

# Humoral immunological kinetics of severe acute respiratory syndrome coronavirus 2 infection and diagnostic performance of serological assays for coronavirus disease 2019: an analysis of global reports

Anthony Uchenna Emeribe pa, Idris Nasir Abdullahi pb,\*, Halima Ali Shuwac, Leonard Uzairued, Sanusi Musab, Abubakar Umar Ankab, Hafeez Aderinsayo Adekolae, Zakariyya Muhammad Bellof, Lawal Dahiru Rogog, Dorcas Aliyua, Shamsuddeen Harunab, Yahaya Usmanb, Habiba Yahaya Muhammadg, Abubakar Muhammad Gwarzoh, Justin Onyebuchi Nwofei, Hassan Musa Chiwari, Chukwudi Crescent Okwumek, Olawale Sunday Animasauni, Samuel Ayobami Fasogbonm, Lawal Olayemin, Christopher Ogara, Chinenye Helen Emeribeo, Peter Elisha Ghambap, Luqman O. Awoniyiq and Bolanle O. P. Musar

<sup>a</sup>Department of Medical Laboratory Science, Faculty of Allied Medical Sciences, University of Calabar, P.M.B 1115, Calabar, Cross River State, Nigeria; <sup>b</sup>Department of Medical Laboratory Science, Faculty of Allied Health Sciences, College of Medical Sciences, Ahmadu Bello University, Zaria, Nigeria; <sup>c</sup>University Health Services, College of Health and Medical Sciences, Federal University, Dutse, Nigeria; <sup>d</sup>Department of Microbiology, Federal University of Agriculture Abeokuta, Nigeria; <sup>e</sup>Department of Microbiology, Olabisi Onabanjo University, Ago-Iwoye, Nigeria; <sup>f</sup>Department of Medical Laboratory Science, Faculty of Allied Health Sciences, College of Medical Sciences, Ahmadu Bello University, Zaria, Nigeria; <sup>g</sup>Department of Medical Laboratory Science, Faculty of Allied Health Sciences, Bayero University, Kano Nigeria; <sup>h</sup>Department of Medical Microbiology and Parasitology, Federal University, Dutse, Nigeria; <sup>h</sup>Department of Medical Laboratory Science, University of Maiduguri, Nigeria; <sup>h</sup>Department of Medical Laboratory Sciences, University of Nigeria Teaching Hospital, Enugu, Nigeria; <sup>h</sup>Nigeria Field Epidemiology and Laboratory Training Programme, African Field Epidemiology Network, Abuja, Nigeria; <sup>m</sup>Public Health In-vitro Diagnostic Control Laboratory, Medical Laboratory Science Council of Nigeria, Lagos, Nigeria; <sup>m</sup>School of Medicine, Faculty of Health Sciences, National University of Samoa, Apia, Samoa; <sup>o</sup>Department of Family Medicine, University of Calabar Teaching Hospital, PMB 1278 Calabar, Cross River, Nigeria; <sup>p</sup>WHO National Polio Reference Laboratory, University of Maiduguri Teaching Hospital, Maiduguri, Nigeria; <sup>q</sup>Institute of Biomedicine, and MediCity Research Laboratories, University, Zaria, Nigeria

\*Corresponding author: Tel: +2348030522324; E-mail: eedris888@yahoo.com; inabdullahi@abu.edu.ng

Received 8 August 2020; revised 23 November 2020; editorial decision 20 January 2021; accepted 25 January 2021

As the coronavirus disease 2019 (COVID-19) pandemic continues to rise and second waves are reported in some countries, serological test kits and strips are being considered to scale up an adequate laboratory response. This study provides an update on the kinetics of humoral immune response to severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) infection and performance characteristics of serological protocols (lateral flow assay [LFA], chemiluminescence immunoassay [CLIA] and ELISA) used for evaluations of recent and past SARS-CoV-2 infection. A thorough and comprehensive review of suitable and eligible full-text articles was performed on PubMed, Scopus, Web of Science, Wordometer and medRxiv from 10 January to 16 July 2020. These articles were searched using the Medical Subject Headings terms 'COVID-19', 'Serological assay', 'Laboratory Diagnosis', 'Performance characteristics', 'POCT', 'LFA', 'CLIA', 'ELISA' and 'SARS-CoV-2'. Data from original research articles on SARS-CoV-2 antibody detection >second day postinfection were included in this study. In total, there were 7938 published articles on humoral immune response and laboratory diagnosis of COVID-19. Of these, 74 were included in this study. The detection, peak and decline period of blood anti-SARS-CoV-2 IgM, IgG and total antibodies for point-of-care testing (POCT), ELISA and CLIA vary widely. The most promising of these assays for POCT detected anti-SARS-CoV-2 at day 3 postinfection and peaked on the 15th day; ELISA products detected anti-SARS-CoV-2 IgM and IgG at days 2 and 6 then peaked on the eighth day; and the most promising CLIA product detected anti-SARS-CoV-2 at day 1 and peaked on the 30th day. The most promising LFA, ELISA and CLIA that had the best performance characteristics were those targeting total SARS-CoV-2 antibodies followed

<sup>©</sup> The Author(s) 2021. Published by Oxford University Press on behalf of Royal Society of Tropical Medicine and Hygiene. This is an Open Access article distributed under the terms of the Creative Commons Attribution-NonCommercial License (http://creativecommons.org/licenses/by-nc/4.0/), which permits non-commercial re-use, distribution, and reproduction in any medium, provided the original work is properly cited. For commercial re-use, please contact journals.permissions@oup.com

by those targeting anti-SARS-CoV-2 IgG then IgM. Essentially, the CLIA-based SARS-CoV-2 tests had the best performance characteristics, followed by ELISA then POCT. Given the varied performance characteristics of all the serological assays, there is a need to continuously improve their detection thresholds, as well as to monitor and re-evaluate their performances to assure their significance and applicability for COVID-19 clinical and epidemiological purposes.

