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Self-collection: An appropriate alternative during the SARS-CoV-2 pandemic

Michael C. Wehrhahn\textsuperscript{a,1,*}, Jennifer Robson\textsuperscript{b,1}, Suzanne Brown\textsuperscript{c}, Evan Bursle\textsuperscript{b}, Shane Byrne\textsuperscript{b}, David New\textsuperscript{d}, Smathi Chong\textsuperscript{d}, James P. Newcombe \textsuperscript{a}, Terri Siversten\textsuperscript{d}, Narelle Hadlow\textsuperscript{d}

\textsuperscript{a} Douglass Hanly Moir Pathology, 14 Giffnock Ave, Macquarie Park, NSW, 2113, Australia
\textsuperscript{b} Sullivan Nicolaides Pathology, 24 Hurworth St, Bowen Hills, QLD, 4006, Australia
\textsuperscript{c} Department of Endocrinology & Diabetes, Sir Charles Gairdner Hospital, Hospital Ave, Nedlands, WA, 6009, Australia
\textsuperscript{d} Clinipath Pathology, 310 Selby St, North Osborne Park, WA, 6017, Australia

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\textbf{ABSTRACT}

\textbf{Objectives:} To evaluate the reliability of self-collection for SARS-CoV-2 and other respiratory viruses because swab collections for SARS-CoV-2 put health workers at risk of infection and require use of personal protective equipment (PPE).

\textbf{Methods:} In a prospective study, patients from two states in Australia attending dedicated COVID-19 collection clinics were offered the option to first self-collect (SC) nasal and throat swabs (SCNT) prior to health worker collect (HC) using throat and nasal swabs (Site 1) or throat and nasopharyngeal swabs (Site 2). Samples were analysed for SARS-CoV-2 as well as common respiratory viruses. Concordance of results between methods was assessed using Cohen's kappa ($\kappa$) and Cycle threshold (Ct) values were recorded for all positive results as a surrogate measure for viral load.

\textbf{Results:} Of 236 patients sampled by HC and SC, 25 had SARS-CoV-2 (24 by HC and 25 by SC) and 63 had other respiratory viruses (56 by HC and 58 by SC). SC was highly concordant with HC ($\kappa=0.890$) for all viruses including SARS-CoV-2 and more concordant than HC to positive results by any method ($\kappa=0.959$ vs 0.933).

Mean SARS-CoV-2 E-gene and N-gene, rhinovirus and parainfluenza Ct values did not differ between HC and SCNT.

\textbf{Conclusions:} Self-collection of nasal and throat swabs offers a reliable alternative to health worker collection for the diagnosis of SARS-CoV-2 and other respiratory viruses and provides patients with easier access to testing, reduces exposure of the community and health workers to those being tested and reduces requirement for PPE.

\section{Introduction}

On the 11th March 2020, the World Health Organisation (WHO) announced COVID-19 as a pandemic.\textsuperscript{[1]} The WHO Director-General issued a call for urgent action and encouraged all countries to ‘innovate and learn’ in their response to this crisis.

Self-collected swabs in the community for SARS-CoV-2, the agent of COVID-19, and for other respiratory viruses offers potential significant benefit in the current pandemic by reducing requirement for PPE, limiting exposure of patients and staff to infection, increased convenience and access for patients and timeliness of a sample receipt.\textsuperscript{[2,3]} Patients report self-collected nasal swabs are easy to perform\textsuperscript{[2,4,5]} and highly acceptable\textsuperscript{[2,4]}. A meta-analysis of 9 studies comparing self-collect (SC) and health worker collect (HC) for influenza testing reported a pooled sensitivity of 87 % and specificity of 99 % for SC compared to HC\textsuperscript{[6]}. Irving et al studied paired samples from 240 adults and found sensitivity using nasal or nasopharyngeal (NP) collection for influenza did not vary significantly when using a highly sensitive molecular test\textsuperscript{[7]}. A study in 230 children reported equivalent sensitivity for all respiratory viruses except respiratory syncytial virus (RSV) when comparing nasal swab and NP aspirate\textsuperscript{[8]}. Larios et al demonstrated that using flocked swabs and sensitive molecular methods, equivalent sensitivity and specificity was obtained for matched self-collected mid-turbinate nasal swabs and NP swabs in 38 individuals for a range of respiratory viruses including human coronaviruses\textsuperscript{[9]}.

Recent reports on SARS-CoV-2 in respiratory specimens indicate early high viral loads in symptomatic and asymptomatic patients in a variety of clinical specimens including nasal and throat swabs, sputum and saliva samples\textsuperscript{[10–14]}. Wang et al reported that in 205 patients
with COVID-19 the highest positive rates were found from bronchial-alveolar lavage fluid, sputum and nasal swabs respectively [15]. Wolfel et al reported that in hospitalized cases of COVID-19 there was no discernible difference between NP and throat swabs with high viral load present in both specimens early in the illness and suggested that simple throat swabs may provide sufficient sensitivity when patients are first tested with mild symptoms of COVID-19 [14].

The aim of this study was to compare prospectively the performance of HC with separate SC nasal (SCN) and throat swabs (SCT) and the combination of the two (SCNT) for respiratory viruses including SARS-CoV-2.

