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Case Study: Prolonged Infectious SARS-CoV-2 Shedding from an Asymptomatic Immunocompromised Individual with Cancer

Highlights
- Persistent SARS-CoV-2 infection and shedding in immunocompromised individual
- Infectious SARS-CoV-2 isolated up to 70 days after diagnosis
- Observed within-host genetic variation with continuous turnover of viral variants
- SARS-CoV-2 isolates from the individual do not display altered replication

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In Brief
This case study describes a female immunocompromised individual with chronic lymphocytic leukemia and acquired hypogammaglobulinemia who became persistently infected with SARS-CoV-2. Although asymptomatic throughout the course of infection, she demonstrated prolonged shedding of infectious SARS-CoV-2 virus and RNA. This study demonstrates that certain individuals may remain infectious for prolonged periods of time and highlights the need for further studies to understand risk factors for prolonged infectious SARS-CoV-2 shedding.
SUMMARY

Long-term severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) shedding was observed from the upper respiratory tract of a female immunocompromised individual with chronic lymphocytic leukemia and acquired hypogammaglobulinemia. Shedding of infectious SARS-CoV-2 was observed up to 70 days, and of genomic and subgenomic RNA up to 105 days, after initial diagnosis. The infection was not cleared after the first treatment with convalescent plasma, suggesting a limited effect on SARS-CoV-2 in the upper respiratory tract of this individual. Several weeks after a second convalescent plasma transfusion, SARS-CoV-2 RNA was no longer detected. We observed marked within-host genomic evolution of SARS-CoV-2 with continuous turnover of dominant viral variants. However, replication kinetics in Vero E6 cells and primary human alveolar epithelial tissues were not affected. Our data indicate that certain immunocompromised individuals may shed infectious virus longer than previously recognized. Detection of subgenomic RNA is recommended in persistently SARS-CoV-2-positive individuals as a proxy for shedding of infectious virus.

INTRODUCTION

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) RNA can be detected at various sites, including samples obtained from the nares, nasopharynx, pharynx, bronchoalveolar lavage (BAL) fluid, feces, and blood (Wang et al., 2020a; Sun et al., 2020; Judson and Munster, 2020). The duration of SARS-CoV-2 RNA shedding is generally between 3 and 46 days after symptom onset (Fu et al., 2020; Qian et al., 2020; Liu et al., 2020c). Asymptomatic individuals shed SARS-CoV-2 RNA comparably with symptomatic individuals regarding duration and viral load (Lee et al., 2020; Long et al., 2020; Zou et al., 2020). Persistent SARS-CoV-2 RNA shedding has been documented, with patients remaining qRT-PCR-positive for up to 63 days (Li et al., 2020; Liu et al., 2020b). In addition, there are reports of symptomatic and asymptomatic individuals testing positive again after a period of negative testing (Lan et al., 2020; Hu et al., 2020). Because qRT-PCR detects viral RNA but does not confirm the presence of infectious SARS-CoV-2, these observations raise questions about the duration of infectious SARS-CoV-2 shedding and transmission potential for symptomatic and asymptomatic individuals. Estimates suggest that infectiousness begins 2.3 days prior to symptom onset and declines within 7 days of symptom onset (He et al., 2020b). Consistent with this, infectious SARS-CoV-2 has been isolated from patient samples taken up to 8 days after symptom onset but typically not thereafter (Wölfel et al., 2020; Bullard et al., 2020). In contrast to prolonged shedding of SARS-CoV-2 RNA, the longest detected shedding of infectious SARS-CoV-2 virus is up to 20 days after the initial positive test result (van Kampen et al., 2020; Liu et al., 2020b). The probability of isolating SARS-CoV-2 decreases with a lower viral load, when the duration of symptoms exceeds 15 days, and upon generation of detectable neutralizing antibodies (van Kampen et al., 2020).

On January 19, 2020, the first case of coronavirus disease 2019 (COVID-19) was identified in the United States, in Snohomish County, Washington, in a traveler returning from Wuhan,
China. Community spread in the Seattle region became evident in late February of 2020 (Bhatraju et al., 2020), with extensive spread in a long-term care facility (McMichael et al., 2020a). Here we describe an asymptomatic, immunocompromised individual persistently testing positive for SARS-CoV-2 by qRT-PCR who was infected during the early phase of SARS-CoV-2 spread in the United States. Infectious SARS-CoV-2 was successfully isolated from nasopharyngeal swabs 49 days and 70 days after the initial positive qRT-PCR test. Convalescent plasma treatment was not immediately successful in clearing the infection, but evidence of SARS-CoV-2 RNA was eventually cleared after 105 days.

RESULTS

Clinical Presentation of an Immunocompromised Individual Persistently Infected with SARS-CoV-2

On February 12, 2020, a 71-year-old woman with a 10-year history of chronic lymphocytic leukemia (CLL), acquired hypogammaglobulinemia, anemia, and chronic leukocytosis presented to the emergency department with low back and lower extremity pain. She underwent surgery for a spinal fracture and stenosis related to her cancer on February 14, 2020 (biopsy results in Table S1) and was subsequently transferred to a rehabilitation facility on February 19, 2020. On February 25, 2020, she was re-hospitalized for anemia and underwent a chest X-ray the following day, which was normal. She could not return to her rehabilitation center because of a confirmed outbreak of COVID-19 at the facility (McMichael et al., 2020a, 2020b). Chest computed tomography (CT), performed on February 28, 2020, was unremarkable. The patient had no respiratory or systemic symptoms during this time. Because she was residing in the rehabilitation facility around the time of the COVID-19 outbreak, she was tested and found positive for SARS-CoV-2 on March 2, 2020 (Figure 1). After the initial SARS-CoV-2 diagnosis, she was kept in an isolation ward in a single room with negative airflow. Attending medical staff were using full personal protective equipment comprised of powered air-purifying respirators (PAPR) or N95 respirators with goggles, gowns, and gloves. Over the course of the next 15 weeks, she was tested another 14 times by several diagnostic companies and remained positive through June 15, 2020, 105 days since the initial positive test. Subsequently, she tested negative on four consecutive swabs from June 16 to July 16, indicating that her infection had cleared.

Because of acquired hypogammaglobulinemia caused by her CLL, the individual received intravenous immunoglobulin (IVIG) every 4–6 weeks as part of her treatment regimen. She received IVIG treatment on April 6 and May 6, 2020. The manufacture date of her specific lot of IVIG preceded January 1, 2020, the beginning of the COVID-19 pandemic, and therefore did not contribute to any SARS-CoV-2 serology results (Table S2). Because of the persistence of her SARS-CoV-2 infection, serum samples were tested for antibodies against the spike glycoprotein through a study at the NIH Clinical Center, and no spike-specific antibodies were detected (Burbelo et al., 2020). On May 12, 2020, she was transfused with 200 mL of SARS-CoV-2 convalescent plasma provided by Bloodworks Northwest under a US Food and Drug Administration (FDA) emergency investigational new drug
In an attempt to treat the persistent SARS-CoV-2 infection, the individual received two doses of convalescent plasma therapy on days 71 and 82. Pre- and post-transfusion serum samples and the transfusion convalescent plasma samples were analyzed for the presence of full-length spike and spike receptor binding domain (RBD) antibodies by ELISA assay and of SARS-CoV-2-neutralizing antibodies in a VN assay (Figures 2B and 2C; Figures S1A and S1B; Table 1; Amanat et al., 2020; Wrapp et al., 2020). The first dose of convalescent plasma (convalescent plasma 1) had an immunoglobulin G (IgG) spike titer of 2,560, RBD titer of 3,840, and VN titer of 60. The second dose of convalescent plasma (convalescent plasma 2) had an IgG spike titer of 5,120, RBD titer of 5,120, and VN titer of 160 (Figure 2B; Figure S1A; Table 1). Prior to the first dose of plasma given on day 71, detectable spike and RBD IgG antibody titers were very low in serum collected from the individual, with IgG titers between 1:10 and 1:40 on days 49 and 71 pre transfusion; no VN titers were detected in these samples. Immediately after the first transfusion on day 71, the spike and RBD IgG antibody titers rose to 1:320 and then decreased to 1:80 and 1:160, respectively, on day 77. No VN titers were detected on days 71 and 77 (Figure 2C; Figure S1B; Table 1). Immediately after the second transfusion on day 82, the spike and RBD IgG titers increased to 1:320 and 1:640, respectively, and remained elevated by day 105 (Figure 2C; Figure S1B). Low neutralizing titers of 1:10 were observed on day 82 and 105 (Table 1).