**Keywords:** COVID-19 serology, diagnostics, laboratory tests, SARS-CoV-2.

## Introduction

The coronavirus disease 2019 (COVID-19) pandemic has caused an unprecedented global health emergency and economic uncertainty. As the incidence of COVID-19 continues to rise, many countries have sought to develop or procure serological test kits and strips with the plan of scaling up laboratory investigations into the COVID-19 pandemic.

Severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) is the etiological agent of COVID-19. It is one of the three highly pathogenic members of the family of *coronaviridae*.<sup>1</sup> Infection with SARS-CoV-2 has been associated with a range of hallmarks that progress from mild to severe clinical presentations before terminating in death in less than 10% of cases.<sup>2</sup>

The WHO has recommended RT-PCR as the gold standard protocol for screening individuals with typical symptoms who are suspected of having COVID-19. Although appropriate use of RT-PCR provides very accurate results, test reagents and consumables are mostly in short supply. Besides, this protocol is laborious, expensive to operate, requiring technical expertise and it has a long test turnaround time. Also, one of the major technical drawbacks in using RT-PCR is the significant number of cases of false-negative results, despite patients having clinical features and radiologic findings that are highly suspicious of SARS-COV-2 infection. The false-negative results could be due to wrong sampling, where SARS-CoV-2 might have been present in the lower respiratory tract rather than upper respiratory tract samples often collected for laboratory diagnosis. This poses a challenge in the proper evaluation of some SARS-CoV-2-infected people.<sup>3</sup>

It has been observed that the transmission dynamics of COVID-19 have made it an arduous task and challenge in the control of the pandemic, despite WHO-proposed measures having already been introduced.<sup>4</sup> Consequently, the COVID-19 pandemic has seriously challenged the operation of the entire healthcare system, including hospitals, laboratory diagnosis, the management of patients and every other aspect of human endeavor.<sup>5,6</sup>

In the quest to augment several lapses in the use of RT-PCR testing for COVID-19, serological assays that detect and /or measure antibodies (immunoglobulins) against SARS-CoV-2 have been developed and evaluated for performance by many institutions and private biotechnology firms. Global efforts to scale up the testing and diagnosis of COVID-19 has led to the commercial production of serological kits and devices. Some of these products have gained executive approval in some countries. For instance, the US Food and Drug Administration (FDA) gave expedient approval for some COVID-19 serological kits based on their accuracy and reliability.<sup>7</sup>

Instances have arisen where massive production and the use of finger-prick assays and in vitro testing have been encouraged in the UK and the USA to scale up COVID-19 surveillance through rapid testing and measurement of either antigens or antibodies to SARS-CoV-2. These rapid testing protocols adopted in these countries are point-of-care testing (POCT), which are designed as lateral flow devices (colloid gold-based immunochromatographic cassettes or test strips) with a diverse range of performance characteristics. These devices require a small sample volume (in microliters), are conducted within a short period (a few seconds to minutes), and are easier to perform as their use requires less technical expertise and equipment compared with protocols that detect nucleic acid.<sup>8</sup>

The transmission dynamics of the COVID-19 pandemic make it very challenging to control despite measures put in place in various countries of Africa and elsewhere outside the continent. Adequate laboratory diagnosis of COVID-19 plays a highly significant role in the control and prevention of the pandemic. However, some of the emerging challenges of testing for SARS-CoV-2 generally include sourcing personal protective equipment, low human capacity, scaling up testing, overwhelming contact tracing and inadequate hospital capacity to accommodate COVID-19 patients, resulting in increased morbidity and mortality. Hence, improved testing capacity, adequate provision of human and material resources, combined with innovative ways of scaling up contact tracing and improved testing capacity, are essential in the control of the COVID-19 pandemic.

This study sought to provide an update on the kinetics of humoral immune response to SARS-CoV-2 infection and performance characteristics of serological protocols (lateral flow assay [LFA], chemiluminescence immunoassay [CLIA] and ELISA) used for evaluations of recent and past SARS-CoV-2 infection. Data from original research articles on SARS-CoV-2 antibody detection ≥second day postinfection were included in this study. Furthermore, this study examined whether these tests could be possible solutions that can ameliorate the constraints of underdiagnosis in resource-limited settings.

This review is conducted under the following sections:

- 1. Virology and structural organization of SARS CoV-2 useful in molecular and serological diagnosis.
- 2. Humoral immune response to SARS CoV-2.
- 3. COVID-19 serological assays.
- 4. Challenges of SARS-CoV-2 serological testing.
- 5. Accuracy and applicability of COVID-19 serological assays.
- 6. Performance characteristics of COVID-19 serological assays.

#### **Article selection criteria**

## Search strategy

A thorough and comprehensive review of suitable and eligible full-text articles was performed on PubMed, Scopus, Web of Science, Wordometer and medRxiv from 10 January to 16 July 2020. These articles were searched using the MeSH terms 'COVID-19', 'Serological assay', 'Laboratory Diagnosis', 'Performance characteristics', 'POCT', 'CLIA', 'ELISA' and 'SARS-CoV-2'.

