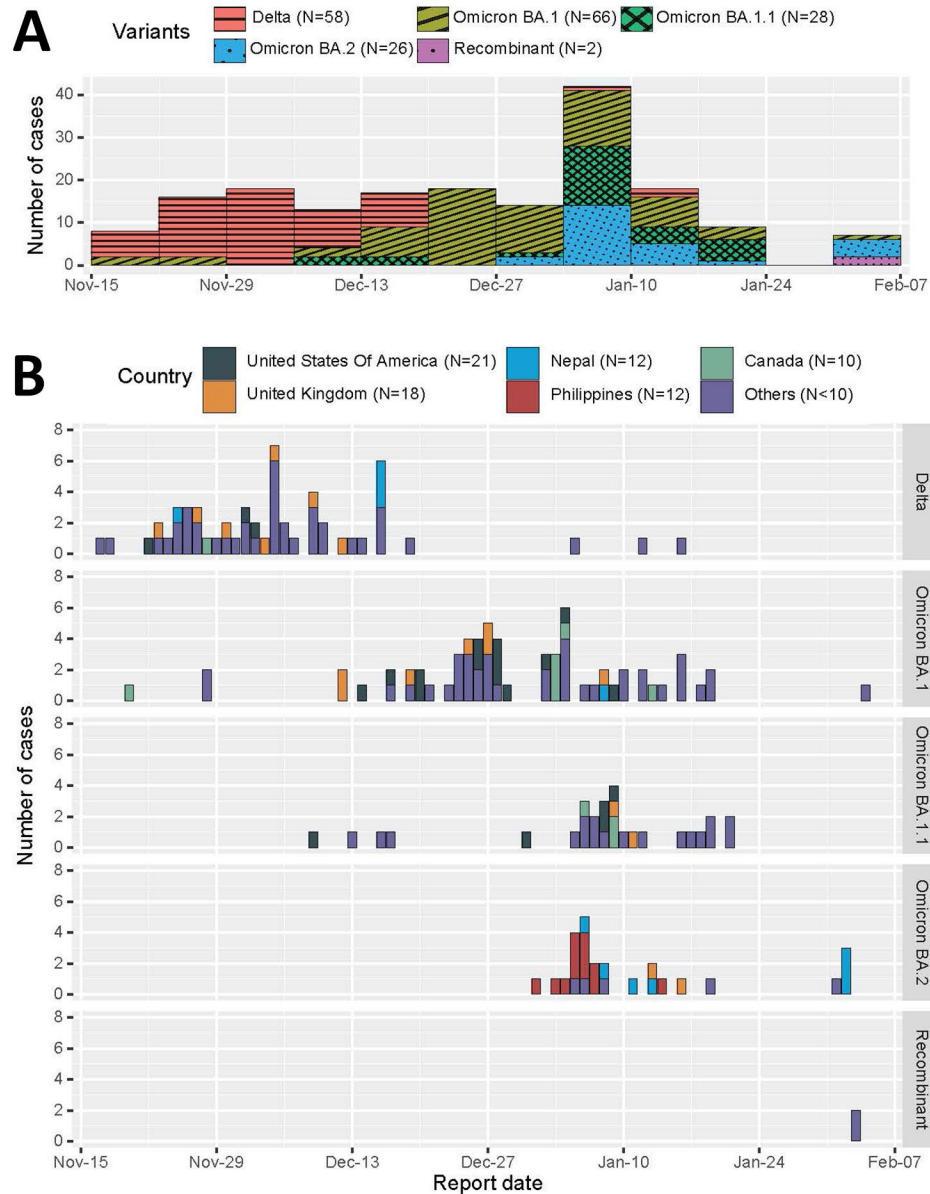
Accession no.	Originating laboratory	Submitting laboratory	Authors
EPI_ISL_1996858, EPI_ISL_2146766	Aegis Sciences Corporation	Centers for Disease Control and Prevention Division of Viral Diseases, Pathogen Discovery	Dakota Howard, Dhvani Batra, Peter W. Cook, Kara Moser, Adrian Paskey, Jason Caravas, Benjamin Rambo-Martin, Shatavia Morrison, Christopher Gulvick, Scott Sammons, Yvette Unoarumhi, Darlene Wagner, Matthew Schmerer, Cyndi Clark, Patrick Campbell, Rob Case, Vikramsinha Ghorpade, Holly Houdeshell, Ola Kvalvaag, Dillon Nall, Ethan Sanders, Alec Vest, Shaun Westlund, Matthew Hardison, Clinton R. Paden, Duncan MacCannell
EPI_ISL_3988521	Colorado Department of Public Health and Environment	Colorado Department of Public Health and Environment	Laura Bankers, Molly C. Hetherington-Rauth, Diana Ir, Alexandria Rossheim, Michael Martin, Mandy Waters, Shannon R. Matzinger, Sarah Elizabeth Totten, Emily A. Travanty
EPI_ISL_3029243	Department of Bacteria, Parasites and Fungi, Statens Serum Institut, Copenhagen, Denmark	Statens Serum Institut Bioinformatics and Microbial Genomics	Danish Covid-19 Genome Consortium
EPI_ISL_9519698	ESPACEBIO	Department of Virology, Henri Mondor University Hospital, Assistance Publique Hôpitaux de Paris, Université Paris-Est Créteil, INSERM U955	Christophe Rodriguez, Slim Fourati, Vanessa Demontant, Guillaume Gricourt, Melissa N'Debi, Alexandre Soulier, Elisabeth Trawinski, Jean-Michel Pawlotsky
EPI_ISL_1557096, EPI_ISL_3348121	Fulgent Genetics	Centers for Disease Control and Prevention Division of Viral Diseases, Pathogen Discovery	Dakota Howard, Dhvani Batra, Peter W. Cook, Kara Moser, Adrian Paskey, Jason Caravas, Benjamin Rambo-Martin, Shatavia Morrison, Christopher Gulvick, Scott Sammons, Yvette Unoarumhi, Darlene Wagner, Matthew Schmerer, Harry Gao, Mickey Li, John Gao, Joseph Fierro, Benafsh Sapra, Becky Tsai, Yan Meng, Doreen Ng, James Xie, Clinton R. Paden, Duncan MacCannell