Despite two transfusions of convalescent plasma, nasopharyngeal swabs on days 85 and 105 remained positive for gRNA and sgRNA, suggesting that the convalescent plasma therapy was not successful in rapidly clearing the infection from the upper respiratory tract in this individual. Although the presence of sgRNA at these time points suggests active viral replication, infectious SARS-CoV-2 could not be cultured after day 70.

### Table 1. Virus Neutralization Titers in Pre- and Post-transfusion Sera from the Individual and Convalescent Plasma Used for Transfusion

| Serum                        | USA/WA1/2020 | Day 49 | Day 70 |
|------------------------------|--------------|--------|--------|
| Day 49                       | <10          | <10    | <10    |
| Day 71                       | <10          | <10    | <10    |
| Day 71 after transfusion      | <10          | <10    | <10    |
| Day 77                       | <10          | <10    | <10    |
| Day 82                       | <10          | 10     | <10    |
| Day 82 after transfusion      | 10           | 10     | 15     |
| Day 105                      | 10           | 10     | <10    |
| Convalescent plasma 1        | 60           | 40     | 40     |
| Convalescent plasma 2        | 160          | 160    | 60     |

Virus neutralization assays were performed for all serum and plasma samples with SARS-CoV-2 strains USA/WA1/2020 and the day 49 and day 70 isolates from the individual. Each serum/plasma sample was tested in duplicate.

Long-Term Shedding of Genomic RNA, Subgenomic RNA, and Infectious SARS-CoV-2

SARS-CoV-2 shedding kinetics in the individual were monitored using detection of genomic RNA (gRNA), subgenomic RNA (sgRNA), and infectious SARS-CoV-2. RNA was extracted from nasopharyngeal and oropharyngeal swabs collected 49, 70, 77, 85, 105, and 136 days after the initial diagnosis and evaluated for the presence of viral gRNA (Corman et al., 2020) and sgRNA (Wöffel et al., 2020). gRNA and sgRNA were detected in nasopharyngeal swabs out to day 105, except for the swab taken on day 77 (Figure 2A), although the test through EvergreenHealth was positive at this time. None of the oropharyngeal swabs were positive for gRNA or sgRNA, suggesting that the infection was confined to the nasopharynx. Absolute quantification of gRNA and sgRNA on positive swabs was performed by droplet digital PCR (ddPCR) (Figure 2A). The highest viral load was detected in the day 70 swab, at $2.2 \times 10^6$ gRNA copies/mL (cycle threshold [Ct] 22.44) and $1.1 \times 10^5$ sgRNA copies/mL (Ct 29.05). Detection of sgRNA in swabs is indicative of active SARS-CoV-2 replication because only actively replicating SARS-CoV-2 initiates RNA synthesis, resulting in replication and transcription of sgRNAs (Wang et al., 2020b; Kim et al., 2020), and sgRNA, unlike gRNA, does not persist in the nasal cavity in the absence of virus replication (Speranza et al., 2020). Virus isolation was attempted on all qRT-PCR-positive samples. Infectious SARS-CoV-2 was successfully cultured from the nasopharyngeal swabs collected on day 49 and day 70. Scanning and transmission electron microscopy on SARS-CoV-2 cultured from the nasopharyngeal swabs collected on days 49 and 70 showed viral particles consistent with coronavirus morphology, supporting persistent SARS-CoV-2 infection with shedding of infectious virus in this individual (Figure 3).

Genetic Analysis of Patient Swab Samples Links Infection to the Primary Washington State Outbreak

SARS-CoV-2 full genome sequences were obtained from nasopharyngeal swabs collected on days 49, 70, 85, and 105 (Table S4). Full genomes were obtained by sequencing using the ARTIC primer set (https://artic.network) and assembling reads to MN985325.1 (USA/WA1/2020) as the reference genome (Harcourt et al., 2020). The SARS-CoV-2 lineage was determined using Pangolin software (https://pangolin.cog-uk.io/), which placed the individual’s viral genomes in lineage A.1, which consists of genomes originating from the primary outbreak in Washington state (Rambaut et al., 2020). A maximum-likelihood tree was generated using representative SARS-CoV-2 genomes from previously described lineages (Rambaut et al., 2020) obtained from the GISAID database (https://www.gisaid.org/; Shu and McCauley, 2017). The individual’s SARS-CoV-2 full-length genomes cluster together within lineage A.1 (Figure 4A). This suggests that she was infected with a virus from the SARS-CoV-2 A.1 lineage, which circulated after the initial import from China, followed by exponential growth and local transmission in Washington state.
To visualize the temporal relationships of the patient isolates, 44 full SARS-CoV-2 genome sequences from Washington state belonging to NextStrain clade 19B (http://clades.nextstrain.org/) were subsampled from the GISAID database (https://www.gisaid.org/; Shu and McCauley, 2017) representing strains collected in Washington state from February to May 2020. A full genome alignment was performed with four of the full genome sequences recovered from the persistently infected individual, the USA/WA1/2020 genome sequence, and the Wuhan-Hu-1/2019 genome sequence with MAFFT v.1.4 (Katoh and Standley, 2013; Katoh et al., 2002) implemented in Geneious Prime v.2020.1.2 (https://www.geneious.com/). A maximum-likelihood tree was reconstructed with PhyML v.3.1 (Guindon et al., 2010), and a tree showing temporal divergence (Figure 4B) was inferred in TreeTime v.0.7.6 (Sagulenko et al., 2018; Hadfield et al., 2018) using the HKY85 model of nucleotide substitution and a fixed molecular clock at 8e−4 with a standard deviation of 4e−4, as implemented in the NextStrain pipeline (https://nextstrain.org/sars-cov-2). Divergence dating estimates place the patient isolates sharing a most recent common ancestor between February 27 and March 31, 2020, within 90% of the marginal probability distribution. This is consistent with the timing of the individual’s first positive test on March 2, 2020. To further evaluate the relationship between the SARS-CoV-2 genomes recovered from the patient swabs and other SARS-CoV-2 genomes circulating in Washington state at the times of sampling (April 20, May 11, May 26, and June 15, 2020), Washington SARS-CoV-2 genomes were downloaded from the GISAID database (Shu and McCauley, 2017). The quality of the sequences was determined by Nextclade v.0.7.5 (https://nextstrain.org/ngcov/global), and 1,789 sequences on April 20, 385 sequences between April 20 and May 11, 268 sequences between May 11 and May 26, and 709 sequences between May 26 and June 15 were kept for further phylogenetic analysis. Maximum likelihood trees using the curated sets of sequences, the four patient genomes, and the USA/WA1/2020 genome were inferred using ModelFinder (Kalyaanamoorthy et al., 2017) and ultrafast bootstrap (Hoang et al., 2018) implemented in IQ-TREE v.1.6.12 (Nguyen et al., 2015). The phylogenetic trees show that the patient genomes in this study cluster as a monophyletic clade consistent with infection in late February/early March, followed by virus persistence (Figure S2).

