SARS-CoV-2 normalized viral loads and subgenomic RNA detection as tools for improving clinical
decision-making and work reincorporation

Marta Santos Bravo¹, David Nicolás², Carla Berenguà³, Mariana Fernandez¹, Juan Carlos Hurtado¹,
Marta Tortajada⁴, Sonia Barroso⁴, Anna Vilella⁵, Mar Mosquera¹, Jordi Vila¹, María Angeles Marcos¹.

1. Department of Microbiology, Hospital Clínic of Barcelona. Institut of Global Health
   (ISGlobal), Barcelona, Spain.
2. Hospital at Home Unit, Internal Medicine Service, Hospital Clínic Barcelona. University of
   Barcelona, Spain
3. Department of Microbiology, Hospital de la Santa Creu i Sant Pau, Barcelona, Spain.
4. Risk Prevention Service, Hospital Clínic of Barcelona, Barcelona, Spain.
5. Epidemiology and Preventive Medicine Service, Hospital Clinic of Barcelona. Institut of
   Global Health (ISGlobal), Barcelona, Spain.

Short summary:

SARS-CoV-2 normalized viral loads and sgRNA detection are two rapid accessible tools that
overcome cycle threshold value and respiratory sample collection variability. They could be easily
implemented in routine hospital practice providing a useful proxy for infectivity and COVID-19
patient follow-up.
Previous presentation

Results showed in table 2 were presented in an oral session (#504) titled “Subgenomic RNA as a surrogate marker for active SARS-CoV-2 replication in health care workers for work reincorporation” at the ECCVID Conference of Coronavirus Disease on September 23-25th of 2020, online.

Corresponding author:

Marta Santos Bravo, PhD; marta.santos@isglobal.org
Department of Clinic Microbiology, Hospital Clinic of Barcelona – University of Barcelona
ISGlobal Barcelona Institute for Global Health (Barcelona, Spain).
Villarroel Street, 170. Stairs 11, Floor 5th. 08036 Barcelona, Spain.

Alternate corresponding author:

Maria Angeles Marcos Maeso, MD; mmarcos@clinic.cat
Department of Clinic Microbiology, Hospital Clinic of Barcelona – University of Barcelona
ISGlobal Barcelona Institute for Global Health (Barcelona, Spain).
Villarroel Street, 170. Stairs 11, Floor 5th. 08036 Barcelona, Spain.
ABSTRACT

Background

SARS-CoV-2 RT-PCR provides a highly variable cycle-threshold (Ct) value that cannot distinguish viral infectivity. Subgenomic RNA (sgRNA) has been used to monitor active replication. Given the importance of long RT-PCR positivity and the need for work reincorporation and discontinuing isolation, we studied the functionality of normalized viral loads (NVL) for patient monitoring and sgRNA for viral infectivity detection.

Methods

NVL measured through the Nucleocapsid and RNA-dependent-RNA-polymerase genes and sgRNA RT-PCRs were performed in 2 consecutive swabs from 84 health-care workers.

Results

NVL provided similar and accurate quantities of both genes of SARS-CoV-2 at two different time-points of infection, overcoming Ct-value and swab collection variability. Among SARS-CoV-2-positive samples, 51.19% were sgRNA-positive in the 1st RT-PCR and 5.95% in the 2nd RT-PCR. All sgRNA-positive samples had >4log_{10}RNAcopies/1000cells, while samples with ≤1log_{10} NVL were sgRNA-negative. Although NVL were positive until 29 days after symptom onset, 84.1% of sgRNA-positive samples were from the first 7 days, which correlated with viral culture viability. Multivariate analyses showed that sgRNA, NVL and days of symptoms were significantly associated (p<0.001)

Conclusions

NVL and sgRNA are two rapid accessible techniques that could be easily implemented in routine hospital practice providing a useful proxy for viral infectivity and COVID-19 patient follow-up.

Keywords: COVID-19; SARS-CoV-2; subgenomic RNA; normalized viral loads; health-care workers.
BACKGROUND

Severe Acute Respiratory Syndrome-Coronavirus-2 (SARS-CoV-2) is an RNA virus that emerged in Wuhan, China, at the end of 2019 and spread rapidly worldwide with 10-15% of severe courses and 2.2% of associated mortality [1, 2]. The main method for SARS-CoV-2 diagnosis is reverse transcription-polymerase chain reaction (RT-PCR). This technique provides a cycle threshold (Ct) value, which is inversely related to viral load, such that every increase in a Ct value of 3.3 corresponds to an approximate 10-fold reduction of RNA amounts. This value has been used to aid in interpretation and clinical decision-making [3]. However, Ct values are not directly comparable across assays [4] and do not reflect the true viral load, which requires standardization using reference curves. Moreover, further variations may occur due to differences in specimen collection quality and reaction conditions [5]. Sample quality should be verified by cell quantification and normalization of viral loads to complement the RT-PCR results in the diagnosis [6].

