Evaluation of a home-based 7-day infection control strategy for healthcare workers following high-risk exposure to severe acute respiratory coronavirus virus 2 (SARS-CoV-2): A cohort study

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Abstract

Objective: Evidence-based infection control strategies are needed for healthcare workers (HCWs) following high-risk exposure to severe acute respiratory coronavirus virus 2 (SARS-CoV-2). In this study, we evaluated the negative predictive value (NPV) of a home-based 7-day infection control strategy.

Methods: HCWs advised by their infection control or occupational health officer to self-isolate due to a high-risk SARS-CoV-2 exposure were enrolled between May and October 2020. The strategy consisted of symptom-triggered nasopharyngeal SARS-CoV-2 RNA testing from day 0 to day 7 after exposure and standardized home-based nasopharyngeal swab and saliva testing on day 7. The NPV of this strategy was calculated for (1) clinical coronavirus disease 2019 (COVID-19) diagnosis from day 8 to day 14 after exposure, and for (2) asymptomatic SARS-CoV-2 detected by standardized nasopharyngeal swab and saliva specimens collected at days 9, 10, and 14 after exposure. Interim results are reported in the context of a second wave threatening this essential workforce.

Results: Among 30 HCWs enrolled, the mean age was 31 years (SD, ±9), and 24 (80%) were female. Moreover, 3 were diagnosed with COVID-19 by day 14 after exposure (secondary attack rate, 10.0%), and all cases were detected using the 7-day infection control strategy: the NPV for subsequent clinical COVID-19 or asymptomatic SARS-CoV-2 detection by day 14 was 100.0% (95% CI, 93.1%–100.0%).

Conclusions: Among HCWs with high-risk exposure to SARS-CoV-2, a home-based 7-day infection control strategy may have a high NPV for subsequent COVID-19 and asymptomatic SARS-CoV-2 detection. Ongoing data collection and data sharing are needed to improve the precision of the estimated NPV, and here we report interim results to inform infection control strategies in light of a second wave threatening this essential workforce.

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(NPV) for subsequent COVID-19 diagnosis and for asymptomatic SARS-CoV-2 detection by day 14 after exposure.

We report the interim results of a cohort study evaluating this testing strategy to provide evidence that may help inform infection control strategies in the event of a second wave.

Methods

Study design

A cohort study was initiated May 18, 2020. The study was approved by the Research Ethics Board of the McGill University Health Centre (no. 2020-6565), and written informed consent was obtained from all participants.

Participants and setting

We enrolled HCWs employed at hospital and nursing residences in the greater Montreal metropolitan area (2016 census population, 4,098,927), which had the highest COVID-19 infection rate in Canada throughout the study period (1,078 per 100,000 on May 4, 2020, which had the highest COVID-19 infection rate in the greater Montreal metropolitan area (2016 census population, 4,098,927), which had the highest COVID-19 infection rate in Canada throughout the study period (1,078 per 100,000 on May 4, 2020, which had the highest COVID-19 infection rate in the greater Montreal metropolitan area (2016 census population, 4,098,927), which had the highest COVID-19 infection rate in Canada throughout the study period (1,078 per 100,000 on May 4, 2020, which had the highest COVID-19 infection rate in the greater Montreal metropolitan area (2016 census population, 4,098,927), which had the highest COVID-19 infection rate in Canada throughout the study period (1,078 per 100,000 on May 4, 2020).14

HCWs were eligible if they had had a high-risk exposure to SARS-CoV-2 that required self-isolation within 7 days of eligibility screening. High-risk SARS-CoV-2 exposure requiring self-isolation was defined as (1) being within 2 m of a person with COVID-19 for at least 10 minutes without face mask or eye shield or (2) being determined by the HCW’s institutional infection control or occupational health officer to have been exposed. During the study period, the participants were required to self-isolate for 14 days after exposure. Participants were not eligible if they were actively using anticoagulant medication or were unable to communicate in English or French.