Next, full genome sequences from the two SARS-CoV-2 isolates were obtained (Table S4), and the consensus level variants in the sequences obtained from nasopharyngeal swabs and SARS-CoV-2 isolates cultured from those swabs were
compared with the reference strain USA/WA1/2020 (MN985325.1) (Harcourt et al., 2020). Several single-nucleotide (nt) substitutions were observed within the ORF1ab, spike, M, and ORF8 coding sequence in the full-genome sequences obtained directly from the individual's swabs and the SARS-CoV-2 isolates. In addition, a 3-nt deletion leading to loss of a methionine residue was observed in nsp1 in day 49 and day 70 samples (Table 2). Within the genomes of the two SARS-CoV-2 isolates, two in-frame deletions were observed in the spike glycoprotein coding region. A 21-nt in-frame deletion (residues 21,975–21,995) was found in the N-terminal domain (NTD) of S1, leading to a 7-amino-acid deletion (aa 139–145) in the spike glycoprotein of the SARS-CoV-2 day 49 isolate. A smaller, 12-nt deletion (residues 21,982–21,993) was detected in the day 70 isolate, leading to a 4-aa deletion (aa 141–144) in the NTD, which falls within the 7-aa deletion found in the day 49 isolate (Figure 5A). These observed deletions in the spike glycoprotein map to a region in the NTD that is partially solvent exposed and forms a β strand in a compact conformation of the spike (Wrobel et al., 2020; Figures 5B and 5C). This region is unmodelled in other structures representing additional conformational states of the spike and, thus, is likely flexible (Wrapp et al., 2020; Walls et al., 2020). It is possible that the apparent plasticity in this region of the molecule may contribute to the structural permissibility of the identified deletions. The position of these deletions is distinct from those observed in other SARS-CoV-2 isolates, which locate to the S1/S2 and S2’ cleavage sites (Andrés et al., 2020; Lau et al., 2020; Liu et al., 2020).

Comparison of the full genome sequences obtained directly from the individual’s samples with the genome data obtained from the two SARS-CoV-2 isolates showed that the 21-nt deletion was present in a minority of sequencing reads (1%) in the genome obtained from the individual’s sample from day 49 (Table 2) and was selected for upon passage in cell culture. The 12-nt deletion on day 70 was present in 100% of the reads in the clinical sample and tissue culture isolate. Notably, neither spike deletion was detected in the genome sequences from the day 85 and day 105 swabs (Table 2). It is possible that other minor variants exist at low levels that were undetected by the depth of sequencing coverage or were not reflected in the sampling at that time point. The variation observed between the different full-length genomes obtained at various time points during the course of infection points to a quasispecies complex with continuous turnover of dominant viral species.

**Growth Kinetics of SARS-CoV-2 Patient Isolates**

The replication kinetics of the day 49 isolate SARS-CoV-2 were compared with those of the reference strain USA/WA1/2020 in Vero E6 cells. Despite the observed mutations in the day 49 isolate, no difference in replication kinetics were observed between the day 49 isolate and the reference strain (Figure 6A). To determine growth kinetics in a more functionally relevant cell type, growth curves were also performed on primary human alveolar epithelial tissues (EpiAlveolar; MatTek, Ashland, MA, USA). No significant differences were observed between the individual’s isolate and the reference strain in these cells (Figure 6B).

**DISCUSSION**

In this report, we describe long-term SARS-CoV-2 shedding in an immunocompromised individual with CLL and acquired hypogammaglobulinemia out to 105 days after the initial positive test. Although the exact time point when the individual acquired SARS-CoV-2 is unknown, it is likely that the exposure occurred in the long-term care facility where she resided between February 19–25, 2020, shortly before a large COVID-19 outbreak was identified in that facility on February 28, 2020. The individual remained asymptomatic throughout the course of infection despite isolation of infectious SARS-CoV-2 49 and 70 days after the initial diagnosis, much longer than shedding of infectious virus up to day 20, as reported previously (van Kampen et al., 2020). The information available to date about SARS-CoV-2 infection in immunocompromised individuals, including those with cancers such as CLL, is limited and mostly focuses on disease severity and outcome (He et al., 2020a; Paneesha et al., 2020).
Although it is difficult to extrapolate from a single individual, our data suggest that long-term shedding of infectious virus may be a concern in certain immunocompromised people. Given that immunocompromised individuals could have prolonged shedding and may not have typical symptoms of COVID-19, symptom-based strategies for testing and discontinuing transmission-based precautions, as recommended by the Centers for Disease Control and Prevention (CDC) (CDC, 2020b), may fail to detect whether certain individuals are shedding infectious virus.

The individual eventually cleared the SARS-CoV-2 infection from the upper respiratory tract after developing low neutralizing antibody titers. How the virus was cleared and the effect of convalescent plasma on clearance of the virus is unknown. The initial administration of convalescent plasma was followed by a decreased viral load in nasal swabs, but viral loads subsequently increased, despite administration of a second dose of convalescent plasma comprising higher antibody titers. Therapeutic administration of convalescent plasma is focused on treatment of severe or life-threatening COVID-19. Several clinical trials are investigating the efficacy of convalescent plasma, but currently the effect of convalescent plasma therapy on COVID-19 outcome remains equivocal (Mira et al., 2020; Salazar et al., 2020). The limited effect of convalescent plasma treatment on clearance of SARS-CoV-2 could be due to the fact that intravenously (i.v.) administered antibodies do not distribute well to the nasal epithelium (Ikegami et al., 2020) compared with the lower respiratory tract (Mira et al., 2020).

Throughout the course of infection, there was marked within-host genomic evolution of SARS-CoV-2. Deep sequencing revealed a continuously changing virus population structure with turnover in the relative frequency of the observed genotypes over the course of infection. With SARS-CoV-2, there is generally relatively limited within-host variation reported, and over the course of infection, the major SARS-CoV-2 population remains identical (Jary et al., 2020; Shen et al., 2020; Capobianchi et al., 2020). Potential factors contributing to the observed within-host evolution is prolonged infection and the compromised immune status of the host, possibly resulting in a different set of selective pressures compared with an immune-competent host. These differential selective pressures may have allowed a larger genetic diversity with continuous turnover of dominant viral species throughout the course of infection. Although some sequence variants remain consistent throughout the duration of infection, we also observed variants unique to individual time points, such as the spike deletions observed on day 49 and day 70. Previously reported spike deletions, distinct from those reported here, were observed at relatively low frequency in clinical samples but were enriched upon virus isolation (Andrés et al., 2020; Liu et al., 2020c). Similar to these reports, the spike
deletion in the isolate on day 49 was observed as a minor variant in the individual’s sample but was also selected for during passage upon virus isolation.

In contrast to the previously reported deletions at the cleavage sites, both spike deletions observed on day 49 and 70 in the individual are located in the NTD of S1, a region distal from the receptor binding site. These deleted residues are not modeled in a number of spike structures (Wrapp et al., 2020; Walls et al., 2020), suggesting that this region is conformationally labile. Although the NTD has been identified as an antigenic target (Brouwer et al., 2020; Chi et al., 2020; Liu et al., 2020a), no clear difference in virus neutralization was observed between the two patient isolates and the prototype USA/WA1/2020 SARS-CoV-2 isolate.