SARS-CoV-2 RT-PCR can remain positive for several weeks after clinical recovery in sicker patients, as well as in asymptomatic and mild ill individuals, leading to prolonged isolation and long work leave [7]. Higher viral load peaks have been found during the first week of the infectious process generally coinciding with the pre- or asymptomatic period, when most transmissions have been described [8]. However, the duration of infectivity after the onset of clinical symptoms remains uncertain.

Viral culture is the gold standard technique for viral viability. Recent studies have shown efficient transmission of SARS-CoV-2 to contact-infected hamsters, and the results have been correlated with the detection of infectious virus by culture but not by RT-PCR viral loads alone [9]. Virus culture might be a good surrogate for transmissibility, however, high biosafety laboratory equipment and the long-term response limit the usefulness of this technique.

Viral subgenomic RNAs (sgRNA) in coronaviruses are only transcribed in infected cells and are poorly incorporated into mature virions, indicating the presence of actively infected cells in the samples [10, 11]. SARS-CoV-2 sgRNA has been used as a surrogate indicator of active virus replication with a
limit of sensitivity of 0.4% of genomic RNA, showing active replication in throat swabs during the first 5 days of symptoms [12], which, however, should be verified with viral culture.

Understanding the duration of SARS-CoV-2 infectivity is crucial for discontinuing isolation, health care worker (HCW) reincorporation, particularly in hospitals facing staff shortages, and save-consumes testing supplies of repeated RT-PCRs. Therefore, strategies aiming an early diagnosis of infected HCWs, preventing nosocomial infections and deciding work reincorporation policies are still needed.

The main objective of this study was to provide two tools for improving clinical decision making and work reincorporation in COVID-19 patients: (i) normalized viral loads (NVL) for patient monitoring and, (ii) sgRNA as a surrogate marker of active replication for discontinuing isolation and work reincorporation in a cohort of HCW.

MATERIAL AND METHODS

Study population

According to hospital protocol, RT-PCR screening for SARS-CoV-2 was performed weekly by the Prevention Service on every HCW working in COVID-19 areas in the Hospital Clinic of Barcelona and also in those HCW with self-reported symptoms suspicious of COVID-19. Every infected HCW was followed by the Home Hospital Unit, with medical and nurse assessment by phone/video calls, home visits and day-hospital visits if needed for further testing.

This study is based on the prospective cohort of infected HCW from the Hospital Clinic of Barcelona from the first diagnosis of COVID-19 (25th February 2020) until the 25th of May 2020 [13]. Only patients with 2 consecutive positive SARS-CoV-2 RT-PCR results were included in the analysis. Work reinstatement was decided by the Human Resources and Work Health Department according to governmental protocols.
Laboratory testing

Inactivation and extraction of samples

All samples were inactivated with 1:1 volume of Cobas Omni Lys (Roche, Germany) and total nucleic acid was extracted using MagNA Pure Compact (Roche, Switzerland). Throat/nasopharyngeal swabs and elutes were aliquoted and stored at -80ºC.

RNA RT-PCR quantification and normalization of SARS-CoV-2

SARS-CoV-2 RT-PCR of the Nucleocapsid (N) and the RNA dependent RNA polymerase (RdRp) genes were used to quantify the number of RNA copies per PCR using the SARS-COV-2 r-gene kit (Biomérieux, France) following the manufacturer’s instructions. This test has a sensitivity limit of 380 copies/mL. Viral load was quantified in RNA copies per PCR using a standard curve made with AMPLIRUN SARS-CoV-2 RNA CONTROL (Vircell, Spain). Nasopharyngeal swab quality was checked by the CELL Control r-gene kit (Biomérieux, France) that provides a quantified plasmid with the HPRT1 housekeeping gene for cellular quantification. Samples with <100 cells/PCR were discarded. Viral loads were normalized according to cellular quantification as the number of RNA copies per 1000 cells. Extraction and both amplifications were checked with positive and negative controls for each reaction.