#### Article evaluation and data extraction

Eight authors independently evaluated and scrutinized titles and abstracts to prospective studies to check for potentially eligible articles and to acquire full texts from credible databases. Articles that were unavailable, incomplete or contained duplicate data were excluded. Furthermore, data from review articles were not considered for computing the antibody kinetics and performance characteristics of the serological assays.

Data were extracted from all eligible studies using the following criteria: (1) author, title, published date, the countries where studies were conducted, study design, sampling technique, participant inclusion criteria, number of participants enrolled and number of participants with known and available results; (2) main data, consisting of the results of serologic tests and RT-PCR for COVID-19 (sensitivity, specificity, positive predictive value [PPV] and negative predictive value [NPV]), number of days after the onset of symptoms, days of detection, peak and decline of antibodies; and (3) the test protocol used for serology and SARS-CoV-2 RNA detection.

#### Search outcome

In total, there were 7938 published articles on humoral immune response and laboratory diagnosis of COVID-19. Of these, 74 were included in this study based on selection criteria.

# Main findings

# Structural organization of SARS-COV-2 useful in serological diagnosis

SARS-CoV-2 is a single-stranded RNA virus with positive polarity.  $^{9,10}$ 

The SARS-CoV-2 genome consists of 14 open reading frames (ORFs) that code for 27 viral proteins, where the longest ORF coding for the 15 non-structural proteins plays an important role in viral propagation and immune evasion; the ORF codings for structural and accessory proteins are located on the 5' end and 3' end, respectively. 11 The first ORF code encompasses two-thirds of the viral genome and translates the polyproteins pp1a and pp1ab, which are implicated in the encoding of the 16 non-structural proteins. However, the remaining ORFs code for the viral structural and accessory proteins. The structural protein nucleocapsid (N) proteins, spike (S) glycoprotein, matrix (M) protein and small envelope (E) complete the remaining one third of the viral genome. 12 These proteins and RNA-dependent RNA polymerase have been substantially harnessed for primers and antigens in the molecular and serological assays used for COVID-19, respectively. 13

## Humoral immune response to SARS-COV-2 infection

There is ongoing research into better understanding the viral genome assembly, replication and mutation of SARS-CoV-2. These viral attributes drastically influence the diagnostic performance of both molecular and serological assays as well as the transmissibility of SARS-CoV-2 and its immune responses.<sup>14</sup>

Prior to SARS-CoV-2 infection, an unexposed individual was expected to have a negative test for either anti-SARS-CoV-2 IgM or IgG (Figure 1). However, following exposure to the infection, SARS-CoV-2 now induces a humoral immune response, which commences with the development of IgM, indicating an acute or ongoing infection from the third day of the first week of infection, as reflected by a positive outcome in either the IgM or IgM/IgG serological test.<sup>15</sup> The level of IgM in an individual with the activated humoral immune response against SARS-CoV-2 continues to rise until it peaks during the third week following infection.<sup>15</sup> By the end of the third week, IgM levels decrease with a concomitant elevation in the level of IgG from the third to the seventh week post symptom onset (PSO), which is revealed by a positive outcome in either the IgG or IgM/IgG serological test (Figure 2).<sup>15</sup>

The median period for the development of all the classes of immunoglobulins following the activation of humoral immune response is 13 d.<sup>16</sup> Individually, IgM, IgG and total immunoglobulins have an average duration of 11, 12 and 14 d, respectively. 16 These immunoalobulins can be measured and monitored by a diverse range of antibody-based serological testing techniques, which include rapid diagnostic assay (e.g. lateral flow immunoassay [LFIA] with colloidal gold], CLIA, ELISA and neutralization assay with various diagnostic performance ratings (e.g. sensitivity, specificity, accuracy, PPV and NPV), sampling methods (e.g. finger prick, venipuncture), turnaround time and setting. 16 Previous studies have demonstrated the diagnostic roles of these antibody-based serological testing techniques based on their performance. Zhao et al.<sup>17</sup> demonstrated that within the first 7 d PSO of COVID-19 infection, the sensitivity of total antibody, IgM and IgG were 38.3%, 28.7% and 19.1%, respectively, which was lower compared with the RNA-based test of 66.7% sensitivity. As the duration of PSO increased, the sensitivity of the RNA-based test decreased by 21.2%, while those of the total antibody, IgM and IgG increased by 61.7%, 65.6% and 60.78%, respectively, within the 15th to 39th day PSO. When the RNA- and antibodybased tests were combined, sensitivity significantly improved to 78.7%, 97.0% and 100% within 1-7, 8-14 and 15-39 d PSO, respectively. The implication of this study indicates the unreliability and unsuitability of serology within the window period of infection, but also reveals an impressive sensitivity for total antibodybased assay in detection of SARS-CoV-2 as the PSO period progresses.

The study further revealed that the percentage of patients with undetectable RNA but with detectable immunoglobulin increased from 28.7% within the first 3 d to 100% within 15–39 d of PSO. This is where the total Ab (which is better than testing IgM and IgG individually) comes in, to rule out people with undetectable RNA.<sup>17</sup> The same study recommended the combination of both RNA- and antibody-based tests to scale up the sensitivity of RNA during the course of the infection. This combined approach was observed to attain timely diagnosis of SARS-CoV-2 infection, prevent multiple sampling several days

