Persistence of infectious SARS-CoV-2 particles for up to 37 days in patients with mild COVID-19

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Abstract

Background: People suffering from COVID-19 are typically considered non-infectious 14 days after diagnosis if symptoms have disappeared for at least 48 h. We describe three patients who independently acquired their infection. These three patients experienced mild COVID-19 and completely recovered symptomatically within 10 days, but remained PCR-positive in deep pharyngeal samples for at least 38 days. We attempted to isolate virus from pharyngeal swabs to investigate whether these patients still carried infectious virus.

Methods: Infectious virus was amplified in Vero E6 cells and characterized by electron microscopy and WGS. The immune response was investigated by ELISA and peptide arrays.

Results: In all three cases, infectious and replication-competent virus was isolated and amplified in Vero E6 cells. Virus replication was detected by RT-PCR and immunofluorescence microscopy. Electron microscopy confirmed the formation of intact SARS-CoV-2 particles. For a more detailed analysis, all three isolates were characterized by whole-genome sequencing (WGS). The sequence data revealed that the isolates belonged to the 20A or 20C clade, and two mutations in ORF8 were identified among other mutations that could be relevant for establishing a long-term infection. Characterization of the humoral immune response in comparison to patients that had fully recovered from mild COVID-19 revealed a lack of antibodies binding to sequential epitopes of the receptor-binding domain (RBD) for the long-term infected patients.

Conclusion: Thus, a small portion of COVID-19 patients displays long-term infectivity and termination of quarantine periods after 14 days, without PCR-based testing, should be reconsidered critically.

KEYWORDS
Persistence, SARS-CoV-2, SARS-CoV2, mutations, whole-genome sequencing, antibodies


1 | BACKGROUND

In December 2019, a novel lung disease was first reported to have emerged in Wuhan, Hubei Province, China, and showed very severe disease courses, including some with fatal outcomes. The new infectious disease rapidly spread throughout China and other countries. As the disease progressed, it was recognized to be a coronavirus infection, closely related to SARS (Severe Acute Respiratory Syndrome) and MERS (Middle East Respiratory Syndrome Coronavirus). The disease has been referred to as COVID (Corona Virus Disease)-19 and the causative virus as SARS-CoV-2. The course of the disease can vary substantially in COVID-19 patients. Approximately 80 percent of detected infections are mild to moderate. In severe cases, respiratory distress and lung failure may occur. Severe courses mainly affect older people and individuals with underlying health conditions. However, severe cases can also be observed in people who are young or without pre-existing illness. The virus is primarily transmitted via droplet infection (i.e., when speaking, sneezing, or coughing) and when there is little distance to other people. Transmission via aerosols is also possible. At the time of writing this report, 225,778,167 total cases have been reported worldwide, with 4,648,145 deaths (source: COVID-19 Dashboard by the Center for Systems Science and Engineering [CSSE] at Johns Hopkins University [JHU]). Currently, contact restrictions, distance and hygiene rules, wearing a surgical mask in everyday life, testing for SARS-CoV-2 infection, and vaccination are the most powerful tools to keep infection events under control. There are reports describing that infectious virus could be readily isolated during the first week of symptoms from a considerable number of patients. To the best knowledge of the authors, so far no isolation of infectious virus was possible with samples of moderately or asymptotically infected patients taken after day 10, although significant loads of viral genomes were detected. Based on these findings, a common practice is to isolate COVID-19 patients for 14 days and to discontinue the quarantine without further testing given the patient is asymptomatic for at least 48 h. There are also reports that describe infectivity for at least 24 days after disease onset in patients with severe COVID-19 but showing no infectivity at later stages of a prolonged COVID-19 infection despite positivity in RNA testing. Hence, because of the discrepancy of this finding to the experiments aimed at isolating active virus and the current time periods for quarantine, the timeline for discontinuation of transmission-based precautions is extensively debated.

This study aimed at addressing the question whether persistence of infectious virus particles for prolonged periods of more than 4 weeks is restricted to patients with severe COVID-19 or might also occur in patients with mild symptoms as well. In the present study, we show that in three patients who already had overcome a mild symptomatic COVID-19 disease active virus particles were still present in deep pharyngeal area for up to 37 days. Thus, there is an urgent medical need to understand the pathophysiology of the prolonged stay of active SARS-CoV-2 particles in some patients and to re-evaluate the procedures for discontinuation of transmission-based precautions of formerly infected COVID-19 patients.
2 | MATERIALS AND METHODS

2.1 | Cell culture and incubation with patients' swab

For this study, Vero E6 cells (ATCC \(^\text{®} \) CRL-1586 \(^\text{™} \)) were used. Cells were cultured in Dulbecco's Modified Eagle's Medium (DMEM from BioWest) and supplemented with 2 mM L-glutamine, 100 \(\mu\)g/ml streptomycin, 100 U/ml penicillin, and 5% v/v fetal bovine serum (FBS superior; Sigma-Aldrich) under atmospheric conditions of 95% relative humidity and 5% \(\text{CO}_2\) at 37°C.