RT-PCR for sgRNA detection for SARS-CoV-2

All samples were tested for the presence of Envelope (E) sgRNA using the leader-specific primer described by Wölfel et al [12] as well as primers and probes targeting sequences downstream of the start codons of the E gene [14]. RT-PCR of E genomic RNA was performed only in sgRNA positive samples to estimate the amount of sgRNA through $2^{(Ct \text{ sgRNA} - Ct \text{ gRNA})}$ [12]. RT-PCRs were performed using the SuperScript™ III Platinum™ One-Step RT-PCR kit (Invitrogen) with 400nM primers concentration and 200nM probe concentration. Cycling involved 15 min at 50ºC for reverse transcription, 3 min at 95ºC for Taq activation and 45 cycles of 10s at 95ºC, 15s at 60ºC (where the fluorescence was quantified), and 5s at 72ºC in the thermocycler StepOne (Applied Biosystems).
SARS-CoV-2 serology

Specific anti-SARS-CoV-2 immunoglobulin G (IgG) was determined using the qualitative Elecsys Anti-SARS-CoV-2 immunoassay (Roche, United Kingdom).

Statistical analysis

We report the number and percentage of patients for categorical variables and the median (first quartile; third quartile) for continuous variables. Categorical variables were compared using the chi-squared or Fisher exact test, whereas two groups of continuous variables were compared using the nonparametric Mann-Whitney U test. Spearman's correlation coefficient was used to measure the relationship between two continuous variables. RdRp RNA copies/1000 cells and N RNA copies/1000 cells were log10 transformed for normalization before analysis. Linear regression analyses were also used to examine the associations between continuous clinical parameters and RdRp RNA copies/1000 cells, N RNA copies/1000 cells and sgRNA, whereas logistic regression analyses were performed to determine the relationship between dichotomous clinical parameters and RdRp RNA copies/1000 cells, N RNA copies/1000 cells and sgRNA. The associations were further assessed with regression models adjusting for age, sex and comorbidities. The effect sizes of association were shown as regression coefficients and 95% confidence intervals (CIs) for continuous parameters and odds ratios and 95% CIs for dichotomous parameters. The level of significance was set at 0.05 (2-tailed). All analyses were performed using IBM SPSS Version 26.0 (IBM Corp., Armonk, NY, USA).

Ethical approval

The study protocol was evaluated and approved by the Ethical Board of the Hospital Clínic of Barcelona (HCB/2020/0444). The informed consent waiver was provided due to the state of infectious disease emergency. Admission to the Home Hospitalization program was voluntary, as was every medical procedure performed.
RESULTS

Clinical description

A total of 93 HCW were selected for having 2 consecutive positive SARS-CoV-2 RT-PCRs with at least a 7-day gap between tests. Assays could not be performed in 9 subjects due to lack of samples, negative RT-PCR repetition or no clinical data collected. The 84 HCW analyzed had a median age of 33.5 (27; 50.5) years and 58 (69%) of them were females. Clinical characteristics are shown in Table 1.

SARS-CoV-2 infection in this cohort of HCW was detected by routine hospital screening in 33.8%, due to symptom reporting in 64.6% and by direct contact in 1.5%. Symptoms were reported at some point during the infection in 91.7% of the total study population, with 89.3% being symptomatic at the time of diagnosis. Only 7 (8.3%) patients remained asymptomatic.

A month after the HCW screening started, specific anti-SARS-CoV-2 IgG serology was performed. All participants presented positive serology, except one that remained negative even 43 days after the first detection. Serology was not performed in 3 subjects.

During follow-up, 6 (7%) patients required chest x-ray, 3 presented bilateral infiltrates and 3 were normal. Five (6%) subjects required treatment according to the local guidelines at that time (azithromycin, hydroxychloroquine, lopinavir/ritonavir and prednisone). None of the HCWs in the present study required oxygen supplementation or hospital admission.

Microbiologic results

A total of 168 nasopharyngeal/throat swabs from the 84 patients were selected to quantify viral loads based on the N and the RdRp genes and normalized per 1000 cells by RT-PCR. All samples were SARS-CoV-2 positive for at least one of the target genes, and had >100 cells per swab to be considered.

RT-PCR Ct values and NVL of the N and RdRp genes of the 1st and 2nd samples are shown in Table 2.
Results demonstrated that Ct values were very variable when testing different genes in the 2nd RT-PCR, whereas NVL results provided similar results for the RdRp and N genes in the 1st and 2nd detections.