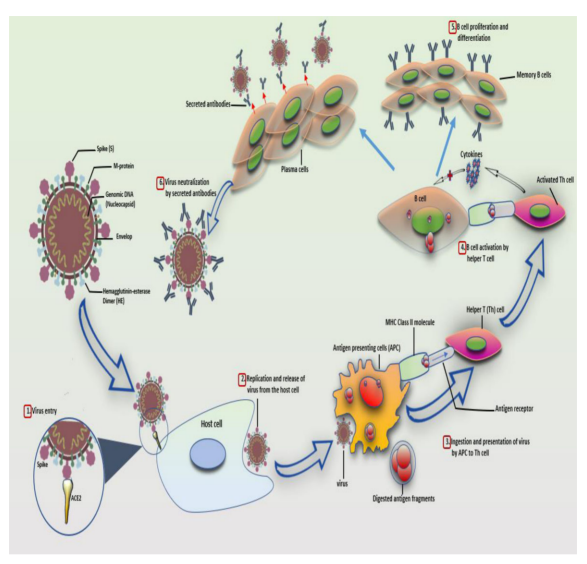


Figure 1. Kinetics of antibody response in SARS-CoV-2 infection. The entry of the SARS-CoV-2 virus into the host cell through interaction and binding between the host's angiotensin-converting enzyme 2 (ACE2) proteins (receptor) and the viral spike (S) protein (ligand) (1). Following replication and release from the host cells (2), antigen-presenting cells (APCs) like macrophages and dendritic cells engulf some of the viruses, digest and present the digested antigen fragments on their class II MHC molecules to the helper T (Th) cells (3). Th cells, in turn, activate B cells (4), activated B cells proliferate and differentiate into memory B cells or plasma cells with high affinity to the SARS-CoV-2 antigens (5). Plasma cells release SARS-CoV-2-specific antibodies (IgM, IgG or IgA) that bind and neutralize the viruses, thus preventing the viral entry into the host cell (6).

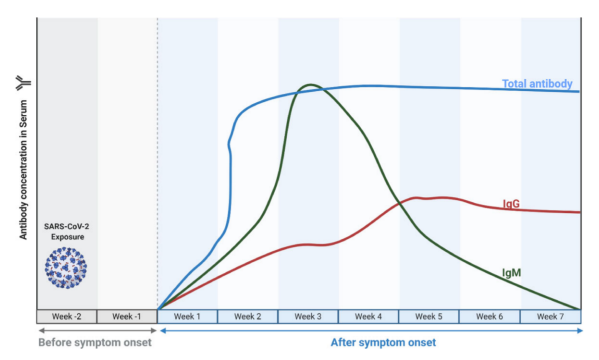
for infection status confirmation, and enhance the ability to prioritize relevant treatments and isolation management.  $^{18-20}$ 

The changes of the antibody response against SARS-CoV-2 are presently under study, as antibodies may be regarded as potent diagnostic tools to complement RT-PCR-based findings. The SARS-CoV-triggered humoral S- and N-specific IgM response reached a climax within 4 wk and was no more detectable at 3 mo PSO; the switch to IgG often occurred about day 14 and IgG were demonstrated up to 36 mo.<sup>21,22</sup>

In another study, the authors demonstrated that in 34 SARS-CoV-2 laboratories established, the cases studied were positive for IgM and IgG at week 3 PSO.<sup>15</sup> Therefore, in the majority of those patients, the acute phase of infection persisted for

>30 d. In an inverse relation, as IgM levels decrease, IgG levels rise gradually from the third to the seventh week, signifying the activation of the humoral immune response against the virus.<sup>15</sup> Thus, the humoral response activated by SARS-CoV-2 may be similar to that elicited by SARS-CoV.<sup>15,16</sup>

In an immunodynamics study reported by Zhao et al.,  $^{17}$  it was observed that the antibody profile in COVID-19 patients showed that seroconversion sequentially appeared for total antibodies, IgM and IgG with a median time of 11, 12 and 14 d, respectively. Full concentrations of SARS-CoV-2 antibodies were detected by double recombinant antigen sandwich immunoassay, which utilized the receptor-binding domain (RBD) of S1 protein and the horse raddish peroxidase-conjugated antigen; IgM  $\mu$ -chain



**Figure 2.** Timeline of IgM, IgG and total antibody kinetics during SARS-CoV-2 infection. The level of IgM in an individual with the activated humoral immune response against SARS-CoV-2 continues to rise until it peaks at the third week following infection. By the end of the third week, IgM levels decrease with a concomitant elevation in the level of IgG from the third to the seventh week postonset symptom (POS). For the total antibody, it peaks at the middle of the second week and reaches a plateau in the middle of the third week. The blood concentration persists for several weeks and months postinfection (image made with Biorender.com).

capture immunoassay was used for anti-SARS-CoV-2 IgM detection. On the other hand, an indirect ELISA kit based on recombinant NP antigen was used for anti-SARS-CoV-2 IgG detection.<sup>17</sup> The seroconversion rates recorded were 93.1%, 82.7% and 64.7% for total antibodies, IgM and IgG, respectively, and no significant difference was observed between severely and mildly affected COVID-19 patients.

The sensitivity of serum anti-SARS-CoV-2 detection was lower than the RT-PCR RNA assay within 7 d from the onset of illness (38.3% vs 66.7% for serological vs RT-PCR). However, the sensitivity increased steadily from the eighth to the 39th day PSO and overtook that of the RT-PCR test. More significantly, detectable and measurable levels of total anti-SARS-CoV-2 in the sera were found in COVID-19 patients with undetectable SARS-CoV-2 RNA in their respiratory tract samples. These results highlighted the importance of combining molecular and serological tests for the correct diagnosis of COVID-19 patients at different stages of the disease. In agreement with these reports, Jin et al. recorded the specificity of serum anti-SARS-CoV IgM and IgG as 90% compared with that of the RT-PCR test. Market services and services as the services of the RT-PCR test.