Despite genetic changes in the SARS-CoV-2 isolated from the individual, the replication kinetics did not change significantly compared with the USA/WA1/2020 virus in Vero E6 cells and primary human alveolar epithelial tissues. This indicates that, most likely, the infectious virus shed by the individual would still be able to establish productive infection in contacts upon transmission, assuming that viral growth kinetics in vitro are a suitable surrogate for virus fitness in vivo. Moreover, despite prolonged replication exclusively in the upper respiratory tract, the virus was still able to replicate in epithelial cells derived from the lower respiratory tract, suggesting that it could still cause pneumonia.

Many current infection control guidelines assume that persistent PCR-positive individuals are shedding residual RNA and not infectious virus, with immunocompromised people thought to remain infectious for no longer than 20 days after symptom onset (CDC, 2020a). Here we show that certain individuals may shed infectious, replication-competent virus for much longer than previously recognized (van Kampen et al., 2020). Although infectious virus could be detected up to day 70, sgRNA, a molecular marker for active SARS-CoV-2 replication (Speranza et al., 2020), could be detected up until day 105. An immunocompromised state has been identified as a risk factor for development of severe disease and complications from COVID-19 (CDC, 2020b). A wide variety of conditions and treatments can alter the immune system and cause immunodeficiency, creating opportunities for prolonged viral replication and shedding of infectious SARS-CoV-2. Although this report focuses on long-term shedding of one immunocompromised individual, an estimated 3 million people in the United States have some form of immunocompromising condition, including individuals with HIV infection, solid organ transplant recipients, hematopoietic stem cell transplant recipients, and individuals receiving chemotherapy and corticosteroids (Kunisaki and Janoff, 2009). This transient or chronic immunocompromised population is at higher risk of respiratory disease complications with respiratory infections such as influenza A virus and SARS-CoV-2 (Kunisaki and Janoff, 2009). Prolonged shedding of pH1N1 shedding was observed in immunocompromised individuals with a variety of immunocompromising conditions during the previous pandemic in 2009, such as people with cancer on chemotherapy and solid organ transplant recipients (van der Vries et al., 2013). For the SARS-CoV-2 related Middle East respiratory syndrome CoV (MERS-CoV), prolonged shedding up to 38 days was observed in individuals with myelodysplastic syndrome, autologous peripheral blood stem cell transplantation for treatment of large B cell lymphoma, and an individual with peripheral T cell lymphoma (Kim et al., 2017). MERS-CoV shedding was higher and longer in experimentally
infected non-human primates immunosuppressed with cyclophosphamide and dexamethasone, providing experimental support for the effect of immunosuppression on virus-host dynamics observed here (Prescott et al., 2018).

**Limitations of Study**
A limitation of the present study is that it comprises only a single case, making it difficult to draw general conclusions regarding use of convalescent plasma for clearance of the virus, potential alternative mechanisms involved in virus clearance, and the frequency of persistent SARS-CoV-2 infection and shedding in individuals with other immunocompromising conditions. Identification of additional cases of persistent infection and long-term shedding of infectious virus are needed so the infection dynamics can be studied in more detail in this diverse population.

Understanding the mechanism of virus persistence and eventual clearance will be essential for providing appropriate treatment and preventing transmission of SARS-CoV-2 because persistent infection and prolonged shedding of infectious SARS-CoV-2 might occur more frequently. Because immunocompromised individuals are often cohorted in hospital settings, a more nuanced approach to testing these individuals is warranted, and the presence of persistently positive people by performing SARS-CoV-2 gRNA and sgRNA analyses on clinical samples should be investigated.

**STAR★METHODS**
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SUPPLEMENTAL INFORMATION

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**AUTHOR CONTRIBUTIONS**

Conceptualization, F.X.R. and V.J.M.; Resources, E.R.F., C.M., and F.X.R.; Methodology, V.A.A., M.J.M., S.N.S., B.N.W., E.R.F., C.M., and T.A.B., and E.D.W.; Investigation, V.A.A., M.J.M., S.N.S., R.P., B.N.W., S.L.A., K.B., E.R.F., and E.D.W.; Writing – Original Draft, V.A.A., M.J.M., F.X.R., and V.J.M; Writing – Review & Editing, S.N.S., S.D.J., C.M., T.A.B., and E.D.W.; Data Curation, C.M.; Supervision, T.A.B., E.d.W., F.X.R., and V.J.M.

**DECLARATION OF INTERESTS**

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### Key Resources Table

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| Rabbit anti-Human IgG Fc fragment Secondary Antibody (HRP) | Novus biologicals | Cat# NBP1-73529; Lot 34900 |
| **Bacterial and Virus Strains** |        |            |
| hCoV-19/USA/WA1/2020 | CDC, Atlanta, USA | GenBank: MN985325.1 |
| hCoV-19/USA/WA-RML-5/2020 (SARS-CoV-2 patient genome d49 isolate) | This paper | GenBank: MT982401 |
| hCoV-19/USA/WA-RML-6/2020 (SARS-CoV-2 patient genome d70 isolate) | This paper | GenBank: MT982404 |
| **Biological Samples** |        |            |
| Patient nasopharyngeal swabs | This paper | EvergreenHealth |
| Patient oropharyngeal swabs | This paper | EvergreenHealth |
| Patient serum and plasma | This paper | EvergreenHealth |
| Donor convalescent plasma | This paper | EvergreenHealth |
| **Chemicals, Peptides, and Recombinant Proteins** |        |            |
| SARS-CoV-2 spike protein | This paper | N/A |
| (Wrapp et al., 2020) |        |            |
| SARS-CoV-2 receptor binding domain protein | This paper | N/A |
| (Amanat et al., 2020) |        |            |
| PEI transfection reagent | Polysciences | Cat# 23966-1 |
| Blocker Casein in PBS | ThermoFisher | Cat# 37528 |
| TMB 2-Component Microwell Peroxidase Substrate Kit | SeraCare | Cat# 5120-0047 |
| KPL TMB Stop Solution | SeraCare | Cat# 5150-0020 |
| Trizol Reagent | Invitrogen | Cat# 15596026 |
| Karnovsky’s EM fixative | Electron Microscopy Sciences | Cat#15720 |
| Sodium Cacodylate | Sigma | Cat#C4945-10G; CAS#6131-99-3 |
| Osmium Tetroxide | Electron Microscopy Sciences | Cat#19190; CAS#20816-12-0 |
| Potassium Ferrocyanide | Sigma | Cat#P-3289; CAS#14459-95-1 |
| Uranyl Acetate | Ted Pella | Cat#19481; CAS#6159-44-0 |
| **Critical Commercial Assays** |        |            |
| PureLink RNA Mini Kit | Invitrogen | Cat# 12183018A |
| SuperScript IV First-Strand Synthesis System | Invitrogen | Cat# 18091050 |
| Quantifast Probe RT-PCR Kit (for Rotorgene) | QIAGEN | Cat# 204556 |
| ddPCR Supermix for Probes (no dUTP) | Biorad | Cat# 1863024 |
| Q5 Hot Start High-Fidelity DNA Polymerase – 100U | New England Biolabs | Cat#M0493S |
| ARTIC nCoV-2019 V3 Panel, 500rxn | Integrated DNA Technologies | Cat#10006788 |
| TruSeq DNA PCR-Free LT Library Prep | Illumina | Cat#20015962 |
| TruSeq DNA Idx Kit Set A | Illumina | Cat#20015960 |
| MiSeq Reagent Nano Kit, v2 (500 cycles) (1M) | Illumina | Cat#MS-103-1003 |
| **Deposited Data** |        |            |
| Data used to generate figures | This paper | Mendeley Data at [https://dx.doi.org/10.17832/3n377gv8kb](https://dx.doi.org/10.17832/3n377gv8kb) |
| hCoV-19/USA/WA-RML-1/2020 (SARS-CoV-2 patient genome d49 NP swab) | This paper | GenBank: MT982403 |