Testing strategy

Enrolled participants were advised to seek clinical testing for SARS-CoV-2 if they developed the following symptoms of COVID-19 from day 0 to day 7 after exposure: fever, significant loss of appetite, major fatigue, general muscle pain unrelated to physical exertion, sudden loss of smell, new or worsening cough, shortness of breath, sore throat, runny or stuffy nose, nausea, vomiting diarrhea, or abdominal pain. On day 7 after exposure, a research staff visited the participant’s self-isolation residence to collect a nasopharyngeal swab and saliva specimen for subsequent SARS-CoV-2 RNA detection by RT-PCR. The testing strategy was considered negative if SARS-CoV-2 RNA was not detected by symptom-triggered clinical testing from day 0 to 7 or the standardized home-based day 7 nasopharyngeal swab and saliva specimens.

Outcome

The primary outcome was clinical COVID-19, defined by the presence of COVID-19 symptoms (defined above) and SARS-CoV-2 RNA detection by RT-PCR from day 8 to day 14 after exposure. The secondary outcome was asymptomatic SARS-CoV-2 RNA detection by RT-PCR on standardized nasopharyngeal swab or saliva specimens, both collected on day 9 or 10, and on day 14 after exposure.

SARS-CoV-2 RNA detection

Symptom-triggered or self-initiated nasopharyngeal swab specimens were collected at public testing centers, and RT-PCR for SARS-CoV-2 RNA was performed within 24 hours. The home-based nasopharyngeal swab and saliva specimens collected on day 7, day 9 or 10, and day 14 after exposure were placed immediately in universal transport media (UTM) and transported on ice to a −80°C freezer within 6 hours of collection. These specimens were thawed once after a median of 74 days, and SARS-CoV-2 RNA RT-PCR was performed in the clinical microbiology laboratory of the McGill University Health Centre using the Cobas SARS-CoV2 Test on the Cobas 6800 System (Roche Diagnostics, Indianapolis, IN), the same system used for clinical care. A pilot-tested, home-collected specimen subjected to the same freeze–thaw duration confirmed that the detection of SARS-CoV-2 RNA was possible.

Statistical analysis

The NPV of the testing strategy was calculated as the number of participants with a negative testing strategy result who did not experience the outcome divided by the number of participants with a negative testing strategy and expressed as a percentage. The confidence interval was calculated with an α of 0.05 using the binom.confint function in the R version 4.0.2 statistical software package (R Foundation for Statistical Computing, Vienna, Austria).

Sample size

To inform infection control strategies in the context of a pandemic that threatens an essential workforce, we sought to maximize the precision of the lower bound of the estimated NPV confidence interval. Assuming a secondary attack rate of 10%, a testing strategy NPV of 100%, and an α of 0.05, enrollment of 50 participants would yield a lower-bound NPV of 96%, and 100 participants would yield a lower-bound NPV of 98%.

Results

Interim study results are presented in the context of an anticipated imminent second wave and the lack of prospective data assessing the performance of infection control strategies for essential service providers following high-risk exposure to SARS-CoV-2.

To date, 46 participants have been screened and 16 were ineligible: 5 participants were outside the 7-day postexposure enrollment period, 2 did not have a high-risk exposure, and 8 expressed interest via the online screen function but did not provide contact information. One participant was eligible and consented to participating but withdrew prior to home testing on day 7. Among the 30 HCWs enrolled, the mean age was 31 years (SD ±9) and 24 (80.0%) were female. Furthermore, 28 of the high-risk exposures (93.3%) occurred at the workplace.

In total, 151 specimens (nasopharyngeal swab and saliva) were collected by research staff as part of the testing strategy, and 30 symptom-triggered or self-initiated clinical tests were sought by participants (Fig. 1). From day 0 to day 7, 18 symptom-triggered or self-initiated clinical nasopharyngeal swab tests were obtained from 14 participants. Also, 49 home-based nasopharyngeal swab and saliva specimens were collected on day 7 by the research staff. Furthermore, 2 participants were symptomatic on day 7 and sought clinical nasopharyngeal swab testing, and 1 participant declined the nasopharyngeal swab but provided a saliva specimen. Primary and secondary outcome ascertainment from day 8 to day 14 included 13 symptom-triggered or self-initiated clinical nasopharyngeal swab specimens, 50 standardized nasopharyngeal swab and saliva specimens collected on days 9 and 10, and 52 specimens were collected on day 14. Furthermore, 3 participants declined the swab on day 9 or 10 and 2 participants declined the swab day 14.
but they provided saliva specimens, and 1 participant provided a home-based nasopharyngeal swab but not a saliva specimen on day 9 or 10.

