Performance of an Antigen Rapid Test Compared to RT-PCR for the Detection of SARS-CoV-2

Bandar A. Suliman a, Ahmed Aljuhani b, Zuhur Ramadan Almohammadi c, Yasser A. Abdou d, Sultan Saud Alahmadi e, Amin Khattab f, Mohannad Almikhlafi g and Hossein M. Elbadawy g*

a Department of Medical Laboratory Technology, College of Applied Medical Sciences, Taibah University, Madinah, Kingdom of Saudi Arabia.
b Intensive Care Unit, Ohud Hospital, Madinah, Kingdom of Saudi Arabia.
c Nursing, Ohud Hospital, Madinah, Kingdom of Saudi Arabia.
d Internal Medicine, Ohud Hospital, Madinah, Kingdom of Saudi Arabia.
e Family Medicine, Ohud Hospital, Madinah, Kingdom of Saudi Arabia.
f Ohud Hospital, Madinah, Kingdom of Saudi Arabia.
g Department of Pharmacology and Toxicology, College of Pharmacy, Taibah University, Madinah, Kingdom of Saudi Arabia.

Authors’ contributions
This work was carried out in collaboration among all authors. All authors read and approved the final manuscript.

Article Information
DOI: 10.9734/JPRI/2023/v35i157379

Open Peer Review History:
This journal follows the Advanced Open Peer Review policy. Identity of the Reviewers, Editor(s) and additional Reviewers, peer review comments, different versions of the manuscript, comments of the editors, etc are available here:
https://www.sdiarticle5.com/review-history/101344

Received: 03/04/2023
Accepted: 05/06/2023
Published: 15/06/2023

Original Research Article

*Corresponding author: E-mail: hmbadawy@taibahu.edu.sa;

INTRODUCTION

In the month of March 2020, the World Health Organization declared Coronavirus disease 19 (COVID-19) is a global pandemic [1]. This pandemic outbreak was caused by the exposure to severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), which first appeared in December 2019 in Wuhan, Hubei Province, China then spread to the rest of the world [2]. However, recently the WHO declared that COVID-19 is no longer a pandemic or an emergency. Infected patients were suffering primarily from acute atypical respiratory symptoms including fever, dry cough, dyspnea, and hypoxia. In addition, other organ systems were also involved [3,4].

Worldwide, approximately 241 million patients were infected with the virus with a total death of around 6.8 million [5]. To control the spread of the infection and react quickly to new cases, faster and cheaper diagnostics are required. Currently, testing approaches fall into two main categories; either nucleic acid or serological [6-8]. Nucleic acid methods directly probe for the viral RNA of a swab taken from the patient throat or nasal cavity [9]. Real time polymerase chain reaction (RT-PCR) a was retained as the gold-standard for clinical diagnosis by the Centers for Disease Control and Prevention (CDC) [10,11].

However, running this method requires the use of special equipment, reagents, and well-trained personnel [12].

The Novel Coronavirus (SARS-CoV-2) Antigen Rapid Test (ART) Cassette (INVBIO) is an in vitro immunochromatographic membrane diagnostic test that detect coronavirus antigen using sensitive monoclonal antibodies [13]. Samples can be collected using throat swab, sputum sample, nasal swab, and nasal aspiration [14]. The ART can provide fast (takes less than 10 minutes to develop) and simple alternative to RT-PCR especially for routine screening. In this article, a side-by-side comparison was done to compare the sensitivity of ART to CDC-recommended RT-PCR protocol.

MATERIALS AND METHODS

The study was reviewed and approved by the general ethical committee for medical research in the ministry of health (MOH) in Madinah (approval number: IRB48-2021). Written consent was taken from all patients who agreed to
participate in the study. All subjects were assigned a study identification number and stayed anonymous and information which identifies patients was not used in this work. The study was performed between May and July 2021.

To study the performance of the rapid antigen test compared to RT-PCR, two nasopharyngeal swabs were taken from 103 COVID-19 hospitalized patients, in addition to 9 positive controls (COVID-19 confirmed cases) and 23 negative controls (healthy individuals). On the day when routine swabbing of patients is normally carried out for RT-PCR screening, an additional swab was taken from each patient to be used for the ART (INVBIO, Beijing, China). Results for the first swab were obtained from the hospital record, while the second swab was used on the rapid test onsite.

The kit contains individually sealed strips with two lanes, a bottom small slot for applying the sample and a top lane where the appearance of two bands indicated a positive result, while one band indicated a negative result. To use the kit, the test device was removed from the sterile foil pouch and placed on a clean and level surface. The nasopharyngeal swabs taken from a patient were inserted in the supplied disposable dropper containing 10 drops of the provided extraction buffer, mixed by squeezing and shaking well, before applying three drops onto the trip. Results were obtained between 2 and 5 minutes. Strips were maintained for further 10 minutes before disposal, to make sure no further changes will occur. Statistical analysis was carried out with a confidence interval (CI) of 95%. The 95% confidence interval was used for the positive and negative likelihood ratios.

3. RESULTS

3.1 Patients QPCR and ART Parallel Tests

Participants in this study were 126 patients, including 23 negative healthy control individuals, 9 positive controls with confirmed COVID-19 positivity by QPCR and 103 patients all from the COVID-19 isolation ward in the local COVID-19 reference hospital. Among those patients, 92.5% were in normal rooms, 4.7% in ICU and 2.8 on mechanical ventilators (MV) in ICU units. Two swabs were taken form each patient in hospital, out of which, one was sent for QPCR and the other for ART. 67.9% of QPCR results were positive while 27.4% of ART were positive (Fig. 1). Comparing QPCR results to ART results, when both tests were negative or positive, this was recorded as ‘agreement’. Agreement was found in 56% of cases showing a good correlation between the QPCR and ART results (Fig. 1), however, ART was less sensitive.