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### Experimental Models: Cell Lines

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Experimental Models: Cell Lines | | |
| Freestyle 293-F | ThermoFisher | Cat# R79007; RRID CVCL_D603 |
| VeroE6 | Ralph Baric | ATCC CRL-1586 |
| MatTek EpiAlveolar | MatTek Life Sciences ([https://www.mattek.com/products/epialveolar/](https://www.mattek.com/products/epialveolar/)) | Cat# ALV-100-FT-PE12 |

### Oligonucleotides

- **Primer to E genomic (E Sarcheco F1)**
  - AACAGGTACGTTAATAGTTAATAGCGT
  - Corman et al., 2020; Integrated DNA Technologies [https://www.who.int/docs/default-source/coronaviruse/wuhan-virus-assay-v1991527e5122341d99287a1b17c111902.pdf](https://www.who.int/docs/default-source/coronaviruse/wuhan-virus-assay-v1991527e5122341d99287a1b17c111902.pdf)

- **Primer to E subgenomic (sgLeadSARS2-F)**
  - CGATCTCTTGATGATGTTTCTC
  - Wölfel et al., 2020; Integrated DNA Technologies N/A

- **Reverse primer to E (E Sarcheco R2)**
  - ATATTGCAGCAGCAGCACA
  - Corman et al., 2020; Integrated DNA Technologies [https://www.who.int/docs/default-source/coronaviruse/wuhan-virus-assay-v1991527e5122341d99287a1b17c111902.pdf](https://www.who.int/docs/default-source/coronaviruse/wuhan-virus-assay-v1991527e5122341d99287a1b17c111902.pdf)

- **Probe for E (E Sarcheco P1)**
  - FAM-ACACTAGCCATCCTTACTGCGCTTCG-ZEN-IBHQ
  - Corman et al., 2020; Integrated DNA Technologies [https://www.who.int/docs/default-source/coronaviruse/wuhan-virus-assay-v1991527e5122341d99287a1b17c111902.pdf](https://www.who.int/docs/default-source/coronaviruse/wuhan-virus-assay-v1991527e5122341d99287a1b17c111902.pdf)

### Recombinant DNA

- **pCXH SARS-CoV-2 spike plasmid**
  - Kizzemekia Corbett and Barney Graham Vaccine Research Center, NIH, Bethesda, USA (Wrapp et al., 2020)
  - N/A

- **pCAGGS SARS-CoV-2 receptor binding domain plasmid**
  - Florian Kramerlcahn School of Medicine at Mt. Sinai, New York, USA (Amanat et al., 2020)
  - N/A

### Software and Algorithms

- **MAFFT align**
  - (Katoh and Standley, 2013, Katoh et al., 2002)
  - Geneious Prime 2020.1.2 plugin

- **Multalin sequence alignment**
  - (Corpet, 1988)
  - [http://multalin.toulouse.inra.fr/multalin/](http://multalin.toulouse.inra.fr/multalin/)

- **ESPript 3.0**
  - (Robert and Gouet, 2014)
  - [http://espiro.ibcp.fr/ESPript/ESPript/](http://espiro.ibcp.fr/ESPript/ESPript/)

- **Geneious Prime 2020.1.2**
  - Geneious
  - [https://www.geneious.com](https://www.geneious.com)

- **PhyML 3.320180621**
  - Guindon et al., 2010
  - Geneious Prime 2020.1.2 plugin

- **FigTree v1.4.4**
  - [http://tree.bio.ed.ac.uk/software/figtree/](http://tree.bio.ed.ac.uk/software/figtree/)
  - [https://github.com/rambaut/figtree/](https://github.com/rambaut/figtree/)

- **Pangolin COVID-19 Lineage Assigner**
  - Rambaut et al., 2020
  - [https://pangolin.cog-uk.io](https://pangolin.cog-uk.io)

- **Prism 8.2.0**
  - GraphPad
  - [https://www.graphpad.com/443/](https://www.graphpad.com/443/)

- **NextClade v0.7.5**
  - [https://github.com/nextstrain/nextclade](https://github.com/nextstrain/nextclade)
  - [https://nextstrain.org/ncov/global?c=region](https://nextstrain.org/ncov/global?c=region)

- **ModelFinder**
  - Kalyaanamoorthy et al., 2017
  - [http://www.iqtree.org/](http://www.iqtree.org/)

- **Ultrafast bootstrap**
  - Hoang et al., 2018
  - [http://www.iqtree.org/](http://www.iqtree.org/)

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RESOURCE AVAILABILITY

Lead Contact
Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Vincent Munster (Vincent.munster@nih.gov).

Materials Availability
This study did not generate new reagents.

Data Availability
The data and the Supplementary Tables from this study have been deposited to Mendeley Data at https://dx.doi.org/10.17632/3n377gv8kb.
Genome sequences have been deposited to GenBank: MT982403, MT982402, MT982405, MT982406, MT982401 and MT982404.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Human Patient
The patient described in this case study is a 71 year old female with a 10 year history of chronic lymphocytic leukemia (CLL), acquired hypogammaglobulinemia, anemia, and chronic leukocytosis. The patient tested positive for SARS-CoV-2 on March 2, 2020, and...
remained positive through June 15, 2020. During the course of the study, the patient was transfused with intravenous immunoglobulin (IVIG, 25 g) on April 6 and May 6, 2020, and convalescent plasma against SARS-CoV-2 on May 12 and May 23, 2020. After the initial SARS-CoV-2 diagnosis, the patient was kept in isolation in an isolation ward in a single room with negative airflow. Anonymized plasma, serum and swabs from a patient at EvergreenHealth, Kirkland, Washington were obtained under an NIH Institutional Review Board exemption. Verbal and signed consent were obtained from the patient to allow analyses of the samples.

**Cells**

Vero E6 is a female African green monkey kidney epithelial cell line. Vero E6 cells were maintained at 37°C and 5% CO₂ in DMEM supplemented with 10% fetal bovine serum, 1 mM L-glutamine, 50 U/mL penicillin and 50 μg/mL streptomycin. Vero E6 cells were provided by Dr. Ralph Baric. Cells were authenticated by cytochrome B sequencing. Mycoplasma testing was performed monthly, and no mycoplasma was detected.

FreeStyle 293-F (RRID: CVCL_D603) is a female human embryonic cell line adapted for growth in suspension culture. FreeStyle 293-F cells were grown in Freestyle 293 Expression Medium (GIBCO) at 37°C and 8% CO₂, shaking at 130 rpm. Cells were not authenticated in house. Mycoplasma testing was performed monthly, and no mycoplasma was detected.

MatTek EpiAlveolar is a 3D co-culture model of the air-blood barrier produced from primary human alveolar epithelial cells, pulmonary endothelial cells and fibroblasts, and maintained according to manufacturers instructions (https://www.mattek.com/products/epialveolar/). Cells were not authenticated in house. Mycoplasma testing was performed monthly, and no mycoplasma was detected.

**SARS-CoV-2 Virus**

SARS-CoV-2 strain nCoV-WA1-2020 (MN985325.1) (Harcourt et al., 2020) was provided by CDC, Atlanta, USA. SARS-CoV-2 isolates were propagated on Vero E6 cells grown in DMEM supplemented with 2% fetal bovine serum (GIBCO), 1 mM L-glutamine (GIBCO), 50 U/mL penicillin and 50 μg/mL streptomycin (GIBCO) (virus isolation medium), at 37°C and 5% CO₂.