Contents of SARS-CoV-2 positive or negative patients' swabs were resuspended in 1 ml of Vero E6 culture medium immediately after collection. The swab suspension was sterile-filtered through a 0.22 \(\mu\)m pore size sterile filter and added to one well of a 12-well cell culture plate with 1.5 \(\times\) \(10^5\) Vero E6 cells per well. After 48 h, supernatant and cells were visually checked for indications of infections, such as color change of medium and especially microscopic examination of generated plaques, which reflect the cytopathic effect of SARS-CoV-2. Supernatant was removed and added to one well of a 6-well cell culture plate containing 3 \(\times\) \(10^5\) Vero E6 cells per well. Additionally, 1 ml of Vero E6 cell culture medium was added to the well for a final volume of 2 ml. After 48 h (96 h after initial incubation), supernatant (2 ml) of one well of a 6-well plate was split into six wells of a 6-well plate containing 3 \(\times\) \(10^5\) cells per well and filled with medium for a total volume of 1 ml per well. An additional 48 h later, supernatant was removed and inactivated by addition of TRIzol LS Reagent (ThermoFisher Scientific) for further sequencing analysis. Cells were washed with PBS and treated with TRIzol \(^\text{™} \) Reagent (ThermoFisher Scientific) for quantitative RT-PCR or washed with PBS and fixed with 3.7% formaldehyde for immunofluorescence staining and confocal laser scanning microscopy (CLSM).

2.2 | Titer quantification and plaque reduction assay

To determine viral titers in the supernatant of Vero E6 cells after inoculation with patients' swabs, supernatant after five passages was added as serial dilutions of 1:20–1:640 to Vero E6 cells. Cells were initially seeded as 2.5 \(\times\) \(10^5\) cells per well in a 12-well plate and grown for 16 h at 37°C, 5% \(\text{CO}_2\) and 90% humidity in DMEM supplemented with 2 mM L-glutamine, 100 \(\mu\)g/ml streptomycin, 100 U/ml penicillin, and 10% v/v FBS before inoculation. Vero E6 cells were inoculated for 1 h at 37°C in aforesaid medium without FBS. After that, medium was removed and 1 ml/well fresh medium containing 5% FBS was mixed with prewarmed liquid agarose (final concentration of 0.4% agarose) and added to the plates. Plates were left at room temperature for 20 min until agar got solid. Afterward, plates were incubated at 37°C for 72 h. For plaque visualization, 1 ml of 8% formaldehyde in PBS was added to each well and incubated for 20 min at 37°C. Agarose and formaldehyde were removed and cells were washed with PBS. Finally, 250 \(\mu\)l 0.1% crystal violet solution (Sigma-Aldrich) in 20% ethanol was added to the cells and incubated for 15 min at room temperature. Crystal violet solution was removed and cells were washed once with ddH\(_2\)O to quantify developed infection plaques.

For determination of neutralizing activity of sera of vaccinated individuals to isolates of long-term positive patients, sera dilutions of 1:20–1:640 were mixed with virus (final concentration 75 PFU/well). Virus was neutralized for 1 h at 37°C in medium without FBS. Afterward, the mixture was added to Vero E6 cells for 1 h at 37°C. An isolate of alpha-variant B.1.1.7 served as a control. Plaque quantification was performed as described above.

2.3 | RNA extraction and cDNA synthesis

RNA was extracted using Direct-zol \(^\text{™} \) RNA MiniPrep (Zymo Research). Reverse transcription of RNA to cDNA was performed by using M-MuLV Reverse Transcriptase RNase H negative (Genaxxon bioscience). RNA extraction and cDNA synthesis were performed according to manufacturer's protocols.

2.4 | Quantitative RT-PCR

Analysis of SARS-CoV-2 gene expression via q-RT-PCR was achieved by using a Maxima SYBR-Green qPCR Kit (ThermoScientific) and primers flanking SARS-CoV-2 N1 (nucleocapsid; FWD: 5′ GACCCAAAAATCAGCGAAAT-3′, REV: 5′-TCTGGTACTGCCAGTGATCTG-3′), N2 (FWD: 5′ TTACAAACATTGCCGCAA-3′, REV: 5′-GCGGCACATTCCGAAGAA-3′), N3 (FWD: 5′ GGGACCTTGAATAACAAAGTTG-3′, REV: 5′-TGTACGCACTGACTG-3′), and RdRP (RNA-dependent RNA polymerase; FWD: 5′ CAACTGGGTAAGCTAGCTTTG-3′, REV: 5′-ACTTGGATAATCCCAACCAT-3′) genes.\(^{13}\) Measurements were performed with a LightCycler 480 Instrument II and analysis via LightCycler 480 Software 1.5.1.62 SP3 using absolute quantification fit points method (Roche).