Fig. 1/A represents the distribution of gender (males/females) participating in the study. Fig. 1/B represents the distribution of patients according to ward and condition; inpatient are patients maintained at isolation rooms at normal room air, ICU represents patients with severe infections maintained in intensive care unit, while MV represents patients with critical conditions, maintained on mechanical ventilators.

Column representation of percentage positive results in all COVID-19 patients in QPCR and ART. The agreement percentage of patients with QPCR and ART identical results is shown in the third column. The percentage included two negative or two positive results for the same patient using QPCR and ART.

3.2 Analysis of Agreement between QPCR and ART

The first hypothesis to be tested was to measure weather QPCR test results is confirmatory of COVID-19 not clinical features. Clinical features included any signs or symptoms of COVID19, which were seen in all hospitalized patients in this study. The likelihood of agreement is shown in Fig. 3 using Fagan nomogram. The figures show the change in posterior probability after the NIRS VOT. The test was considered positive if the delta tissue oxygen index was < 15.2. The positive and negative log-likelihood ratios were 3.67 and 0.51. Panel A. The ‘prior’ was set at 0.8. The 95% confidence interval for the positive and negative log-likelihood ratios were (1.1-12) and (0.35-0.73). Panel B. The ‘prior’ was set at 0.001. The 95% confidence interval for the positive and negative log likelihood ratios were (0.01-1901) and (0.00-0.00). Results are shown in Table 1.

Graphical representation showing prior and posterior probability of COVID-19 test results and the likelihood ration. Fagan nomogram negative test (red) prior probability was set at 26%. The 95% confidence interval was (2-20) with a negative likelihood of 0.18. Posterior probability was 0.1. Positive test (blue) prior probability was set at 26%. The 95% confidence interval was (31-41) with a positive likelihood of 1.56. Posterior probability was 0.6.
Fig. 1. Participants by gender and percentage of patients

Fig. 2. Positivity of QPCR and ART
3.3 Investigating Clinical Features versus QPCR

To investigate whether clinical features will necessarily result in positive QPCR result, the second hypothesis was to determine if clinical features are confirmatory to COVID-19 rather than QPCR tests. Fig. 4 shows the results of the second hypothesis to investigate whether clinical features are confirmatory of COVID-19 infection and not QPCR. It shows a higher percentage within the 95% confidence level.

4. DISCUSSION

SARS-CoV-2 is a member of a large family known as coronavirus causing the COVID19 pandemic. The pandemic is now over as the virus has infected around 841 thousand in Saudi Arabia according to the ministry of health statistics. The gold standard for diagnosis of infected patients with the virus is PCR test, however, faster, and cheaper methods are urgently required to help control the spread of the infection. The Novel Coronavirus (SARS-CoV-2) Antigen Rapid Test Cassette (INVIBIO) is an in-vitro diagnostic test for qualitative detection of coronavirus antigens in nasal Swab and nasal aspirate samples, using the rapid immunochromatographic method [15]. The identification is based on coronavirus antigen specific monoclonal antibody. The assay will provide an easy and fast option especially for healthcare workers routine screening and is a promising tool for combatting the infection [16]. While PCR is currently the gold standard for the detection of the infection, new testing platforms were introduced based on the detection of antigens in nasopharyngeal swabs. Those tests are cheaper and can provide results within minutes. The PCR tests require certified laboratories, expensive equipment, and well-trained technicians to operate the instrument. In addition, false negative results have been reported when RT-PCR was used in detecting SARS-CoV-2 [17,18]. These limitations make RT-PCR inappropriate for use when rapid and simple diagnosis is necessary especially in the case of screening healthcare workers, travelers, and patients. Rapid and onsite detection methods can significantly improve the outbreak containment effort [19,20]. Therefore, there is an urgent need for a rapid, simple, sensitive, and accurate test to identify infected patients of SARS-CoV-2 to prevent virus transmission and to assure timely treatment of patients. The novel coronavirus (SARS-CoV-2) antigen rapid test cassette concept is to employ monoclonal antibodies with specificity for the novel coronavirus antigen [21]. The test is simple and takes 10 minutes to get results. Point-of-care diagnostic tests (POCTs) for detecting viral antigens in clinical samples would be very helpful for the diagnosis of COVID-19 either as mass-screening or first aid tests in the emergency room [22].
Fig. 4. Results of the second hypothesis to investigate whether clinical features

The SARS-CoV-2 ART kit used to detect SARS-CoV-2 antigens in nasopharyngeal swabs employs a rapid immunochromatographic method to identify the viral antigens using specific monoclonal antibodies. The relative sensitivity of the test was around 96.17% and the accuracy 98.79% as reported by the manufacturer. To get precise results using the CDC protocol, the test takes about three hours to complete and costs around $10 [7]. Samples taken from a swab of the nasopharyngeal cavity can harbor approximately 1 million viral particles [23]. In addition, serological tests quantify antibodies in the patient’s serum, which tend to be high during the first few days after infection [24].

5. CONCLUSION

The rapid test used in this research has many advantages over PCR due to its high sensitivity and accuracy. Besides, the cost of the test is much cheaper than PCR and it does not need training to collect or run the sample.

CONSENT

As per international standard or university standard, patient(s) written consent has been collected and preserved by the author(s).

ETHICAL APPROVAL

As per international standard or university standard written ethical approval has been collected and preserved by the author(s).

COMPETING INTERESTS

Authors have declared that no competing interests exist.

REFERENCES


Access on 20 October, 2021.