Subgenomic RNA RT-PCR was performed in the 168 samples as an indicator of active replication. The results showed that sgRNA was positive in 43 (51.19%) HCW in the first RT-PCR and in 5 (5.95%) in the second, after a 12-day interval (11; 16). Subgenomic RNA results were compared with N and RdRp RT-PCR Ct values and NVL (RNA copies/1000 cells) in linear and logarithmic scales demonstrating a highly significant correlation with both parameters (Table 3). All samples with N and RdRp RT-PCR Ct values <24.38 were sgRNA positive, whereas samples with Ct values > 27.85 were sgRNA negative. In relation to NVL, all samples with >4 log10 RNA copies/1000 cells were sgRNA positive, 1-3 log10 samples were variable and ≤1 log10 samples were sgRNA negative (Figure 1).

**Microbiologic and clinical variables association**

NVL of the 74 HCWs symptomatic at diagnosis were compared with days after symptom onset (Figure 2A). NVL of the N and RdRp genes decreased over time. SARS-CoV-2 samples remained positive until 29 days after symptom onset and only one sample 38 days afterwards.

Results showed that sgRNA was positive in 3 asymptomatic HCWs in the 1st RT-PCR and in 1 subject who presented symptoms after the 1st RT-PCR. Of the rest of positive sgRNA samples, 84.1% were from ≤7 days after symptom onset, and this percentage declined over infection-time, as shown in Figure 2B. Dividing the presence of sgRNA by different intervals of days of symptoms, the highest proportion of sgRNA detection was in samples from ≤7 days, being only 7 samples sgRNA-positive from the 7th day onwards (Table 4). Statistical analyses showed that all three parameters (sgRNA, RdRp and N NVL) were significantly associated with days of symptoms (see Supplementary Table 1).

Univariant and multivariant analyses of microbiological results and the remaining clinical data were carried out to determine potential risk factors (see Supplementary Table 2). Only significant associations (p<0.05) with a 95% confidence interval (CI) were presented (Table 5). The only two symptoms that individually correlated with SARS-CoV-2 viral load in the first detection were fever
and gastrointestinal disorders (GID). Each symptom appeared in patients with viral loads $\approx 2 \log_{10}$ higher, and positive sgRNA in the case of fever. In contrast, dysgeusia was associated with lower NVL in the 2nd detection.

**DISCUSSION**

This study presents two tools for improving clinical decision-making and work reincorporation after SARS-CoV-2 infection: (i) NVL that provide robust and precise SARS-CoV-2 RNA measurements which overcome Ct values and respiratory sample collection variations; and (ii) sgRNA as a possible surrogate marker of active viral replication, as it presented a significant association with NVL and days after symptom onset.

Ct values have been used as a prognostic indicator for clinical decision-making [15], however they could vary significantly between and within methods [16]. The College of American Pathologists surveyed more than 700 laboratories and demonstrated that different methods using identical control material varied by as much as 14.0 cycles and by up to 12.0 cycle differences within a single gene target using a single method [17]. Therefore, Ct values do not have a linear relation with viral loads, and the dispersion of the measurements is understated. The use of reference standard curves is recommended to calibrate every target and primer/probe design in order to provide accurate viral load determination for patient follow-up.

Nasopharyngeal/throat swabs also have an intrinsic variability that depends on the operator and the tolerance of the patients [18]. Therefore, swab collection quality of every sample must be checked in order to avoid false negatives and provide reliable values. Our results demonstrate that NVL showed robust and similar quantities of SARS-CoV-2 of the $N$ and $RdRp$ genes at two different time-points of the infection in 84 HCWs. Normalization overcame the limitations of Ct value and sample collection variability, providing more accurate values that can guide the monitoring of patients and treatment management.
The gold standard to determine viral infective capacity is viral culture; however, this requires high bio-safety laboratories, fresh samples and long time for response. SARS-CoV-2 culture and sgRNA have been used to detect replication-competent virus longitudinally in respiratory samples [12], presenting a high association between the two techniques [19].

Our study aimed to determine the potential utility of sgRNA for monitoring actively replicating virus in the context of HCW to improve work reincorporation policies. Our findings showed sgRNA was mainly detectable in specimens collected ≤7 days after symptom onset. This result is in correlation with several culture studies describing SARS-CoV-2 culture recovery from respiratory specimens obtained 1-9 days after onset in patients with mild to moderate disease [20, 21]. Our results also showed that after 7 days of symptoms only 7 patients presented sgRNA, indicating that a reduction of isolation protocols should be evaluated to ensure adherence and to vacate COVID-19 hospital isolation rooms.