In a study by Guo et al., which was carried out on two cohorts of SARS-CoV-2-infected patients, the early antibody response to NP protein was evaluated. Of 208 patients, 90.4% and 93.3% harbored plasma IgM and IgA, respectively. Also, 77.9% of plasma samples were IgG positive, and the median time for both IgM and IgA detection was on day 5 PSO (IQR 3–6) and day 14 PSO (IQR 10–18) for IgG.<sup>24</sup> The authors observed that swift and unanticipated IgA seroconversion might be an upshot of the

cytokine storm promoting the germline transcription of  $\alpha$  and  $\mu$  genes of the heavy chain constant.

Furthermore, it has been reported and established that T-cell-independent antibody responses can cause excitation of a specialized B cell subset to produce both IgA and IgM throughout the infection of some pathogens. Although T-cell-independent antibody response against viruses is still controversial, some viruses can act in vivo as T-cell-independent antigens and therefore cause eliciting protective isotype-switched antibodies in the non-appearance of conventional T-cell help. Inactivated virus or virus-like particles can also elicit IgM response, but factors induced when an active virus infection is ongoing seem very important and are required before there can be induction of the isotype switch and then IgG or IgA responses. <sup>26</sup>

In another study of 214 COVID-19 patients, 68.2% and 70.1% were positive for rN-specific IgM and IgG, respectively; and 77.1% and 74.3% were positive for rS-specific IgM and IgG, respectively.<sup>27</sup> These findings indicated that the detection of rS-specific IgM was more sensitive compared with that of rN-specific IgM, which may be because of the lower immunogenicity of the N protein compared with that of the S protein. A bioinformatics study reported a lower number of B cell epitopes in the NP protein of SARS-CoV-2 than in the S protein, especially as the positive rates of IgM and IgG were low during the early stages of the disease (0–10 days post-disease onset (DPO)). On the other hand, IgM and/or IgG specific for rN and rS reached a climax at 11–15 DPO.<sup>27</sup>

The sensitivity of the tests and the epitope on which the test is based are significant factors for the well-organized detection of specific SARS-CoV-2 antibodies and timing of the humoral

response. Consequently, several tests are rapidly being developed in many laboratories. For example, Li et al. developed a point-ofcare LFIA test based on the RBD antigen of the SARS-CoV-2 S1 protein that can help in the concomitant detection of IgM and IgG in human blood within 15 min, with higher sensitivity than the individual IgG and IgM tests; however, the detection limit of the test was not determined. 19 Also, Amanat et al. developed sensitive and specific ELISA assays based on the recombinant fulllength S protein and RBD epitope, permitting the screening and detection of seroconversion upon SARS-CoV-2 infection 3 d PSO.<sup>28</sup> Of note, no cross-reactivity from other human coronaviruses was noted, in agreement with another study highlighting that S1 is a specific antigen for SARS-CoV-2 diagnosis, as cross-reactive antibodies against the S protein of Middle East respiratory syndrome-related coronavirus (MERS-CoV) were not detected in a COVID-19 patient.<sup>29</sup> Additionally, strong IgA and IgM responses were discovered and the IaG3 response was stronger than that of IgG1.<sup>28</sup> The sensitivity of the test may create challenges for the early detection of IgM. Several patients were more positive for IgG than IgM during the time of hospital stay and 5 d later; likewise, they had an earlier IgG than IgM seroconversion.<sup>30</sup>

Furthermore, SARS-CoV-2-specific antibodies were detected in the sera of six infants born to mothers with COVID-19. Five of the six infants and their mothers had elevated levels of IgG and two of them also had elevated levels of anti-SARS-CoV-2 IgM. Three of the six infants who had elevated levels of IgG also had normal levels of IgM. However, two of their mothers displayed elevated levels of IgM. How the newborns that developed IgM require additional investigation. Undeniably, due to its large magnitude, IgM is not typically transferred through the placenta; however, it is affected by some pathology that compromises its configuration. The newborn might be in contact with the virus if the latter crosses the placenta, although no virus was detected from RT-PCR analysis.<sup>31</sup>

Currently, several studies are investigating the connection between antigen-specific antibodies and the clinical characteristics of COVID-19 patients, but interestingly, among people with comorbidities, lesser anti-RBD IgG, but not anti-NP IgM or IgG, have been reported, although the difference was not significant when compared with people without comorbidities.

#### COVID-19 serological assays

The recent pandemic outbreak of the SARS-CoV-2 virus and its rapid spread poses an urgent need for both diagnostic and therapeutic interventions to manage the infection and the outcome of the disease. The diminishment or absence of IgG and persistence of IgM are considered biomarkers for recent infection. As the epidemic progresses more individuals could get infected. The measurement of these antibodies is a good differential that helps to distinguish between recent and older infections. The detection of IgM (from days 1 to 7) in the absence of IgG represent an acute/recent infection, whereas the simultaneous detection of IgM and IgG could represent acute reinfection. Route other hand, the detection of IgG in the absence of IgM denotes a past infection.

The increasing number of confirmed COVID-19 cases has resulted in an unprecedented rise in demand for antibody-based tests from researchers and healthcare policymakers. Recently, a

list of >200 serological products was released by the Foundation for Innovative New Diagnostics (FIND); these products, which are predominately from China, are currently either available for use or are in industrial development and evaluation. However, only 12 have received emergency use authorization from the FDA. Serological products from a host of other countries, including South Korea, Germany, the USA and the UK, were also present on the FIND list.