Infectious titer of SARS-CoV-2 virus stocks was determined by end-point titration and is reported as log₁₀ 50% tissue culture infective dose (TCID₅₀/mL). 1.5 × 10⁴ Vero E6 cells were seeded into each well in 96-well plates in DMEM supplemented with 10% fetal bovine serum, 1 mM L-glutamine, 50 U/mL penicillin and 50 μg/mL streptomycin and incubated overnight at 37°C and 5% CO₂. The following morning, when the cells were at approximately 90% confluency, the wells were inoculated with ten-fold serial dilutions of virus stock diluted in virus isolation medium (100 μL per well, with 10 replicate wells for each dilution). The plates were incubated at 37°C and 5% CO₂, and the cytopathic effect (CPE) was assessed for each well after 5 days. Wells that demonstrated CPE were counted, and the titer was determined by the method of Spearman and Kärber using 10 replicates as follows:

\[
\text{Log}_{10}\,	ext{TCID}_{50}/\text{mL} = \left(\frac{X - d}{2} + \left|d \cdot S\right|\right)
\]

where X is log₁₀ of the lowest dilution with all wells positive for CPE, d is log₁₀ of the dilution factor (10 in these titrations), and S is the sum of the fraction of wells positive for CPE at all tested dilutions.

**METHOD DETAILS**

**Clinical Sample RNA Extraction and qRT-PCR**

Clinical samples were deidentified as part of their analyses. Nasopharyngeal and oropharyngeal swabs were shipped on wet ice in viral transport medium (VTM) to Rocky Mountain Laboratories (NIH). RNA was extracted using Trizol (Invitrogen), Phasedmaker tubes (Invitrogen) and the PureLink RNA Mini Kit (Invitrogen) according to manufacturer’s instructions and eluted in 100 μL RNase-free H₂O. First strand cDNA synthesis was performed with the SuperScript IV First Strand Synthesis System (Invitrogen), using 11 μL input RNA and random hexamers. qRT-PCR was performed using 5 μL of cDNA using the QuantiFast Probe kit (QIAGEN) using E gRNA (Corman et al., 2020) and sgRNA specific assays (Wölfel et al., 2020). To quantify viral load within the patient samples, 5 μL of cDNA was analyzed using droplet digital PCR (Biorad) using the same E gRNA and sgRNA assays. The SARS-CoV-2 testing through EvergreenHealth were performed by University of Washington, LabCrop, Cepheid, and GenMark. Kashi clinical laboratories and Magnolia diagnostics performed the negative tests taken at the care facilities.

**Virus Isolation**

Virus isolation of the clinical specimen was performed on Vero E6 cells in 96 well plates. In brief, media was removed from wells and replaced with 100 μL of undiluted swab sample, or swab sample diluted 1:10 in DMEM supplemented with 2% fetal bovine serum (GIBCO), 1 mM L-glutamine (GIBCO), 50 U/mL penicillin and 50 μg/mL streptomycin (GIBCO) (virus isolation medium). Diluted and undiluted samples were inoculated onto 7 wells. Spin inoculation was performed at 1000 x g for 1 hour at 35°C. Inoculum was removed and wells were washed twice with and replaced with 100 μL of virus isolation medium and incubated at 37°C and 5% CO₂. After 5 days, replicate wells were pooled, diluted 10x in virus isolation medium, and used to inoculate T25 flasks of Vero E6 cells in virus isolation medium and incubated at 37°C and 5% CO₂. Flasks were observed for cytopathic effect. RNA was extracted, as described above, for confirmation of SARS-CoV-2 by qRT-PCR and next generation sequencing.
**Growth kinetics of SARS-CoV-2 isolates**

Vero E6 cells were seeded in 6 well plates at a density of 4 × 10^5 cells/well in DMEM supplemented with 2% fetal bovine serum ( Gibco), 1 mM L-glutamine ( Gibco), 50 U/mL penicillin and 50 µg/mL streptomycin (Gibco) (virus isolation medium) and incubated overnight at 37 °C and 5% CO₂. The following day, the media was removed from the wells and replaced with 1 mL of virus isolation medium containing virus at a MOI of 0.01. The patient day49 isolate and the USA/WA1/2020 strain were tested in triplicate, with mock control wells in triplicate. After a 1-hour incubation at 37 °C and 5% CO₂, the inoculum was removed, and wells were washed 3x with PBS and replaced with a fresh 2 mL of virus isolation medium. Supernatant samples were taken at 0, 12, 24, 48, 72, 96, and 120 hours post inoculation. Titer of infectious virus from supernatant was determined by endpoint titration in Vero E6 cells, as described above, but using 4 replicates per sample to determine the TCID₅₀/mL using the Spearman-Karber method. The EpiAlveolar cell growth kinetic experiment was set up similar to the Vero E6 cells but with the following differences. Cells were provided by MatTek with 2.5 × 10⁵ cells/transwell insert. Cells were infected by adding 75 µL of ALI medium containing virus at an MOI of 0.01 to the apical side of the transwell insert. After the above outlined incubation, the inoculum was removed, wells were washed 1x with PBS and replaced with 75 µL of ALI medium upon the apical surface. During sampling of the EpiAlveolar cells, 500 µL of DMEM medium was added to the apical side, gently pipetted to mix, removed, and 75 µL of fresh ALI medium replaced on the apical surface.

**Expression and Purification of SARS-CoV-2 Spike and Receptor Binding Domain**

Expression plasmids encoding the codon optimized SARS-CoV-2 full length spike and receptor binding domain (RBD) were kindly provided Kizzmekia Corbett and Barney Graham (Vaccine Research Center, Bethesda, USA) and Florian Krammer (Icahn School of Medicine at Mt. Sinai, New York, USA), respectively (Wrapp et al., 2020; Amanat et al., 2020). Both plasmids were expressed in Freestyle 293-F cells (Thermofisher), maintained in Freestyle 293 Expression Medium (GIBCO/ThermoFisher) at 37 °C and 8% CO₂ in a humidified incubator shaking at 130 rpm. Cultures totaling 500 mL were transfected with PEI at a density of one million cells per mL. Supernatant was harvested 7 days post transfection, clarified by centrifugation and sterile filtered through a 0.22 µM membrane. The protein was purified using Ni-NTA immobilized metal-affinity chromatography (IMAC) using Ni Sepharose Fast Flow Resin (GE Lifesciences) or NINTA Agarose (QIAGEN) and gravity flow. After elution the protein was buffer exchanged into 10 mM Tris pH 8, 150 mM NaCl buffer before further use or frozen at –80 °C for storage.

**Enzyme-Linked Immunosorbent Assay (ELISA)**

Purified SARS-CoV-2 full length spike or RBD protein was diluted to 1 µg/mL in PBS. Maxisorp plates (Nunc) were coated with 100 µL per well (100 ng protein per well) and incubated overnight at 4 °C. Plates were washed 3x with PBST (0.1% Tween) and blocked with 100 µL casein in PBS blocking buffer (ThermoFisher) for 1 hour at room temp. Plates were again washed 3x with PBST (0.1% Tween), and 100 µL of serum samples, serially diluted 2 fold in casein in PBS blocking buffer, in duplicate, was added to the wells and incubated at room temperature for 1 hour. Plates were washed 4x with PBST (0.1% Tween), and 100 µL secondary antibody, rabbit anti-human IgG Fc HRP (Novus Biologicals, NB51-73529) diluted 1:4000 in casein in PBS blocking buffer, was added to the wells and incubated for 1 hour at room temperature. The wells were washed 5x with PBST (0.1% Tween) and developed with the KPL TMP 2-component peroxidase substrate kit (SeraCare, 5120-0047). The reaction was stopped with KPL stop solution (SeraCare, 5150-0020) and read at 450 nm. The threshold for positivity was calculated as the average plus 3 times the standard deviation of negative control sera. Reported titers are the reciprocal value of the highest dilution at which signal was observed above the calculated threshold.