Subgenomic RNA was also compared with Ct values and NVL obtained by RT-PCR standardized for the N and RdRp genes. Both genes were tested simultaneously together with the cellular quantification by multiplex RT-PCR always using the same procedure and thermocycler. This explains why sgRNA results significantly correlated with Ct values in this assay and should be standardized in every laboratory before using. To achieve less sample and technique variability, it is more optimal to use NVL. Considering the significant association between samples with ≥4log10 RNA copies being sgRNA-positive and <1log10 being sgRNA-negative, a viral load cut point between 1-3log10 RNA copies/1000cells could be setted to request sgRNA detection.

A correlation between chest x-rays, comorbidities or serology with NVL or sgRNA has not previously been reported, but our multivariant analysis showed they are not significantly correlated. Our results confirmed that fever and GID are the only two clinical parameters associated with higher viral loads and replication-competent virus (sgRNA positivity) in the case of fever. Furthermore, dysgeusia is likely to be a delayed symptom associated with lower copies of the virus and a tendency to less
replicative capacity. No other COVID-19 related symptoms, gender or age were significantly correlated with SARS-CoV-2 viral load, in accordance with previous reports [15].

In spite of satisfactory sgRNA results, further viral culture studies are required to verify the association of sgRNA with viral replicative capacity and to define the timing of infectivity. In addition, the study population analyzed was not wide enough and did not include immunocompromised or severely ill patients, in where NVL and sgRNA should be tested to improve patient monitoring and infection control in the hospital setting, and to verify treatment efficacy.

NVL and sgRNA detection are two rapid and accessible techniques that can be easily implemented in routine hospital practice providing a useful proxy for infectivity and patient follow-up. These tools can overcome the obstacles of Ct values, inadequately used in clinical specimens to establish criteria for active infection and transmission, and the costs of high biosafety laboratory equipment and long-term of response limitation of viral culture. Subgenomic RNA RT-PCR can provide essential information for safe work reincorporation of infected workers, patient assessment, discontinuing isolation precautions and optimization of vaccines and treatment approaches.
ACKNOWLEDGMENT

We thank Albert Gabarrus from the Fundació Clínic per la Recerca Biomèdica (FCRB), Hospital Clínic of Barcelona (Spain) for his contribution to statistical analysis; and Donna Pringle for her contribution to English language editing.

Potential conflict of interest

All no reported conflicts. All authors have submitted the ICMJE Form for Disclosure of Conflicts of Interest.

Financial support

No funding was received.
REFERENCES

1. Zhu N, Zhang D, Wang W, Li X, et al. A Novel Coronavirus from Patients with Pneumonia in China, 2019. N Engl J Med 2020; 382:727-733. doi: 10.1056/NEJMoa2001017.

2. World Health Organization Coronavirus disease (COVID-19) situation report. 9 March 2021.

3. Tom MR, Mina MJ. To Interpret the SARS-CoV-2 Test, Consider the Cycle Threshold Value. Clin Infect Dis 2020 [Online] doi: 10.1093/cid/ciaa619

4. Van Kasteren PB, Van der Veer B, Van den Brink S, et al. Comparison of seven commercial RT-PCR diagnostic kits for COVID-19. J Clin Virol, 2020; 128: 104412. doi: 10.1016/j.jcv.2020.104412

5. Rogers AA, Baumann RE, Borillo GA, et al. Evaluation of Transport Media and Specimen Transport Conditions for the Detection of SARS-CoV-2 Using Real Time Reverse Transcription PCR. J Clin Microbiol, 2020; 58 (8). doi: 10.1128/JCM.00708-20.

6. Piralla A, Giardina F, Rovida F, Campanini G, Baldanti F. Cellular DNA quantification in respiratory samples for the normalization of viral load: a real need?. Clin Infect Dis. 2018; 107: 6-10. doi: 10.1016/j.jinf.2018.07.010

7. Xiao AT, Tong YX, Gao C, Zhu L, Zhang YJ, Zhang S. Dynamic profile of RT-PCR findings from 301 COVID-19 patients in Wuhan, China: a descriptive study. J Clin Virol. 2020;127:104346 [Online] doi: 10.1016/j.jcv.2020.104346.