In addition, 3 participants were diagnosed with COVID-19 by day 14 after exposure, resulting in a secondary attack rate of 10.0%, with all cases detected by the testing strategy (all via symptom-triggered from day 0 to day 7). The NPV of the testing strategy for COVID-19 diagnosis between days 8 and day 14 was 100.0% (95% CI, 93.1%–100.0%), and for asymptomatic SARS-CoV-2 detection, the NPV was 100.0% (95% CI, 93.1%–100.0%).

Discussion

A home-based 7-day infection control strategy for HCWs following high-risk exposure to SARS-CoV-2 may have high NPV for subsequent COVID-19 and asymptomatic SARS-CoV-2 detection. Acknowledging the limited precision of the NPV estimate obtained from an interim analysis, this finding may inform local infection control policies in the event that the second wave threatens this essential workforce.

Infection control strategies for HCWs exposed to SARS-CoV-2 are guided by transmission risk and incubation time, but few have been prospectively evaluated using standardized outcome ascertainment. The present study begins to address this knowledge gap by showing that a relatively simple testing strategy may have sufficiently high NPV to allow HCWs with high-risk exposure to SARS-CoV-2 to return to work with low risk of exposure-related COVID-19 or asymptomatic SARS-CoV-2 transmission.

The present study must be considered in light of its weaknesses. First, the interim analysis resulted in a confidence interval surrounding the NPV estimate and may include unacceptably low values to merit implementation as an infection control strategy. We nevertheless present these interim results due to the lack of prospective data evaluating postexposure HCW testing strategies and because a second wave threatens this essential workforce. With ongoing data collection and data sharing, we expect the precision of the NPV estimate to increase. Second, although the secondary attack rate in this sample is consistent with earlier close-contact estimates, evolving HCW personal protective equipment policies may lower secondary attack rates in the second wave. If true, this would make our NPV estimate conservative. Third, SARS-CoV-2 immunity was not assessed but may influence the NPV of the testing strategy and the generalizability of our findings. Although the research has not yet been peer reviewed, the prevalence of SARS-CoV-2 IgG antibody detection among HCWs at another Canadian tertiary-care center during the same study period was low (1.4%–3.4%). If immunity can be achieved through infection or vaccination, immunity will likely increase with time, making our NPV estimate conservative. Fourth, we did not perform viral culture to confirm the infectivity of participants with SARS-CoV-2 RNA detected by RT-PCR. Although we designed our pragmatic study design to evaluate the testing strategy under ‘real-world’ conditions (ie, with widely available clinical tools), the assumption that any participant with SARS-CoV-2 RNA detected is infectious would make our NPV estimate conservative in terms of transmission risk. Fifth, the single and prolonged freeze–thaw cycle in this study, required due to pandemic prioritization of clinical specimen processing, may have influenced RT-PCR sensitivity. We think this is unlikely (1) because prolonged storage at −80°C with a single freeze-thaw cycle has generally not been considered deleterious to detection of enveloped RNA viruses from nasopharyngeal specimens and (2) because a recent evaluation of a single freeze–thaw cycle on SARS-CoV-2 RNA detection using the same assay resulted in 0 of 66 positive samples converting to negative and only 4 of 66 (6%) converting to inconclusive.

Finally, household exposure risk may vary over time, as may adherence to self-isolation and risk of secondary infection to infection control recommendations from public health authorities. The NPV was estimated from a sample of HCWs studied during a period when schools and businesses were closed, possibly lowering the risk of household exposure during self-isolation. Further evaluation or application of this infection control strategy should consider adherence to self-isolation and risk of secondary high-risk SARS-CoV-2 exposure.

In conclusion, among HCWs with high-risk exposure to SARS-CoV-2, a home-based 7-day infection control strategy may have a high NPV for subsequent COVID-19 and asymptomatic SARS-CoV-2 detection. These interim results are reported in light of a second wave that threatens healthcare workforce capacity in some regions. Ongoing data collection and data sharing will increase the precision of the estimated NPV and better inform infection control strategies.

Data sharing

The data from this study can be made available, provided appropriate ethics approvals and data use agreements, by contacting the corresponding author. Researchers with datasets that include standardized SARS-CoV-2 RNA testing 7 days post exposure are encouraged to contact the corresponding author to conduct an individual participant data meta-analysis.

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