2.5 | High-throughput sequencing library preparation

The total RNA isolated from the cell culture supernatant was used for Illumina library preparation using a modified NNSR priming method.\(^{14}\) We removed the rRNA from the samples with a QIAseq FastSelect- rRNA HMR kit (Qiagen) using the recommended reaction with the following program: 2 min, 75°C; 2 min, 70°C; 2 min, 65°C; 2 min, 60°C; 2 min, 55°C; 5 min, 37°C; 5 min, 25°C. The above reaction was combined with subsequent reverse transcription with 100 pmol NNSR_RT primer, 10 mM DTT, 0.05 mM dNTPs, 20 Us of RiboLock RNase inhibitor, 4 \(\mu\)g Actinomycin D, and 200 Us Superscript IV (Thermo Fisher Scientific) as follows: 45°C for 30 min; 70°C for 15 min. The cDNA was purified with 1.8 volumes
of magnetic beads, and RNase H digest was performed as recommended (NEB). After repeated bead purification, the cDNA samples were subjected to second-strand synthesis at 37°C for 30 min in a 50 µl final reaction volume containing NEB buffer 2, 0.125 mM dNTPs, 5 Us of 3'-5'exo-Klenow Fragment (NEB), and 200 pmol NNSR-2 primer. The resulting double-stranded DNA was PCR-amplified to obtain the barcoded Illumina libraries with a NEBNext Ultra II Master Mix (NEB), containing 25 pmol NNSRnest_ind_N and NNSR_illumina primers, each, using the following cycling conditions: 98°C 10 s; 5 cycles of 98°C 10 s, 55°C 30 s, 68°C 30 s; 15 cycles of 98°C 10 s, 65°C 30 s, 68°C 30 s. The resulting 400–700-bp-long DNA smears were isolated from 1.5% agarose gels with a Zymoclean Gel DNA Recovery Kit (Zymo Research), quantified with qPCR using a NEBNext Library Quant Kit for Illumina (NEB), and sequenced on an MiSeq instrument with a paired-end 2×301 setting.

The overall quality of the reads were checked with the FastQC tool. Raw reads were trimmed and aligned against the reference sequence of SARS-CoV-2 Wuhan-Hu-1 isolate (Accession number: NC_045512) with Trimmomatic (v0.39) and SNAP, respectively. After sorting, the consensus call was performed with Rsamtools and SAMtools. The variant and phylogenetic analysis was performed with Nextclade (https://clades.nextstrain.org/). The phylogenetic tree was visualized with auspice, which is part of the Nextstrain project.

2.6 Virus concentration, purification, and transmission electron microscopy

After inoculation of Vero E6 cells with swabs of SARS-CoV-2 PCR-positive and -negative patients and viral amplification, 5 ml of virus-containing cell culture supernatant was inactivated overnight by addition of formaldehyde to a final concentration of 4%. Virus concentration, purification, and inactivation was performed according to .

For microscopic, negative stain visualization of viral particles, transmission electron microscopy (TEM) was used. A drop of 20 µl of sample was pipetted on a carbon-coated, gla-m-discharged formvar grid and immobilized for 30 min at RT. After two washing steps with 25 µl ddH2O each, grids were incubated with 1% phosphotungstic acid in ddH2O for 10–30 s at RT. Grids were dried and analyzed using an EM 109 (Zeiss).

2.7 ELISA

Receptor-binding domain protein was produced in HEK293T cells and purified with Ni-NTA affinity chromatography according to the protocol from . Florian Krammer, University of Icahn School of Medicine at Mount Sinai, kindly provided the expression plasmid pCAGGS-sRBD. 50 µl at a concentration of 4 µg/ml RBD in PBS was used to coat 96-well microtiter plates (costar 3590, Corning Incorporation) overnight at 4°C. Afterward, plates were blocked for 1 h at RT with 10% FCS in PBS. Between each step, plates were washed three times with PBS-T (PBS supplemented with 0.05% Tween). The sera were pre-diluted 1:100 in 10% FCS in PBS and incubated for 1.5 h at RT. For detection, anti-human IgG HRP-linked ECL antibodies (Cytiva) were used in a 1:3000 dilution. After 1 h of incubation at RT, ELISA plates were developed with 75 µl TMB ELISA Substrate Solution (eBioscience) for 5 min, stopped with 75 µl 1 N sulfuric acid, and analyzed directly at 450 nm on a Tecan reader (Tecan Group). As a negative control, sera of patients without SARS-CoV-2 infection nor vaccination, collected 2017, were used. All values are given as absorbance at 450 nm. The dotted line, representing positivity cut-off, was determined in respect of negative control.