8. WHO Coronavirus disease (COVID-19) situation report – 73. 2 April 2020.

9. Sia SF, Yan LM, Chin AW, Fung K, Choy KT, Wong AY, et al. Pathogenesis and transmission of SARS-CoV-2 in golden Syriam hamsters. Nature. 2020; 583 (7818): 834-838. doi: 10.1038/s41586-020-2342-5

10. Escors, D., Izeta, A., Capiscol, C. & Enjuanes, L. Transmissible gastroenteritis coronavirus packaging signal is located at the 5' end of the virus genome. J Virol. 2003; 77: 7890-7902. Doi: 10.1128/JVI.77.14.7890-7902.2003
11. Dagotto G, Mercadoa N, Martinezc D, Houc Y, Nkololaa J, Carnahand R. Comparison of Subgenomic and Total RNA in SARS-CoV-2 Challenged Rhesus Macaques. J Virol. 2021. [Online] doi:10.1128/JVI.02370-20

12. Wölfel R, Corman VM, Guggemos W, Seilmaier M, Zange S, Müller MA, et al.. Virological assessment of hospitalized patients with COVID-2019. Nature. 2020; 581 (7809): 465-469. doi: 10.1038/s41586-020-2196-x.

13. Nicolás D, Camós-Carreras A, Spencer F, Arenas A, Butori E, Maymó P. A prospective cohort of SARS-COV2 infected health care workers: Clinical characteristics, outcomes and follow up strategy. Open Forum Infect Dis. 2020 [Online] doi: 10.1093/ofid/ofaa592

14. Corman VM, Landt O, Kaiser M, Molenkamp R, Meijer A, Chu DK, et al. Detection of 2019 novel coronavirus (2019-nCoV) by real-time RT-PCR. Euro Surveill. 2020; 25(3):2000045. doi: 10.2807/1560-7917.ES.2020.25.3.2000045

15. Tom MR, Mina MJ. To interpret the SARS-CoV-2 test, consider the cycle threshold value. Clin Infect Dis. 2020 [Online] doi: 10.1093/cid/ciaa619

16. Infectious Diseases Society of America (IDSA) and Association for Molecular Pathology (AMP) joint statement on the use of SARS-CoV-2 PCR cycle threshold (Ct) values for clinical decision-making - Updated March 12, 2021

17. Rhoads D, Peaper DR, She RC, Nolte FS, Wojewoda CM, Anderson NW, Pritt BS. College of American Pathologists (CAP) Microbiology Committee Perspective: Caution Must Used in Interpreting the Cycle Threshold (Ct) Value. Clin Infect Dis 2020. doi: 10.1093/cid/ciaa1199

18. Zhou Y, Pei F, Wang L, Zhao H, Li H, Ji M, et al. Sensitivity Evaluation of 2019 Novel Coronavirus (SARS-CoV-2) RT-PCR Detection Kits and Strategy to Reduce False Negative. PloS One, 2020; 15 (11) [Online] doi: 10.1371/journal.pone.0241469

19. Perera RAPM, Tso E, Tsang OTY, Tsang DNC, Fung K, Leung YWY. SARS-CoV-2 Virus Culture and Subgenomic RNA for Respiratory Specimens from Patients with Mild Coronavirus
Disease. Emerg Infect Dis. 2020 Nov; 26(11):2701-2704. doi: 10.3201/eid2611.203219. Epub 2020.

20. The COVID-19 Investigation Team. Clinical and virologic characteristics of the first 12 patients with coronavirus disease 2019 (COVID-19) in the United States. Nat Med. 2020; 26(6):861–868 [Online] Available from: http://www.nature.com/articles/s41591-020-0877-5. doi: 10.1038/s41591-020-0877-5.

21. Arons MM, Hatfield KM, Reddy SC, et al. Presymptomatic SARS-CoV-2 infections and transmission in a skilled nursing facility. N Engl J Med. 2020; 382(22):2081–2090. doi: 10.1056/NEJMoa2008457.
Table 1. Clinical description of SARS-CoV-2 infected Health-care workers cohort.