EPI_ISL_4359169, EPI_ISL_4454918	Fulgent Genetics	Centers for Disease Control and Prevention Division of Viral Diseases, Pathogen Discovery	Dakota Howard, Dhvani Batra, Peter Cook, Jason Caravas, Benjamin Rambo-Martin, Scott Sammons, Yvette Unoarumhi, Matthew Schmerer, Kristine Lacey, Tymeckia Kendall, Victoria Caban Figueroa, Shatavia Morrison, Christopher Gulvick, Erisa Sula, Harry Gao, Mickey Li, John Gao, Joseph Fierro, Benafsh Sapra, Becky Tsai, Yan Meng, Doreen Ng, James Xie, Clinton Paden, Duncan MacCannell
EPI_ISL_4017432	Fulgent Genetics	Centers for Disease Control and Prevention Division of Viral Diseases, Pathogen Discovery	Dakota Howard, Dhvani Batra, Peter Cook, Kara Moser, Adrian Paskey, Jason Caravas, Benjamin Rambo-Martin, Shatavia Morrison, Christopher Gulvick, Scott Sammons, Yvette Unoarumhi, Darlene Wagner, Matthew Schmerer, Harry Gao, Mickey Li, John Gao, Joseph Fierro, Benafsh Sapra, Becky Tsai, Yan Meng, Doreen Ng, James Xie, Clinton Paden, Duncan MacCannell
EPI_ISL_2614026	Gravity Diagnostics, LLC	Gravity Diagnostics, LLC	Gravity Diagnostics
EPI_ISL_9879325	Hospital	National Reference Center for Viruses of Respiratory Infections, Institut Pasteur, Paris	Marion Barbet, Sylvie Behillil, Méline Bizard, Angela Brisebarre, Camille Capel, Vincent Enouf, Louise Lefrançois, Frédéric Lemoine, Christophe Malabat, Corinne Maufrais, Slim El Khiari, Julien Fumey, Etienne Simon-Lorière, Maud Vanpeene, Sylvie Van der Werf, Benedicte LUREAU