2.8 Peptide array

For mapping of sequential stretches of the SARS-CoV-2 proteins full-length spike (S), nucleocapsid (N), membrane (M), and envelope (E) protein recognized by antibodies in sera of SARS-CoV-2 PCR-positive and -negative patients were used. In approximation of epitopes, 253 (S), 52 (N), 27 (M), or 9 (E) synthetic overlapping peptides of 15 aa in length with an offset of 5 (S) or 8 (N, M, E) aa between each peptide, representing the respective proteins’ sequences, were synthesized according to the principle of Merrifield using a MultiPep RSI automated peptide synthesizer (Intavis AG). Using a slide-spotting robot (Intavis AG) peptides were immobilized in a duplicate on a cellulose membrane (Intavis AG) attached to a 76 mm × 26 mm slide.

Slides were pre-incubated with 10% 10× Casein Blocking Buffer in TBS-T, pH 7.4, supplemented with 5% sucrose for blocking purposes and incubated with patients’ sera, 1:40 diluted in casein blocking buffer. For antibody detection, slides were incubated with LI-COR antibody anti-human-IgG-800 secondary antibody (1:10.000 diluted in 1:10 Casein Blocking Buffer) and bound antibodies were detected via an LI-COR Odyssey Infrared Imager (Biosciences). The peptide array was validated with sera of convalescent individuals. Sera of these persons were diluted to comparable ODs of the lowest measured IgG OD of the long-term infected ones (~OD 0.4) and compared directly to each other in an individual approach. The results of these peptide arrays showed much stronger binding to defined epitopes in case of the adjusted convalescent sera as compared to the sera derived from the long-term infected patients.

2.9 Immunofluorescence staining and CLSM analysis

Incubated cells were fixed with 3.7% formaldehyde and permeabilized with 0.5% Triton X-100 in PBS. Afterward, cells were blocked with 1% (w/v) bovine serum albumin in TBS-T (20 mM Tris, 150 mM sodium chloride pH 8.8, 0.05% [v/v] Tween 20) and incubated with primary antibody rabbit-anti-SARS-CoV-2 spike protein (40150-T62-COV2-100, Sino Biological; diluted 1:800 in PBS). For detection of primary antibodies, Cy3-coupled donkey-anti-rabbit antibody (Jackson ImmunoResearch Laboratories, Inc; diluted 1:400 in PBS) was used. Nuclei were stained with 4′,6-Diamidin-2-phenylindole (DAPI; diluted
CLSM analysis was performed using a Leica SP8 confocal laser scanning microscope (Leica).

2.10 | Ethics statement

The study (PEI-SARS-CoV2) was approved by the local ethics committee (Ethik-Kommission, Landesärztekammer Hessen 60314 Frankfurt am Main) and written informed consent was obtained from all patients (2020-1664_2-evBO). Patients provided a written agreement to publish their medical data. The study was performed in accordance with the provisions of the Declaration of Helsinki and good clinical practice guidelines.

3 | RESULTS

3.1 | Identification of long-term infected COVID-19 patients

A physician-led medical service (Internal Medical Service, IMS) was established at the Paul Ehrlich Institute (PEI) in Langen/Hessen/Germany in April 2020 to conduct routine and occasion-based testing of employees and to provide counseling and care for employees and families affected by COVID-19. As part of the testing, personal information and possible symptoms are also collected with the consent of the person being tested. Formerly infected employees can only return to the PEI after being tested negative for SARS-CoV-2 using PCR run on samples obtained from throat swabs. At the time of writing, 26 cases of COVID-19 were registered among employees at the PEI. Three infected employees (patients 1–3) were found to still be positive for SARS-CoV-2 at day 37, 33 and 31, respectively, after the initial PCR-based diagnosis. The local health agencies had already released all three employees without any restrictions from home quarantine 14 days after the initial diagnosis. An overview of the infection periods of all long-term infected patients and sample taking is given in the timeline (Figure 1).

3.2 | Patient’s medical history and anamnesis

Patient 1 is male and 27 years old. The patient was tested positive on pharyngeal SARS-CoV-2-RNA using RT-PCR with $C_T = 16$ on 11/30/2020 by a general practitioner at the Institute for Medical Diagnostics Bioscientia GmbH. Symptoms with tightness in the chest and increased temperature of up to 38°C had already existed 2 days earlier since 11/29/2020. The symptoms lasted for 11 days and mainly consisted of tightness in the chest, cough, headache, loss...
of smell and taste, and toward the end of the illness, increased tiredness. Apart from a known hypothyroidism since 2010 under substitution with iodide 200 µg once a day, the patient reported no further diseases. The patient did not take any antivirals, immunosuppressive drugs or other medication before or during his infection. The last collection of material from a deep pharyngeal swab (IMS, PEI) and positive reactivity (Ct = 35) in the RT-PCR for SARS-CoV-2 was conducted on 01/06/2021 (Bioscientia). The PCR assay performed on 01/08/2021 already showed a negative reactivity. Thus, the patient had been documentably infected with SARS-CoV-2 for at least 37 days. Considering the initial symptomatic stage and a possible SARS-CoV-2-positivity even after 01/07/2021, the probable duration of the infection was approx. 39–40 days. On 01/08/2021, a swab was collected for in vitro replication of SARS-CoV-2 in cell culture.