| Characteristics                        |       |
|----------------------------------------|-------|
| Age, years (median, IQR)               | 33.5  |
| Female, sex, n (%)                     | 58 (69%) |
| Current smokers, n (%)                 | 3 (3.6%) |
| Comorbidities*, n (%)                  | 12 (14.3%) |

| Clinical features                        |       |
|----------------------------------------|-------|
| Symptomatic at diagnosis, n (%)         | 74 (88.1%) |
| Days from symptoms onset until 1<sup>st</sup> PCR (median, IQR) | 3 (1; 6) |
| Days from symptoms onset until 2<sup>nd</sup> PCR (median, IQR) | 15 (14; 19) |
| Symptoms n (%)                          | 77 (91.7%) |
| - Cough, n (%)                          | 56 (66.7%) |
| - Fever, n (%)                          | 23 (27.4%) |
| - Dyspnea, n (%)                        | 10 (11.9%) |
| - Hyposmia/anosmia, n (%)               | 35 (41.7%) |
| - Dysgeusia, n (%)                      | 26 (31%) |
| - Gastrointestinal disorders, n (%)     | 17 (20.2%) |
| Asymptomatic, n (%)                     | 7 (8.3%) |
| Treatment, n (%)                        | 5 (6%) |

Serology SARS-CoV-2 IgG
- Positive, n (%) 80 (98.8%)
- Negative, n (%) 1 (1.2%)

Days from symptom onset until positive serology (median, IQR) 44 (36; 52)

IQR: Interquartile Range

n: number of patients

*a Comorbidities include dyslipidemia, hypertension, chronic lung disease, ischemic heart disease, chronic kidney disease and neoplasm.
Table 2. RT-PCR Ct values and normalized viral loads of the SARS-CoV-2 RdRp and N genes.

|            | RdRp gene |                  | N gene |                  |
|------------|-----------|------------------|--------|------------------|
|            | Ct value  | NVL              | Ct value| NVL              |
| 1st RT-PCR | 26.58 (22.36; 28.99) | 2.52E+03 (4.23E+02; 7.96E+04) | 26.65 (22.07; 32.52) | 4.00E+03 (1.38E+02; 1.24E+05) |
| 2nd RT-PCR | 29.99 (27.63; 31.95) | 7.55E+01 (1.85E+01; 2.45E+02) | 34.25 (30.77; 35.56) | 1.61E+01 (7.37E+00; 4.05E+01) |
| Subtraction| 3.71 (0.34; 7.62) | 1.78 log10 (0.59; 3.15) | 5.71 (1.32; 11.05) | 2.1 log10 (0.95; 3.74) |

RdRp: RNA dependent RNA polymerase
N: Nucleocapsid
Ct: Cycle Threshold; median (Interquartile Range)
NVL: Normalized viral load; median of RNA copies / 1000 cells (Interquartile Range).
Subtraction: indicates the difference of the 2nd PCR minus the 1st PCR values.
Table 3. Association between subgenomic RNA and N and RdRp RT-PCR Ct values and normalized viral loads of the 1st and 2nd RT-PCR.

| Detection Gen | n | Variable | Total | Negative sgRNA | Positive sgRNA | P-value |
|---------------|---|----------|-------|----------------|----------------|---------|
|               | n |          |       |                |                |         |
| 1st RT-PCR    | n | 81       | 38    |                |                | <0.001  |
| N             | Ct value | 26.65 (22.07; 32.52) | 32.66 (30.23; 34.53) | 22.54 (17.94; 25.09) | <0.001 |
|              | NVL | 4000 (138.05; 124130.88) | 633.64 | 380403.46 | <0.001 |
|              | Log10 |                     |       |                |                |         |
| 2nd RT-PCR    | n | 79       | 74    |                |                | 0.010   |
| RdR           | Ct value | 34.25 (30.77; 35.56) | 34.46 (31.88; 35.63) | 29.07 (27.5; 30.07) | 0.010  |
|              | NVL | 16.16 (7.37; 40.53) | 15.36 (7.14; 32.29) | 1502.86 (477.19; 2624.29) | 0.001  |
|              | Log10 |                     |       |                |                |         |
|              | NVL | 1.21 (0.87; 1.61) | 1.19 (0.85; 1.51) | 3.18 (2.68; 3.42) | 0.001  |
| RdR           | n | 84       | 79    |                |                |         |
| Detectio n Gen e Variable | Total | Negative sgRNA | Positive sgRNA | p-Ct value | NVL | Log10 NVL | P-value |
|--------------------------|-------|---------------|----------------|-------------|-----|-----------|---------|
| Ct value                 |       |               |                |             |     |           | 0.135   |
| 29.99 (27.63; 31.95)    | 30.06 (27.71; 31.98) | 27.32 (25.68; 30.1) | 0.135       |
| NVL                      |       |               |                |             | 75.55 (18.47; 245.64) | 73 (15.51; 236) | 1247 (25.7; 2914.29) | 0.123   |
| 1.88 (1.26; 2.39)        | 1.86 (1.19; 2.37) | 3.1 (1.41; 3.46) |               |             |     |           | 0.123   |

n: number of samples

Ct value: cycle threshold value; median (interquartile range)