Some commercially available serology-based tests have been considered to be inadequate for COVID-19 diagnosis if used alone, due to their low degrees of sensitivity and specificity. For instance, anti-SARS-CoV-2 IgG takes a relatively long period (not yet reported) for quantification. <sup>18</sup> More details regarding the limitations of COVD-19 serological assay follow later in this article.

Cases of poor performance characteristics of some serological kits/devices underscore the need for re-evaluation and validation before being made available to end-users. This is to prevent clinicians and healthcare professionals from using these serological kits/strips off the shelf for clinical purposes. Furthermore, despite kits' satisfactory diagnostic performance, it is important to include internal quality control and external quality assurance measures in all tests run on human samples to ensure accuracy, precision and reproducibility of test results.

## Challenges of SARS-CoV-2 serological testing

Serological tests rely on the detection of specific anti-viral antibodies (IgM, IgA, IgG or total antibody) in patient sera, plasma or whole blood.<sup>32</sup> Determining the optimal antigenic epitopes to maximize sensitivity, but minimize cross-reactivity, particularly against other human coronaviruses, has meant that the development of high-quality serological testing has been slower than molecular-based diagnostics.<sup>17,32</sup> Initial candidate epitopes have largely focused on the immunogenic viral structural proteins which include nucleocapsid (N) and spike (S) protein, particularly the S1 subunit and the RBD.<sup>32</sup> To date, a range of serological tests for COVID-19 have been developed, each with particular test characteristics. Broadly, these serological tests can be divided into tests that (1) can be performed at the point of care; (2) can be performed in routine diagnostic laboratories; and (3) can only be performed in specialized reference laboratories.

Initial studies have reported that most patients with COVID-19 seroconvert by day 10–14 (approximately 80%), with almost 100% seroconversion by day 20.<sup>6,7</sup> However, comparisons across published studies are challenging due to (1) different antigens used in assays; (2) differences in the complexity of patient populations; and (3) variations in the RT-PCR assays used as the gold standard for determining the sensitivity of serological assays. Further, it is not clear whether the type and number of antibodies correlate with the severity of COVID-19, or more importantly, with immune protection from reinfection.

At present, the most widely available (and most publicized) serological tests are POCT, which involves the detection of anti-SARS-CoV-2 antibodies through binding to immobilized antigen, generally bound to colloidal gold on a test strip. The relatively cheap and simple nature of lateral flow assays means that production is suited to scaling up for increased testing capacity. However, there are limited published data on the performance characteristics of serological POCT, and high-quality data are urgently

Se	96.55 73.44 62.22 97.92 (47/48) a. No additional validation in participants with (28/29) (47/64) (28/45) mild symptoms; this is required to rule out any		) (49/64) (28/43)	73.44 60.47 94.0 (47/50) c.	(47/164) (26/43) 84.09 86.0 86.05 (37/43) d.	(43/49) (37/44) (43/50) specificity for orient gene RDI	83.67 84.09 85.42 82.22 (37/45) (41/49) (37/44) (41/48)	81.82	100.0	(9/2) (6/6) (			(9/6) (1/4/14)	88.89 100.0 100.0 50.0(9/18) (72/81) (9/9) (72/72)	90.63 96.7 72.05 a.	(352/397) (116/128) (352/364) (116/161) SARS-CoV-2	70.53 98.44 99.29 53.09 b. Cross-reactivity studies with other (280/397) (126/128) (280/282) (126/243) coronaviruses and flu viruses were not	performed 82.62 01.71 06.76 62.00 c. Tha lawal of changes in immunacelabilities was
Sample Type size	al 93	IgG 93		IgM 93	Total 93		IgG 93	IgM 93	Total 90			lgG 90		IgM 90	Total 525		IgG 525	I CM
Product name/source	GeurtsvanKessel Rapid SARS -CoV-2 antibody Total et al. <sup>b 43</sup> (IgM/IgG) test from InTec (Test of	lot \$2020021505) I		I		rapid test (GICA) from Cellex Inc. (test lot 0200311WI5513C-3)	I		COVID-19 JaG/JaM rapid test		blood/serum/plasma) from orient aene/Healaen (test lot 2003260)			П		antibodies, rabbit IgG and goat anti-rabbit IgG antibodies were obtained from Sigma-Aldrich		-