Patient 2 is male and 36 years old. The patient was tested positive on pharyngeal SARS-CoV-2-RNA by a general practitioner on 12/07/2020 by RT-PCR (MVZ Dr. Helge Riegel GmbH). Ct value was 28.06. Prior to diagnosis, there was already symptomatology with sinusitis, headache, and increased temperature up to 38°C since 12/01/2020. Symptoms lasted over 14 days and consisted of nausea, loss of smell and taste, muscle pain, and increased fatigue, primarily at the end of the illness period. Apart from migraine, the patient reported no other underlying diseases. No antivirals, immunosuppressive drugs or other medications were administered to this patient before or during his infection. The last collection of material from a deep pharyngeal swab taken by the IMS with positive reactivity at Ct = 31 using RT-PCR for SARS-CoV-2 (Bioscientia) was performed on 01/08/2021. The PCR analysis on 01/11/2021 (Bioscientia) demonstrated a negative reactivity. Thus, the patient had been documentably infected with SARS-CoV-2 for at least 33 days. Including the initial symptomatic stage and a possible SARS-CoV-2-positivity later than 01/08/2021, the probable duration of infection was, therefore, approximately 39–41 days. A swab was also collected for in vitro replication of SARS-CoV-2 in cell culture on 01/08/2021.

Patient 3 is female and 36 years old. The patient was tested positive on SARS-CoV-2-RNA by RT-PCR by the IMS at the PEI on 12/15/2020 (Division of Virology, PEI) during routine testing. A validation of the positive reactivity was obtained by an external diagnostic laboratory (sampling: IMS, analysis: Bioscientia) on 12/16/2020 using PCR with a Ct value of 15. Interestingly, the patient did not develop any symptoms at any time that would have indicated an infection with SARS-CoV-2. The patient reported a lack of underlying diseases. The patient did not take any antivirals, immunosuppressive drugs or other medication before or during her infection. The last collection of material from a deep pharyngeal swab with clearly positive reactivity using RT-PCR for SARS-CoV-2 was conducted on 01/14/2021 (IMS) with Ct = 31. On 01/18/2021, there was a reported ambiguous PCR reactivity to SARS-CoV-2, and on 01/21/2021, only an initial weak reactivity was observed during the PCR run, which was evaluated as questionably negative. The PCR analysis conducted on 01/21/2021 then resulted in a clear negative reactivity. Thus, the patient had been documentably infected with SARS-CoV-2 for least 31 days. Including the possibility of an infection before 12/15/2020 and the unclear terminal reactivity in PCR analysis that was observed until 21/01/2021, the probable duration of infection was approximately 35–40 days. A swab was also collected for in vitro replication of SARS-CoV-2 in cell culture on 01/18/2021.

As control, three documentably SARS-CoV-2-negative, healthy volunteers P4–P6 were included in this study and material from swabs was subjected to in vitro replication following the identical procedure as described for patients 1–3.

### 3.3 Detection of infectious SARS-CoV-2 in pharyngeal swabs of long-term COVID-19 patients

To determine whether the presence of viral nucleic acids in pharyngeal swabs of patients (P1–P3) who had undergone COVID-19 weeks before, recovered, and lacked any symptoms was based on the presence of infectious virus, we attempted to amplify virus from pharyngeal swabs. For this purpose, pharyngeal swabs were dispensed in medium, sterile-filtered, and added to Vero E6 cells. Pharyngeal swabs from PCR-negative patients served as controls (P4–P6). To exclude any signals that may have been based on input material, virus amplification was performed by two passages over 5 days or over 7 days with five passages. Titer determination via plaque assay after five passages revealed titers of $5.46 \times 10^4$ PFU/ml for P1, $5.6 \times 10^3$ PFU/ml for P2, and $5.89 \times 10^3$ PFU/ml for P3.

To detect viral replication, RNA was isolated from cellular lysates and from cell culture supernatant at both time points. After cDNA synthesis, SARS-CoV-2-specific RT-PCR was performed utilizing three gene segments of the nucleocapsid (N)-gene and one of the RNA-dependent RNA polymerase (RdRP) gene. RT-PCR analysis revealed in the case of isolates from P1 low Ct values (approximately 10 for the intracellular RNA and 18 for the extracellular RNA) after 5 days (Figure 2A,B). For P2 and P3, Ct values of about 10 were observed after 7 days for intracellular RNA and roughly 23 for extracellular RNA. In contrast, all negative controls showed high values of approximately 40, comparable to the water control. For all positive samples, detection of the RdRP gene always showed the highest Ct values in all patients. Because the positive RT-PCR data were obtained after several rounds of passaging and reflect a significantly higher genome number as compared to the input, contamination by input RNA can be excluded.