NVL: median (interquartile range) of linear (RNA copies / 1000 cells) and logarithmic scales (log10 RNA copies /1000 cells)

P-values obtained from the Mann-Whitney test. P-values < 0.05 are indicated in bold.
Table 4. Subgenomic RNA of the 1\textsuperscript{st} and 2\textsuperscript{nd} RT-PCR results divided by intervals of days after symptom onset.

| Days of symptoms | Positive sgRNA | Negative sgRNA | Total |
|------------------|----------------|----------------|-------|
| 1-5              | 34 (68\%)     | 16 (32\%)      | 50    |
| 6-7              | 3 (37.5\%)    | 5 (62.5\%)     | 8     |
| 8-10             | 1 (10\%)      | 9 (90\%)       | 10    |
| 11-14            | 2 (5.9\%)     | 32 (94.1\%)    | 34    |
| >14              | 4 (8.7\%)     | 42 (91.3\%)    | 46    |
Table 5. Significant associations between normalized SARS-CoV-2 viral loads and subgenomic RNA with categorical variables in the total study population.

| Symptom | Variable | Statistic | Total | Yes | No | P value |
|---------|----------|-----------|-------|-----|----|---------|
|         |          | n         | n     |     |    |         |
| Fever   | Log10 NVL - 1st RT-PCR | Median | 81  | 58  | 23 | 0.002   |
|         |          | (IQR)     | 3.6 (2.14; 5.09) | 3.19 (2.08; 5.18) | 5.18 (2.71; 5.8) |         |
|         | Positive sgRNA | n (%) | 43 (51.2%) | 26 (46.6%) | 17 (73.9%) | 0.011   |
|         | Negative sgRNA | n (%) | 41 (48.8%) | 35 (57.4%) | 6 (26.1%)  |         |
| GID     | Log10 NVL - 1st RT-PCR | Median | 81  | 65  | 17 | 0.007   |
|         | (IQR)     | 3.6 (2.14; 3.2 (2.08; 5.03 (3.26; 5.82) |         |
|         | Positive sgRNA | n (%) | 43 (51.2%) | 31 (46.3%) | 12 (70.6%) | 0.073   |
|         | Negative sgRNA | n (%) | 41 (48.8%) | 36 (53.7%) | 5 (29.4%)  |         |
| Dysgeusia | Log10 NVL - 2nd RT-PCR | Median | 84  | 58  | 26 | 0.006   |
|         | (IQR)     | 1.88 (1.26; 2.04 (1.56; 1.44 (0.81; 1.91) |         |
|         | Positive sgRNA | n (%) | 5 (6%)  | 5 (8.6%) | 0 (0%)  | 0.318   |
|         | Negative sgRNA | n (%) | 79 (94%) | 53 (91.4%) | 26 (100%) |         |

GID: Gastrointestinal Disorders.
NVL: normalized viral loads (log10 RNA copies /1000 cells)
sgRNA: subgenomic RNA
n: number of samples
IQR: Interquartile Range

P-values obtained from the Mann-Whitney test. P-values<0.05 are indicated in bold.
FIGURE LEGENDS

Figure 1. Subgenomic RNA detection compared with normalized viral loads. Normalized viral loads were quantified as log10 RNA copies per 1000 cells. Subgenomic RNA was qualitatively determined as positive or negative by RT-PCR. The first and second sample determinations of each patient are shown.

Figure 2. Normalized SARS CoV-2 N and RdRp viral loads and sgRNA of the first and second detection compared with days of symptoms. A. Viral loads were measured by RT-PCR and normalized to RNA copies per 1000 cells. Quantification was performed with the RNA dependent RNA polymerase (RdRp) and Nucleocapsid (N) genes in each specimen. B. Subgenomic RNA transcripts in relation to genomic RNA Ct values are expressed as $2^{(Ct_{gRNA} - Ct_{gRNA})}$ and plotted against days of symptoms.
Figure 1

- Positive sgRNA
- Negative sgRNA

Viral Load (RNA copies / 1000 cells)
Figure 2