Citation	Product name/source	Type	Sample	Sensitivity (%)	Specificity (%)	(%) Add	(%) NBN	Limitation of study
Cassaniti	VivaDiagTM COVID-19 IgM/IgG rapid	Total	20	18.42	91.67	87.5 (7/8)	26.19	a. Small sample size
פו מו.	1621	IgG	20	13.16 (5/38)	100 (12/12)	100 (5/5)	26.67	b. Poor sensitivity
		IgM	20	15.79 (6/38)	91.67 (11/12)	85.71 (6/7)	25.58 (11/43)	c. High false-negative value, which can lead to misdiagnosis
orte et al.ª 45	Porte et al.a 45 Fluorescence immunochromatographic SARS-CoV-2 antigen test (Bioeasy Biotechnology Co., Shenzhen, China)	Total	127	93.9 (77/82)	100 (45/45)	100 (77/77)	90.0 (45/50)	a. Use of samples not specifically permitted by the manufacturer of the kit
		IgG IgM	127	N A o	NA° NA°	NA <sup>a</sup>	NA <sup>a</sup> NA <sup>a</sup>	b. Retrospective use of clinical data
Pan et al.ª 46	Colloidal gold-based immunochromatographic strip (Zhuhai Livzon Diaanositic Inc.)	Total	108	68.6 (59/86)	63.64 (14/22)	88.06 (59/67)	34.15 (14/41)	a. Intensity of color bands formed do not correlate with the abundance of immunoglobulin
		IgG	108	54.65	59.09	83.93	25.0	b. Very low probability of having negative outcomes without the infertion
		IgM	108	55.81 (48/86)	36.36 (8/22)	77.42 (48/62)	17.39 (8/46)	
Lassauniere et al.ª 35	2019-nCOV IgG/IgM rapid test (Dynamiker Biotechnology, Tianjin, China Caf # DNK-1419-1)	Total	62	90 (27/30)	100 (32/32)	100 (27/27)	89 (32/35)	a. Small sample size
		IgG	62	NΑ <sup>b</sup>	NAb	NAb	NAb	b. All kits were not tested uniformly with the same number of control sera
		IgM	62	NAb	NA <sup>b</sup>	NAb	NAb	c. Acro Biotech and Alltest Biotech had comparatively poor test performances, which led to the suspension of further testina
	OnSiteTM COVID-19 IgG/IgM rapid test (CTK Biotech, Poway, CA, USA; cat. # R0180C)	Total	62	90 (27/30)	100 (32/32)	100 (27/27)	89 (32/35)	c. Acro Biotech test had a cross-reaction with a control serum of a patient infected with human coronavirus HKU1

Limitation of study				oatients with $\leq 14  \mathrm{d}  \mathrm{F}$						nts with symptoms of infection			Inadequate number of cases, which could not reveal the statistical difference in the performance characteristics for the various POCT	ion for cross-reactivi	ution of COVID-19 vith patients having y infiltration					
Limitatio				a Poor performance in patients with $\leq \! 14$ d POS				Small sample size	מ. טוומו טמוווטופ טוצפ	b. Not reliable for patients with symptoms within the early days of infection		a. Single-center study	<ul> <li>Inadequate number of cases, which could not reveal the statistical difference in the performance characteristics for the variou POCT</li> </ul>	c. Laboratory investigation for cross-reactivity	sucuses were indeequate d. Possible misclassification of COVID-19 pneumonia patient with patients having subclinical pulmonary infiltration					
NPV (%)	93.23 (124/133)	98.4	(123/125)	70.59	(48/68)	(48/74)	(50/75)	76 7.7	(13/17)	NAb	NAb	100 (98/98)	100 (98/98)	97.96	(289/306)	94.44	(289/306) 93.46	(286/306)	82.69 (234/283)	N N o
PPV (%)	100 (20/20)	96.43	(27/28)	98.48	(130/132)	(124/126)	100 (125/125)	QF.O	(19/20)	NAb	NAb	91.67 (22/24)	91.67 (22/24)	83.33	(20/24) 92.95 (145/156)	92.95	(145/156) 91.03	(142/156)	99.68 (312/313)	A N N
Specificity (%)	100 (124/124)	99.19	(123/124)	96.0	(48/50)	(48/50)	100 (50/50)	92 86	(13/14)	NAb	NAb	98 (98/100)	98 (98/100)	96	(387/100) 96.33 (289/300)	96.33	(289/300) 95.33	(286/300)	82.11 (234/285)	NA A
Sensitivity (%)	68.97 (20/29)	93.1	(27/29)	82.67	(124/150)	(124/150)	83.33 (125/150)	82.61	(19/23)	NA <sup>b</sup>	NAb	100 (22/22)	100 (22/22)	90.91	(20/22) 89.51 (145/162)	89.51	(145/162) 87.65	(142/162)	86.43 (312/361)	N Ab
Sample size	153	153		200	200	000	200	37	ñ	37	37	122	122	122	462	462	462	Ĺ	980	596
Туре	IgG	IgM		Total	InG	ָ ס	IgM	To+oT		IgG	IgM	Total	IgG	IgM	Total	IgG	IgM	- - -	וסנמו	IgG IgM
Product name/source				COVID-19 IgG/IgM rapid test	cassettes (OrientGene)			Shoditan Mall 201 Vola-2010	rapid test kit (Beijing Diagreat Biotechnologies Co Ltd)			ALLTEST 2019-nCoV IgG/IgM rapid test (Hangzhou ALLTEST Biotech Co., Ltd. [China])			Dynamiker 2019-nCoV IgG/IgM rapid test (Dynamiker Biotechnology [Tianjin] Co, Ltd. [Chinal)				Wonaro SARS-CoV-2 anuboay test (Guangzhou Wondfo Biotech Co. Ltd [China])	
Citation				Pallet et al.ª	<del>,</del>			Spiciol of	al.a 48			Wu et al. <sup>a</sup> <sup>49</sup>								