To corroborate these data that indicate the presence of infectious replication-competent virus in the pharynx-derived material, Vero E6 cells, inoculated with the pharynx-derived material for 7 days and four passages, were analyzed by immunofluorescence microscopy (CLSM) using spike-specific antibodies (Figure 3). The immunofluorescence microscopy revealed viral replication evidenced by the significant production of spike proteins in the case of cells infected with the isolates from P1 and P2. For P3, only some cells showed spike-specific signals.
Taken together, these data indicate the presence of replication-competent infectious virus in the pharyngeal tract of all tested long-term PCR-positive patients.

3.4 | Detection of coronavirus particles by electron microscopy

To further confirm the formation of intact viral particles, virions were isolated by ultracentrifugation and analyzed by TEM after negative staining. The electron microscopy clearly showed the presence of viral particles with the characteristic morphology of coronaviruses in the amplified isolates of all three tested patients (Figure 4). The size of viral particles, with a diameter of approximately 75 nm, the length of spike structures on the surface of these particles of about 15 nm, and the general morphology corresponded to SARS-CoV-2 particles (Figure 4D). Taken together, the electron microscopy confirmed the findings of the RT-PCR and immunofluorescence microscopy: the pharyngeal swabs obtained from patients 1, 2, and 3 contained infectious, replication-competent SARS-CoV-2.
3.5 | SARS-CoV-2 isolated from long-term infected patients belonged to 20A and 20C clade

To characterize the isolates in more detail and classify the isolates, Next-Generation Sequencing (NGS) analysis was performed. For this purpose, RNA was isolated after a single passage to minimize the risk of in vitro mutations. Based on the sequence analysis, the isolates derived from P1 and P3 could be identified as belonging to the 20C clade, whereas that of P2 belongs to the 20A clade (Figure 5A). (GenBank accession numbers; Patient 1: OK075089, Patient 2: OK075090, Patient 3: OK075091).

The detailed analysis of the whole-genome revealed eleven mutations in P1 and P3 and ten mutations in P2 compared to the aligned strain NC_045512.2 (Table 1). These mutations include the common mutations D614G of the spike protein as well as the often co-occurring mutations P314L in ORF1b/RdRP and T265I in ORF1a.23-26 Additionally, the also already well-known Q57H mutation in ORF3a (not detected in isolate of P2) and the S686G mutation in the spike that was originally described in virus isolates of ferrets were identified.24,27 Besides, the more uncommon mutations H125Y in the M protein, the mutations V818A in ORF1b, D35Y and A51S in ORF8, and T95I and H245R in the spike protein could be identified. No particular abnormalities could be detected within the receptor-binding domain (RBD).

3.6 | Lack of antibodies binding to sequential epitopes in the RBD

To study the humoral immune response of the long-term infected patients, the amount of SARS-CoV-2 RBD antibodies were determined by ELISA and stratified for IgG, IgA, and IgM. Sera were collected at day 38 for patient 1, after 36 days for patient 2, and after 31 days for patient 3 after initial PCR analysis. For all three patients,

![Visualization of isolated SARS-CoV-2 particles via TEM.](image)
the presence of RBD-specific antibodies was detected (Figure 6). For patient 1, a higher IgG response was found as compared to patients 2 and 3. The highest IgM response was found for patient 2, whereas for patients 1 and 3, a moderate level was detected. A low IgA level was found for patients 1 and 2, and for patient 3, the level was moderate. Compared to convalescent sera of SARS-CoV-2 infected subjects, IgG and IgA antibody levels of all long-term PCR-positive individuals showed lower values, but higher IgM levels.

For a more detailed analysis of the antibody response, peptide arrays covering the S, N, M, and E protein were established, and the binding pattern of the sera derived from patients 1–3, from three negative patients, and from three convalescent plasma samples were determined (Figure 7). It should be emphasized that this technique mainly reflects binding to sequential epitopes. To identify specifically recognized peptides, the signals obtained for sera from SARS-CoV-2 negative samples were considered as background (Figure 7A).

If the background signals were subtracted from the specific signals (Figure 7C), there was only one sequential epitope specifically detected in the N protein by serum derived from patient 1. In case of the serum derived from patient 2 two linear epitopes in the S1 domain of the spike, one epitope derived from the N protein...
and one from the M protein were identified. In contrast to this, the linear epitopes recognized by serum derived from patient 3 were completely different from the patterns observed for the other patients. Here, we identified one epitope at the C-terminal part of the spike S2 domain and one in the N protein (Figure 7A,C; Table 2). In contrast, a wide variety of sequential epitopes was recognized with high intensity by convalescent sera (Figure 7B) derived from patients who rapidly eliminated the virus (Figure 7B). These epitopes were not recognized by sera derived from long-term patients (Table 3). In the case of the convalescent sera, epitopes localized within the RBD were recognized. This finding is in clear contrast to the binding pattern obtained for the long-term patients; here, no specific binding to linear epitopes covering the RBD was observed. Taken together, these data indicate that the pattern of sequential epitopes differs for sera derived from long-term infected patients and from patients that eliminated the infection within 10 days.