EPI_ISL_2282971	Infinity Biologix	Centers for Disease Control and Prevention Division of Viral Diseases, Pathogen Discovery	Dakota Howard, Dhvani Batra, Peter W. Cook, Kara Moser, Adrian Paskey, Jason Caravas, Benjamin Rambo-Martin, Shatavia Morrison, Christopher Gulvick, Scott Sammons, Yvette Unoarumhi, Darlene Wagner, Matthew Schmerer, Christian Bixby, Yihe Wang, Jonathan Schultz, Chirayu Goswami, Russ Hager, Robin Grimwood, Clinton R. Paden, Duncan MacCannell

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EPI_ISL_1682200	Laboratory Corporation of America	Centers for Disease Control and Prevention, Division of Viral Diseases, Pathogen Discovery	Dakota Howard, Dhvani Batra, Peter W. Cook, Kara Moser, Adrian Paskey, Jason Caravas, Benjamin Rambo-Martin, Shatavia Morrison, Christopher Gulvick, Scott Sammons, Yvette Unoarumhi, Darlene Wagner, Matthew Schmerer, Minoo Agarwal, Eyad Almasri, Debbie Boles, Ayla Burns, Nuthawin Charoensri, Oren Cohen, Susan Countryman, Mary Ann Cristobal, Bobbi Croy, Suzanne Dale, Hrushikesh Deshmukh, Amanda Douglas, Vincent Drouillon, Marcia Eisenberg, Howard Engler, Rama Ghatti, Prashant Gupta, Susan Hicks, Jake Humphrey, Lax Iyer, Manoj Jain, Mohan Kolli, Brian Krueger, Tim Kuphal, Stanley Letovsky, Michael Levandoski, Craig Lukasik, Jonathan Meltzer, Brian Norvell, Mindy Nye, Scott Parker, Christos Petropoulos, John Pruitt, Steven Ragan, Scott Ryan, Mike Sapeta, Jana Schroth, Suresh Babu Selvaraju, Goran Stevovic, Amanda Suchanek, Andrea Throop, Lyndon Tilson, Thomas Urban, Joe Voshell, Kimberly Wagner, Jonathan Williams, Mary Williamson, Qian Zeng, Tricia Zwiefelhofer, Clinton R. Paden, Duncan MacCannell
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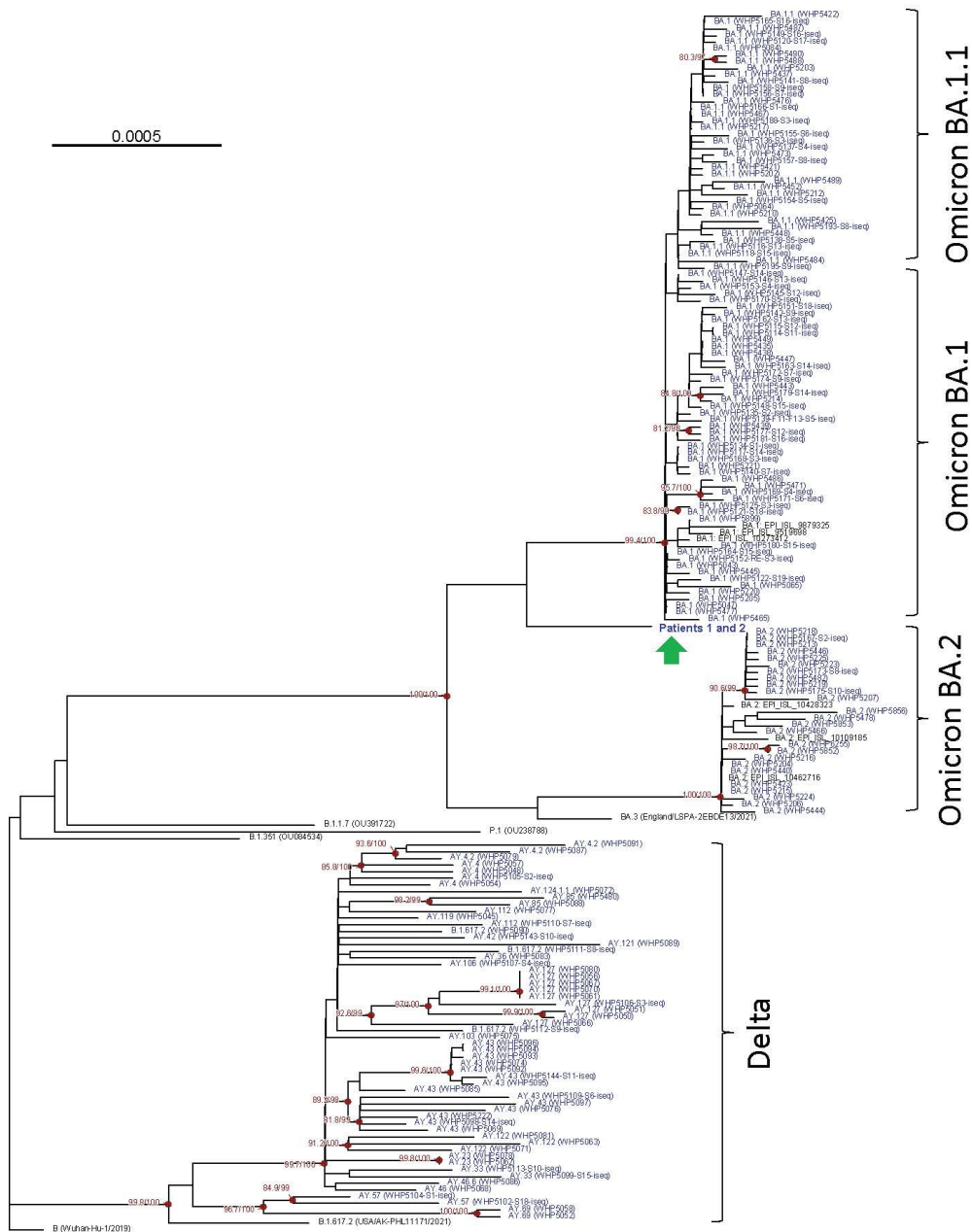