2. Methods

This study was conducted across two laboratory sites (Site 1 and Site 2) and had ethics approval with all participants providing informed consent. For a period of one week in March 2020, patients presenting for SARS-CoV-2 testing at dedicated COVID-19 collection rooms were offered participation in the study. Demographic data was recorded including the address and postcode to assess the Index of Education and Occupation (IEO) which assesses education level based on a scale of 1 to 5 with 5 being the highest level of education. [16] A questionnaire assessing acceptability of SC based on that of Akmatov was provided to patients [4]. Printed instructions including diagrams were provided on how to collect throat and nasal swab (See Supplementary Information). Self-collection kits included two swab packets each containing a single swab and screw-top container with 2ml liquid Amies medium, a tongue depressor and a zip lock sample bag. SC samples were taken immediately prior to trained HC collects to reduce ‘training bias.’ For SC and HC at Site 1 and SC at Site 2, open-cell polyurethane foam swabs (Σ Transwab® ref MW940, Medical Wire & Equipment (MWE), Wiltshire, England) were used. Throat swabs were collected from the posterior throat and tonsil areas while nasal swabs were inserted as far as comfortably possible and at least 2–3 cm inside one nostril, rotating the swab 5 times and leaving in place for 5–10 seconds. For HC at Site 2, a flocked NP swab and a foam throat swab (Σ Transwab® ref MW819 and MW940) were used. In addition, because the expected SARS-CoV-2 positivity rate at the time was estimated to be less than 1%, a subset of 24 patients recently diagnosed with COVID-19 performed SC in their homes.

At site 1, testing for SARS-CoV-2 was on the Alplex™ 2019-nCoV assay (Seegene, Seoul, South Korea) and followed sample extraction using MagNA Pure 96 (Roche, Basel, Switzerland) with amplification utilising CFX96 Touch RT-PCR Detection Systems (BioRad, Hercules, California USA). Samples were confirmed as SARS-CoV-2 positive if all three gene targets (E/RdRp and N genes) were detected within 40 cycles. At site 2, the same extraction method was used. Testing for SARS-CoV-2 was performed using an in-house developed Taqman assay, was detected only on SCN but not the HC. A second positive patient retested 6 days after symptom onset using the screening E-gene assay, was detected only on SCN and not the HC. A positive result on either HC or SCNT was included in the group AP.

Table 2 summarises the respiratory viruses detected by the different methods of collection. At Site 1, co-detection of rhinovirus (Ct 29) + influenza A (Ct 41) was found in one patient by SC only and RSV (Ct 24) + rhinovirus (Ct 35) in one patient by HC only. Two parainfluenza cases and one rhinovirus case were detected only by SC. Overall the detection rate was 6% higher in SC compared with HC swabs for non-SARS-CoV-2 respiratory viruses which equated to 3/20 (15%) additional positive results. At Site 2, no co-detections occurred. Collection of samples for the 13 SARS-CoV-2 positive patients ranged from 2 to 9 days following onset of symptoms with a mean of 4.8 days. One positive patient retested 6 days after symptom onset using the screening E-gene assay, was detected only on SCN but not the HC. The second positive patient was detected using HC and SCT but not SCN. Of the patients with detectable respiratory viruses other than SARS-CoV-2, at site 1, 8/23 (35%) had virus only detectable on one of SCN or SCT while the proportion was 14/35 (40%) at site 2.

When all detections by HC and SCNT were compared with AP, the sensitivity of SCNT and HC to detect SARS-CoV-2 was 1.0 (95% CI: 0.86-1) and 0.96 (95% CI: 0.8-1) respectively; for other respiratory viruses it was 0.94 (95%CI: 0.87-0.98) and 0.9 (95% CI: 0.83-0.96) respectively.

Table 3 summarises concordance between AP and each collection method. Both SCNT and HC showed high concordance with AP at each site and overall, with SCNT slightly higher (κ = 1.0, 0.934, 0.959 at Site 1, Site 2, Combined Sites) than HC (κ = 0.929, 0.934, 0.933). Additionally, SCNT was highly concordant with HC (κ = 0.929, 0.863, 0.890 at Site 1, Site 2, Combined Sites). When Ct values for COVID-19
The mean Ct differed between HC and SCN (β = 4.67, p = 0.014) but not with HC (p = 0.036; α' = 0.017) but was higher in SCN and SCT. The mean Ct value was not significantly higher in SCNT compared with HC and SCNT (p = 0.041; α' = 0.008) but was higher in SCN and SCT (β = 7.31, p < 0.001). Mean N-gene Ct value did not differ against α' = 0.008) but was significantly higher in SCT compared with HC and SCNT (p = 0.041; α' = 0.008). Concordance (Cohen's κ) between (i) AP and HC, SCN, SCT and SCNT; and (ii) HC and SCNT. A value of 1 indicates the method detected all COVID-19 and other respiratory cases, while a value above 0.9 indicates a very high level of detection of all respiratory cases (AP).