3.7 | Reduced neutralization of isolate P1 by sera of vaccinated individuals

To determine how efficiently infection with isolates of long-term positive patients can be neutralized by sera of Comirnaty (BNT162b2, Biontech/Pfizer) vaccinated individuals, a plaque reduction neutralization test (PRNT) was performed and compared to neutralization of an isolate of the SARS-CoV-2 alpha-variant (B.1.1.7; Figure 8). First, neutralization tests showed different neutralizing capacities of the three used sera, with low (S1), medium (S2) and high (S3) neutralizing capacity concerning all isolates. While isolates P2 and P3 could be neutralized by S1 at dilutions lower than 1:320, this serum blocked infection with the alpha-variant and P1 at 1:80 and ≤1:80. Medium neutralizing serum S2 could block infection with alpha at a dilution of about 1:320 and P2 and P3 at a dilution of ≤1:640, and a dilution of 1:40 was needed to neutralize infection with P1 isolate. While S3 neutralized infection with isolates alpha, P2 and P3 in dilutions of 1:640, it shows slightly less efficiency neutralizing P1 with ≤1:640. These data suggest that isolates of P2 and P3 are more efficiently neutralized by sera of vaccinated individuals compared to SARS-CoV-2 variant B.1.1.7. However, infection with isolate of P1 could be less efficiently blocked by all sera compared to the other isolates of long-term positive patients or to the alpha-variant.

4 | DISCUSSION

The data presented in this study demonstrate that in non-immunocompromised patients, SARS-CoV-2 infection may persist for several weeks. Moreover, these results indicate that the detection of viral nucleic acids even weeks after diagnosis of infection does not only reflect remnants of viral genomes. There were still infectious viral particles detectable in the pharynx as demonstrated by amplification and detection the intact and functional virus. Apart from the RT-PCR data, immunofluorescence microscopy, WGS, and electron microscopy unequivocally showed that infectious virus was present in the pharyngeal swabs of the patients, which could be isolated and amplified. Because the quantification of infectious viral particles depends on the quality of the swab, conclusions on the viral load in the respective patients require a critical assessment. However, in all three patients, even more than 4 weeks after COVID-19 diagnosis the positive RT-PCR results obviously correlated with the presence of infectious viral particles, reflecting an ongoing active infection.

A detailed analysis by NGS revealed that SARS-CoV-2 isolated from P1 and P3 belonged to clade 20C, whereas isolate of P2 belonged to clade 20A and sequence analysis revealed several mutations. The 20A clade first attracted attention in beginning of 2020 with increasing outbreaks in Europe, followed by displacement of other lineages in North and South America.28 It is particularly characterized by its D614G mutation in the spike protein, caused by A23403G nucleotide change.29 Clade 20C on the other hand represents a daughter clade of 20A, also harboring the D614G mutation.30 This mutation had already been described as part of a widespread variation, leading to increased infectivity due to facilitated ACE2-binding characteristics.33,34 This D614G mutation is known to be frequently accompanied by the P314L in ORF1b/RdRP that had also been identified in the patient 1 isolate (Table 1).25 Another common mutation that could be detected in these isolates was T265I in ORF1a. The threonine substitution by isoleucine at position 265 leads to the addition of a beta-sheet structure.36 We could identify the Q57H mutation in ORF3a in the isolates of P1 and P3, which was first described in Singapore in February 2020. This mutation coexists in many cases with the mutation T85I in non-structural protein 2 (NSP2) that, in contrast, could not be identified in this isolate.24 One mutation that was originally described in ferrets and was
supposed to lead to immune evasion before receptor binding was the S686G mutation in the spike protein that we could also identify as part of the analyzed sequence. A rare missense mutation that was detected in our study is H125Y in the M protein. It is not clear what impact that mutation might have in the context of stability or immune evasion, but it seems to be a mutation with globally increasing incidence.\textsuperscript{31} Furthermore, next to the well-recognized mutations D614G and P314L in ORF1b a new and so far unknown mutation V818A in ORF1b, was detected. Because ORF8 is known to play a crucial role in inhibiting the type I interferon signaling pathway as well as major histocompatibility complex I (MHC I) degradation, it was interesting to detect two mutations in ORF8.\textsuperscript{32,33} As MHC I degradation facilitates immune evasion, it was an unexpected finding that mutation D35Y that we identified in ORF8 was formerly described as providing protein stability and decreasing disease severity.\textsuperscript{34,35} The second ORF8 mutation A51S, in contrast, is located within two possible CD4+ T cell epitopes.\textsuperscript{36} The consequences arising from this mutation had yet not been investigated. However, these epitope mutations could possibly affect the T cell response against SARS-CoV-2 positive cells and delay virus elimination by the...
cellular immune response. In addition to these mutations affecting T cell epitopes, two further mutations in the spike protein were identified that could affect the T cell response. This includes T95I, a mutation occurring within a predicted CD8+ T cell epitope, and mutation H245R that might be located both within a T cell and/or B cell epitope. Therefore, it can be hypothesized that these mutations contribute to impaired virus elimination, as reflected by a productive long-term SARS-CoV-2 infection.