**Virus Neutralization assay**

Serum and plasma samples were heat inactivated at 56 °C for 30 minutes. Two-fold serial dilutions were prepared in DMEM supplemented with 2% FBS, with each sample diluted in duplicate in 96 well plate format. 100 TCID₅₀ of SARS-CoV-2, in virus isolation medium, was then added to each well. The virus-serum/plasma mixture was incubated at 37 °C for 1 hour to allow for neutralization, then 100 µL per well was added to Vero E6 cells in 96 well plates and incubated at 37 °C and 5% CO₂. After 5 days, wells were observed for cytopathic effect. The virus neutralization titer is displayed as the reciprocal value of the highest dilution of serum/plasma that still inhibited virus replication at which no cytopathic effect was observed.

**Next generation sequencing of patient clinical samples and isolates**

Clinical Samples - Viral RNA was extracted from patient nasopharyngeal swabs using Trizol (Invitrogen) for use with the ARTIC nCoV-2019 sequencing protocol V.1 (Protocols.io; https://www.protocols.io/view/ncov-2019-sequencing-protocol-bbmuik6w). 30-35 PCR cycles were used to generate tiled-PCR amplicons. Primer pools consisted of the ARTIC nCoV-2019 v3 Panel (Integrated DNA Technologies, Belgium) and were designed and used in PCR reactions following the instructions. Products from Pool 1 and Pool 2 were combined, AmPure XP cleaned, and quantitated as per the instructions – through step 16.18. Following assessment on a BioAnalyzer DNA Chip (Agilent Technologies, Santa Clara, CA), a volume consisting of 500 ng of product was taken directly into TruSeq DNA PCR-Free Library Preparation Guide, Revision D. (Illumina, San Diego, CA) beginning with the Repair Ends step (q.s. to 50 µL with RSB) and subsequent cleanup consisted of a single 1:1 AmPure XP/reaction ratio. All downstream steps followed the manufacturer’s instructions. Final libraries were visualized on a BioAnalyzer HS chip (Agilent Technologies, Santa Clara, CA) and quantified using KAPA Library Quant Kit (Illumina) Universal qPCR Mix (Kapa Biosystems, Wilmington, MA) on a CFX96 Real-Time System (BioRad, Hercules, CA).
Isolates - Viral RNA was extracted from clarified cell culture supernatant using Trizol (Invitrogen). Extracted RNA was depleted of rRNA using Ribo-Zero Gold H/M/R (Illumina, San Diego, CA) based on manufacturer’s protocols. After Ampure RNAClean XP (Beckman Coulter, Brea, CA) purification, the enriched RNA was eluted in 6 μL of water and assessed on a BioAnalyzer RNA Pico Chip (Agilent Technologies, Santa Clara, CA). Following the Truseq Stranded mRNA Library Preparation Guide, Revision E., (Illumina, San Diego, CA), the remaining RNA was added to Elute-Frag-Prime Buffer and continued through second-strand cDNA synthesis. The resulting double-stranded cDNAs were treated with a combined mixture of RiboShredder RNase Blend (Lucigen, Middleton, WI) and high concentration DNase-free RNase (Roche Diagnostics, Indianapolis, IN). After Ampure XP purification (Beckman Coulter, Brea, CA), samples were analyzed on a RNA Pico chip to confirm no remaining RNA. Library preparation continued with adenylation of ends following manufacturer’s recommendations. All downstream steps followed the manufacturer’s instructions. Final libraries were visualized on a BioAnalyzer DNA1000 chip (Agilent Technologies, Santa Clara, CA) and quantified using KAPA Library Quant Kit (Illumina) Universal qPCR Mix (Kapa Biosystems, Wilmington, MA) on a CFX96 Real-Time System (BioRad, Hercules, CA).

**Sequencing and bioinformatics**

Libraries were diluted to 2 nM stock, pooled together as needed in equimolar concentrations and sequenced on the MiSeq (Illumina, Inc, San Diego, CA) using on-board cluster generation and 2 × 150 paired-end sequencing. Raw image files were converted to fastq files using bcl2fastq (v2.20.0.422, Illumina, Inc San Diego, CA) and trimmed of adapter sequences using cutadapt version 1.12 (Martin, 2011). Adapter-trimmed reads were trimmed and filtered to remove low quality sequence using fastq_quality_trimmer and fastq_quality_filter tools from the FASTX Toolkit, v 0.0.14 (Gordon, 2018). Singletons were removed and quality filtered reads were coordinate-order sorted using a custom perl script.

Reads were filtered for repeat sequence, rRNA, and PhiX contaminants and then mapped to the SARS-CoV-2 isolate 2019-nCoV/USA_WA1 (MN985325.1) reference genome using bowtie2 with –no-mixed –no-unal -X 1500 options (Langmead and Salzberg, 2012). Aligned SAM files were converted to BAM format, then sorted and indexed using SAMTools (Li et al., 2009). Duplicate reads were removed from the mapped reads using picard’s MarkDuplicates tool (Broad Institute, 2018).

To process the ARTIC data a custom pipeline was developed. Fastq read pairs were first compared to a database of ARTIC primer pairs to identify read pairs that had correct, matching primers on each end. Once identified, the ARTIC primer sequence was trimmed off. Read pairs that did not have the correct ARTIC primer pairs were discarded. Remaining read pairs were collapsed into one sequence using AdapterRemoval (Schubert et al., 2016), requiring a minimum 25 base overlap and 300 base minimum length, generating ARTIC amplicon sequences. Identical amplicon sequences were removed and the unique amplicon sequences were then mapped to the SARS-CoV-2 genome (MN985325.1) using Bowtie2 (Langmead and Salzberg, 2012). Aligned SAM files were converted to BAM format, then sorted and indexed using SAMTools (Li et al., 2009).

Variant calling was performed using Genome Analysis Toolkit (GATK, version 4.1.2) HaplotypeCaller with ploidy set to 2 (McKenna et al., 2010). Single nucleotide polymorphic variants were filtered for QUAL > 200 and quality by depth (QD) > 20 and indels were filtered for QUAL > 500 and QD > 20 using the filter tool in bcftools, v1.9 (Li et al., 2009). The accuracy of the filtered variant calls was manually inspected in Broad’s Integrative Genomics Viewer (IGV) (Robinson et al., 2017). Consensus sequences were generated using bcftools consensus (Li et al., 2009) and subsequently aligned using MAFFT (Kato and Standley, 2013; Katoh et al., 2002) with 2,434 GISAID Washington SARS2 reference sequences in addition to the 2019-nCoV/USA_WA1 genome used for mapping.