Table 3
Concordance (Cohen’s κ) between (i) AP and HC, SCN, SCT and SCNT; and (ii) HC and SCNT. A value of 1 indicates the method detected all COVID-19 and other respiratory cases, while a value above 0.9 indicates a very high level of detection of all respiratory cases (AP).

| Site | HC | SCN | SCT | SCNT | AP |
|------|----|-----|-----|------|----|
| Site 1 | 0.929 | 0.905 | 0.872 | 1 |
| Site 2 | 0.934 | 0.835 | 0.789 | 0.934 |
| Combined Sites | 0.933 | 0.858 | 0.817 | 0.959 |

Discussion
In our group of 236 ambulatory, literate, mostly adult patients, the performance of self-collected nasal and throat swabs was at least equivalent to that of health worker collected swabs for the detection of SARS-CoV-2 and other respiratory viruses.

This study included two different sites using two different methods of HC (combined N + T and combined NP + T) and also employed two different molecular strategies for detection of SARS-CoV-2. As such these findings are more widely applicable.

At Site 1 where SCNT was compared with HC using the same swab and collection methods, for the 12 patients testing positive to SARS-CoV-2 there was complete concordance between HC and SCNT samples even though on average 2.5 days had lapsed. In the remaining SARS-CoV-2 negative patients, SCNT detected 3 additional respiratory viruses, with the overall positivity rate increasing from 34 % to 40 %. However, the additional 3 SC detections were weak positives based on high Ct values (33–40).

At site 2 where comparative HC involving a NP and T swab occurred at the same time as the HC and SCT for the SARS-CoV-2 positive patients, SCNT detected all 13 positive patients while one patient was negative by HC. Detection of other respiratory viruses by SCNT was highly concordant with HC detecting only 1 less respiratory virus and may relate to the fact that SCNT sampling was compared with NP + T sampling.

When data from each site was combined, concordance between SCNT or HC with the All Positive (AP) rate was very high, slightly favouring SCNT. The similar SARS-CoV-2 percent-positivity rate in ongoing comparison data between those having only HC or SC provides further reassurance that SCNT is equivalent to HC.

The advantages of self-collection are even more important at a time of global health crisis. Self-collection greatly reduces the number of patients requiring trained health workers and the necessary PPE to protect them. Access to testing is increased, as swab kits can be provided quickly by clinicians or available at dedicated COVID-19 collection centres aiding timeliness of testing [3] which is critical in the current pandemic. [2,3] Safety for both patients and staff using a SC model is also increased as exposure to others is limited.

Further, data from patients at site 1 suggests that SC is accessible and achievable over a range of education levels with all finding SC acceptable and the majority having a preference for this method over HC as has previously been reported. [2,4,5] This may relate to the ability of patients to control the comfort level of throat and nasal collection better than a trained collector can.

Recent studies suggest there is a high viral load in patients with early COVID-19 across the upper and lower respiratory tracts, including nasal and throat sites [10–12,14] as well as in saliva [13], even in asymptomatic, mild or prodromal states. Wofelel al noted no discernible difference between nasopharyngeal and oropharyngeal viral loads in hospitalized cases of COVID-19 [14]. Given these high viral loads throughout the respiratory tract it may be that requiring NP sampling is not as significant for SARS-CoV-2 as for some other...
respiratory viruses. It may also be that PCR methods for viral detection are improving the sensitivity of a range of sample and collection methods as shown for a range of respiratory viruses but also Group A Streptococcal detection [9,20]. We hypothesize that the high viral load of SARS-CoV-2 and sensitive molecular techniques may explain the equivalent sensitivity of SC to HC samples in COVID-19 patients.

Our data support the decision by the Public Health Laboratory Network of Australia (PHLN) [21] to recommend sampling of both nasal and throat sites for the diagnosis of respiratory viruses including for SARS-CoV-2, due to the concern of a possible missed diagnosis if only one site is sampled. This was the case for two COVID-19 positive patients on SC who were only diagnosed by SCN and another only by SCT. If only one swab site was obtainable, our data suggests the nasal may be the better swab site for the diagnosis of COVID-19 as it had greater concordance with the AP group and showed consistently lower Ct values in the order of 100–1000 fold higher viral load (data not shown). Furthermore, we have instituted use of a single swab to sample both throat then nasal sites. This has the potential to preserve limited supplies of swabs and also provide additional efficiencies in the laboratory as only preparation of a single sample per patient is required.

Limitations of this study include the limited number of positive SARS-CoV-2 patients and modest number of other positive respiratory virus cases with the exception of rhinovirus. Further data on self-collection would be helpful to confirm these findings.

5. Conclusion

The world is facing unprecedented demands on health care services during the COVID-19 pandemic. Innovative ways to address this crisis are required and we believe that this study provides early evidence that self-collection of throat and nasal swabs for SARS-CoV-2 offers an acceptable and reliable alternative to health worker collected samples. This is achieved whilst preserving critically needed PPE supplies, optimizing the time to testing and reducing exposure of health care workers to potentially infected patients.

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Declaration of Competing Interest

All authors declare no competing interests.
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Appendix A. Supplementary data

Supplementary material related to this article can be found, in the online version, at doi:https://doi.org/10.1016/j.jcv.2020.104417.

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