**TABLE 2** Sequential epitopes identified in infected long-term patients

| Patient 1 | Epitopes: (identified outliers) |
|-----------|--------------------------------|
| N         | G-A-L-N-T-P-K-D-H-I-G-T-R-N-P |

| Patient 2 | Epitopes: (identified outliers) |
|-----------|--------------------------------|
| S1        | S-L-I-V-N-N-A-T-N-V-V-I-K-V   |
| N         | L-P-Y-G-A-N-K-D-G-I-W-V-A-T    |
| M         | Q-R-V-A-G-D-S-G-F-A-A-Y-S-R-Y  |

| Patient 3 | Epitopes: (identified outliers) |
|-----------|--------------------------------|
| S2        | S-P-D-V-D-L-G-D-I-S-G-I-N-A-S  |
| N         | T-A-S-W-F-T-A-L-T-Q-H-G-K-E-D  |

Moreover, we found distinct differences in relative antibody concentrations in serum between the three long-term patients as compared to convalescent patients. In the majority of SARS-CoV-2 patients, IgG and IgM is detectable 2–3 weeks after onset of symptoms. For patients with no obvious emerging symptoms or who remained completely asymptomatic, IgG and IgM levels are consistently less pronounced compared to patients with noticeable mild, moderate, or severe symptoms. Patients 1 and 2 in this study both showed SARS-CoV-2 infections with similar symptoms, but only patient 1 demonstrated an IgG seroconversion that resulted in a considerable increase of IgG antibody titers (Figure 6), whereas IgG levels in patient 2 were similar to asymptomatic patient 3. Another striking feature of the antibody response in patient 2 are the levels of IgM clearly exceeding those of IgG after 38 days of symptom onset, albeit IgG levels are typically higher than IgM after such time period. Furthermore, symptomatically infected patients 1 and 2 surprisingly produced only a rather weak IgA response at all, even though other studies described early and rather high IgA titers beginning from approximately 10 days to more than 40 days and even longer in symptomatically infected patients. The lowest IgA titers were observed in both symptomatically infected patients 1 and 2, and in both cases, swab inoculation led to distinct in vitro infections (Figure 3A), whereas the highest IgA titers were measured in asymptomatic patient 3. Here swab inoculation caused only weaker infection in vitro (Figure 3B). These observations strengthen the hypothesis of a correlation of IgA titers with the efficacy of virus neutralization.

**TABLE 3** Epitopes restricted to infected control patient

| Control inf. patient 1 | Epitopes |
|-------------------------|----------|
| S1                      | H-K-N-N-K-S-W-M-E-S-E-F-R-V-Y |
| R-S-Y-L-T-P-G-D-S-S-S-G-W-T-A |
| Q-P-R-T-F-L-L-K-Y-N-E-N-G-T-I |
| T-D-A-V-D-C-A-L-D-P-L-S-E-T-K |
| RBD                     |
| S2                      | G-P-K-K-S-T-N-L-V-K-N-K-C-V-N |
| F-N-F-N-G-L-T-G-G-V-L-T-E-S |
| RBD                     |
| S1                      | S-V-I-T-P-G-N-T-S-N-Q-V-A-V  |
| A-D-Q-L-T-P-T-W-R-V-Y-S-T-G-S |
| N                       | L-P-Y-G-A-N-K-D-G-I-W-V-A-T  |

| Control inf. patient 2 | Epitopes |
|------------------------|----------|
| S1                      | C-T-F-E-Y-V-S-Q-P-F-L-M-D-L-E |
| R-S-Y-L-T-P-G-D-S-S-S-G-W-T-A |
| RBD                     |
| S2                      | L-S-R-L-D-K-V-E-A-E-V-Q-I-D-R |
| L-Q-S-L-Q-T-Y-V-T-Q-Q-L-I-R-A |
| N                       | L-P-Y-G-A-N-K-D-G-I-W-V-A-T  |

| Control inf. patient 3 | Epitopes |
|------------------------|----------|
| S1                      | G-T-N-T-S-N-Q-V-A-V-L-Y-Q-D-V |
| P-T-W-R-V-Y-S-T-G-S-N-V-F-Q-T |
| S2                      | V-T-L-A-D-A-G-F-I-K-Q-Y-G-D-C |
| A-R-D-L-I-C-A-Q-K-F-N-G-L-T-V |
| L-S-S-N-F-G-A-I-S-S-V-L-D-I |
| N                       | L-G-T-G-P-E-A-G-L-P-Y-G-A-N-K |
| L-P-Q-G-T-T-L-P-K-G-F-Y-A-E-G |
| M                       | L-R-G-H-L-R-I-A-G-H-H-L-G-R-C |
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CONFLICT OF INTEREST

The authors declare no conflict of interest.

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