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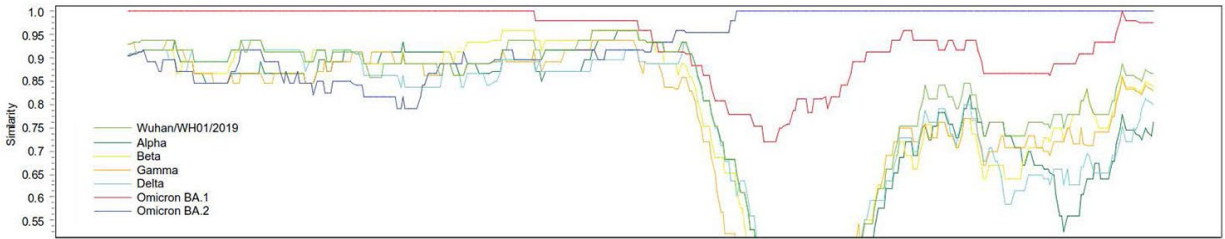
*We gratefully acknowledge the authors from the originating laboratories responsible for obtaining the specimens and the submitting laboratories where genetic sequence data were generated and shared via the GISAID Initiative, on which this research is based.



Appendix Figure 1. Importation of SARS-CoV-2 variants from incoming travelers, Hong Kong, China, November 15, 2021–February 7, 2022. A) Time series of number of patients testing positive for different SARS-CoV-2 variants by RT-PCR. B) Time series divided by SARS-CoV-2 variants, and country of origin. All infections were confirmed by whole-genome sequencing. RT-PCR, reverse transcription PCR.



Appendix Figure 2. Phylogeny of SARS-CoV-2 variants identified in incoming travelers, Hong Kong, China. The maximum-likelihood phylogenetic tree was generated by using IQ-TREE (<http://www.iqtree.org>) and the GTR+F+I nucleotide substitution model with Wuhan-Hu-1 (GenBank accession no. MN908947.3) as the outgroup. Blue text indicates viral genomes generated from this study and references sequences used in the analysis are shown as indicated. Arrow indicates the recombinant virus detected from patients 1 and 2. Red node points show strongly supported branches by SH-aLRT/ultrafast bootstrap. Scale bar indicates estimated nucleotide substitutions per site.



Appendix Figure 3. Simplot analysis of recombinant BA.1/BA.2 SARS-CoV-2 virus, Hong Kong, China, February 2022. Plot shows similarity between full viral genomes of different variants of concern including Alpha, Beta, Gamma, Delta, Omicron BA.1, Omicron BA.2, as well as the prototype Wuhan/WH01/2019 were used in the analysis.