**Phylogenomic Analysis**

Available SARS-CoV-2 full genome sequences were downloaded from the GISAID database (https://gisaid.org/; Shu and McCauley, 2017). The sequences were then aligned to previously described lineages (Rambaut et al., 2020) using Pangolin v2.0.3 (https://pangolin.cog-uk.io/), and aligned using MAFFT v. 1.4 (Kato and Standley, 2013; Katoh et al., 2002). A maximum likelihood tree with the patient SARS-CoV-2 genomes, the Wuhan-Hu-1/2019 genome sequence, the USA/WA-1/2020 genome, and a representative genome from the assigned lineages was inferred using PhyML v.3.3.20180621 (Guindon et al., 2010) implemented in Geneious Prime v.2020.1.2 (https://www.geneious.com/) with a general time reversible model of nucleotide substitution and rooted at the Wuhan-Hu-1/2019 SARS-CoV-2 strain. The final figure was made using FigTree v.1.4.4 (http://tree.bio.ed.ac.uk/software/figtree/). For the time tree, full SARS-CoV-2 genomes were subsampled from Washington state representing NextStrain clade 19B, including the four patient genomes and the Wuhan-Hu-1/2019 genome sequence. The sequences were aligned using MAFFT v. 1.4 (Kato and Standley, 2013; Katoh et al., 2002) implemented in Geneious Prime v. 2020.1.2 (https://www.geneious.com/), a maximum likelihood tree reconstructed with PhyML v.3.1 (Guindon et al., 2010), and the time tree showing temporal divergence inferred in TreeTime v.0.7.6 (Hadfield et al., 2018) using the HKY85 model of nucleotide substitution and a fixed molecular clock at 8e-4 with a standard deviation of 4e-4 as implemented in the NextStrain pipeline (https://nextstrain.org/sars-cov-2/).

To evaluate the relationship between the SARS-CoV-2 genomes recovered from the patient swabs with other SARS-CoV-2 genomes from Washington state, genomes at the times of sampling (April 20, May 11, May 26, and June 15, 2020) from Washington state were downloaded from the GISAID database (https://gisaid.org/; Shu and McCauley, 2017). The sequences were aligned by MAFFT (Kato and Standley, 2013; Katoh et al., 2002). The sequences were analyzed by the Nextclade server v0.7.5 (https://clades.nextstrain.org/) for quality and sequences that were not of sufficient quality were discarded. 1,789 sequences at April 20, 385 sequences between April 20 and May 11, 268 sequences between May 11 and May 26, and 709 sequences between May 26 and June 15 were kept for further phylogenetic analysis. Maximum likelihood trees using the curated sets of genomes, the four patient
Electron Microscopy
Vero E6 cells cultured in DMEM supplemented with 10% fetal bovine serum, 1 mM L-glutamine, 50 U/mL penicillin and 50 μg/mL streptomycin were plated at 5 x 10⁴ cells/well in 24 well plates containing Thermanox coverslips (Ted Pella, Redding, CA) for transmission electron microscopy or silicon chips (Ted Pella, Redding, CA) for scanning electron microscopy in the wells, and incubated overnight at 37°C and 5% CO₂. The next day, media was carefully aspirated from the wells and replaced with 1 mL of virus isolation medium containing SARS-CoV-2 virus at a MOI of 1 and incubated for 1 hour at 37°C and 5% CO₂. Wells were washed three times with PBS, then replaced with 1 mL fresh virus isolation medium and incubated at 37°C and 5% CO₂. At 24 and 48 hours post-infection, wells were washed three times with PBS, then fixed as described below.

Scanning electron microscopy
Cells were fixed with Karnovsky’s formulation of 2% paraformaldehyde/2.5% glutaraldehyde in 0.1 M Sorenson’s phosphate buffer, and then post-fixed with 1.0% osmium tetroxide/0.8% potassium ferrocyanide in 0.1 M sodium cacodylate buffer washed with 0.1 M sodium cacodylate buffer then stained with 1% tannic acid in dH₂O. After additional buffer washes, the samples were further osmicated with 2% osmium tetroxide in 0.1M sodium cacodylate, then washed with dH₂O. Specimens were dehydrated with a graded ethanol series from 50%, 75%, 100% x 3 for 5 minutes each, critical point dried under CO₂ in a Bal-Tec model CPD 030 Drier (Balzers, Liechtenstein), mounted with double sided carbon tape on aluminum specimen mounts (Ted Pella), and sputter coated with 35 Å of iridium in a Quorum EMS300T D sputter coater (Electron Microscopy Sciences, Hatfield, PA) prior to viewing at 5 kV in a Hitachi SU-8000 field emission scanning electron microscope (Hitachi, Tokyo, Japan).

Transmission electron microscopy
Specimens were fixed as described above for scanning electron microscopy and additionally stained overnight with 1% uranyl acetate at 4°C after the second osmium staining and then dehydrated with the same graded ethanol series and embedded in Spurr’s resin. Thin sections were cut with a Leica UC7 ultramicrotome (Buffalo Grove, IL) prior to viewing at 120 kV on a FEI BT Tecnai transmission electron microscope (Thermofisher/FEI, Hillsboro, OR). Digital images were acquired with a Gatan Rio camera (Gatan, Pleasanton, CA).

Structure Mapping
The Pymol Molecular Graphics System (https://www.schrodinger.com/pymol) was used to map the location of the observed deletions onto a SARS-CoV-2 spike structure (PDB: 6ZGE; Wrobel et al., 2020). Nucleotide sequence alignments were generated using MAAFT align (Katoh and Standley, 2013; Katoh et al., 2002) implemented in Geneious Prime v.2020.1.2 (https://www.geneious.com) and amino acid sequence alignments were generated with Multalin (Corpet, 1988) and plotted with ESPript (Robert and Gouet, 2014).

QUANTIFICATION AND STATISTICAL ANALYSIS
Data and statistical analysis was performed using GraphPad Prism 8.2.0. Replicates and statistical details can also be found in the methods and figure legends. For ELISA and virus neutralization assays, the serum/plasma samples were diluted and tested in duplicate. For the growth curves, both virus isolates (day 49 patient isolate and USA/WA1/2020) were tested in three replicate wells for both Vero E6 cells and the primary human alveolar epithelial cells. The growth curve data shown are the mean and standard error of the mean for the three independent replicates. The statistical analysis was performed using a 2-way ANOVA in Graphpad Prism 8.2.0. Further methods to determine whether the data met assumptions of the statistical approach were not relevant for these analyses.
Figure S1. ELISA Titers against the SARS-CoV-2 RBD, Related to Figure 2
(A) IgG titers against SARS-CoV-2 receptor binding domain (RBD) were determined in ELISA on convalescent plasma used for transfusion. The light gray bar is the IgG titer of the first donor (convalescent plasma 1) and the dark gray is the second donor (convalescent plasma 2). (B) IgG titers against SARS-CoV-2 (RBD) were determined in ELISA on patient serum collected on several time points, including immediately before and after transfusion with convalescent plasma at day 71 (light gray) and day 82 (dark gray). Each serum/plasma sample was tested in duplicate.
Figure S2. Maximum-Likelihood Trees of the Individual with SARS-CoV-2 with Other SARS-CoV-2 Genomes Circulating in Washington State at the Times of Sampling (April 20, May 11, May 26, and June 15, 2020), Related to Figure 4 and Table 2

(A) Maximum likelihood tree using 1789 full genome SARS-CoV-2 sequences deposited to GISAID until 20 April 2020. Inset shows a close up of the monophyletic clade of the genomes directly obtained from the patient samples (cyan). (B) Maximum likelihood tree using 385 full genome SARS-CoV-2 sequences deposited to GISAID between 20 April and 11 May, 2020. The monophyletic clade of the genomes directly obtained from the patient samples is shown in cyan. (C) Maximum likelihood tree using 268 full genome SARS-CoV-2 sequences deposited to GISAID between 11 May and 26 May, 2020. The monophyletic clade of the genomes directly obtained from the patient samples is shown in cyan. (D) Maximum likelihood tree using 709 full genome SARS-CoV-2 sequences deposited to GISAID between 26 May and 15 June, 2020. The monophyletic clade of the genomes directly obtained from the patient samples is shown in cyan.