Citation	Product name/source	Type	Sample	Sensitivity (%)	Specificity (%)	PPV (%)	NPV (%)	Limitation of study
Green et al. <sup>a</sup> 50	COVID-19 IgM-IgG Rapid Test (BioMedomics, BD, USA)	Total	525	88.66	90.63	96.7 (352/364)	72.05 (116/161)	a. No detail on the diagnostic performance of both IgG and IgM for most POC diagnostic devices that were evaluated.
		IgG	525	NAb	NA <sup>b</sup>	NAb	NA <sup>b</sup>	
		IgM	525	NAb	NΑ <sub>ρ</sub>	NΑ <sub>ρ</sub>	NΑ <sub>ρ</sub>	
	Xpert SARS- CoV-2 (Cepheid	Total	65	100	100	100	100	
	[USA/worldwide distribution])	(-1	Ļ	(30/30)	(35/35)	(30/30)	(35/35)	
		Dgr Mol	65 65	A N	NA P	N A N	A N	
	VitaPCR COVID-19 assay (Credo	Total	180	100		100	100	
	[Singapore])			(120/120)		(120/120)	(09/09)	
		IgG	180	NAb	NA <sup>p</sup>	NAb	NAb	
		IgM	180	NAb	NAb	NAb	NAb	
	Accula SARS- CoV-2 (Mesa	Total	80	100	100	100	100	
	Biotech [USA])			(20/20)	(30/30)	(20/20)	(30/30)	
		IgG	80	NA♭	NAb	NAb	NAb	
		IgM	80	NAb	NAb	NAb	NAb	
	ID NOW COVID-19 (Abbott	Total	09	100	100	100	100	
	Diagnostics [worldwide])			(30/30)	(30/30)	(30/30)	(30/30)	
		IgG	09	NAb	NA <sup>b</sup>	NAb	NAb	
		IgM	30	NAb	NAb	NAb	NAb	
	GT-100 SARS-CoV-2 IgG/IgM kit	Total	70	100	(49/20)	6.06	100	
	(Goldsite Diagnostics Inc.			(20/20)		(20/22)	(64/64)	
	[China])	(	1	0		0	7	
		1gc	0	100	98 (49/50)	90.9	100	
		McI	02	(20/20)	06 (/,8/50)	(20/22)	(45/45)	
		I gla	2	(07//1) (0	(00/04) 06	(17/19)	(48/51)	
MIcochova	SAMBA II SARS-CoV-2 point of	Total	45	79.17	100	100	80.77	a. Small sample size
et al.ª 51	care testing			(19/24)	(21/21)	(19/19)	(21/26)	-
	n	IqG	45	50.0	100	100	63.62	b. Recommendation of combined rapid testing
		)		(12/24)	(21/21)	(12/12)	(21/33)	protocol with PCR in order to ensure:
		IgM	45	87.5	100	100	87.5	i. Expansive testing in areas where diagnostic centers
		)		(21/24)	(21/21)	(21/21)	(21/24)	are sparse, and transmission is rapid
	COVIDIX 2019 SARS-CoV-2	Total	45	95.83	85.71	88.46	94.74	ii. That repeated sampling is avoided, which can
	IgG/IgM test (COVIDIX			(23/24)	(18/21)	(23/26)	(18/19)	generate aerosols and encourage transmission
	Healthcare, Cambridge, UK)	רַט	57	100	80 Q5	85 71	100	iii That nationts are safely and anickly recruited for
		ילע	)	)	)		)	וווי ווומר מתוכוונט מול נמוכים מיום ממוכיה וליים

Citation		Van Elslande et al.ª <sup>52</sup>																														
Product name/source		Clungene COVID-19 IgG/IgM rapid test				OrientGene COVID-19 IqG/IqM rapid test	- n			VivaDiag COVID-19 IgG/IgM	rapid test			StrongStrep COVID-19	IgG/IgM rapid test				Dynammiker COVID-19	IgG/IgM rapid test				Multi-G COVID-19 IgG/IgM	lapia test			Prima COVID-19 InG/IaM	rapid test			
Туре	IgM	Total	IgG	MgI	- 	Total	IgG	Mol		Total	707	D fi	IgM	Total		IgG	Mol	<u> </u>	Total		IgG	IgM		Total	IaG	) )	IgM	Total		IgG	MgI	
Sample size	45	256	256	256	L	256	256	256	000	256	256	7.70	256	256		256	256	007	256		256	256		256	256	0	256	756	2	256	256	
Sensitivity (%)	95.83 (23/24)	35.95 (55/153)	62.09	39.22	(60/153)	64.05 (98/153)	67.97	(104/153)	(111/153)	62.75	(96/153) 62.75	(96/153)	65.36	(100/153)	(46/153)	64.71	(99/153) 32 03	(49/153)	61.44	(94/153)	61.44	69.28	(106/153)	37.25	(57/155)	(99/153)	43.79	(67/153)	(74/153)	71.24	(109/153) 56.21	(86/153)
Specificity (%)	90.48 (19/21)	99.03 (102/103)	98.06	(101/103) 91.26	(94/103)	97.09 (100/103)	93.2	(96/103)	(98/103)	100	(103/103)	(102/103)	100	(103/103)	(103/103)	99.03	(102/103)	(102/103)	99.03	(102/103)	99.03	95.15	(98/103)	100	(103/103)	(100/103)	91.26	(94/103) 98.06	(101/103)	90.29	(93/103) 93.2	(96/103)
(%) Add	92.0 (23/25)	98.21 (55/56)	97.94	(95/97)	(69/09)	97.03 (98/101)	93.69	(104/111)	(111/116)	100	(96/96)	(26/96)	100	(100/100)	(94/94)	99.0	(99/100)	(49/50)	98.94	(96/46)	98.94	95.5	(106/111)	100	(75775) 97.06	(99/102)	88.16	(67/76)	(74/76)	91.6	(109/119) 68.25	(00/100)
(%) NAN	95.0 (19/20)	51.0 (102/200)	63.52	(101/139)	(94/187)	64.52 (100/155)	66.21	(96/145)	(98/140)	64.38	(103/160)	(102/159)	66.03	(103/156)	(103/210)	65.38	(102/156)	(102/206)	63.35	(102/161)	63.35	(102/101)	(98/145)	51.76	(103/199)	(100/154)	52.22	(94/180) 56.11	(101/180)	67.88	(95/157) 71.67	(0,0,0)
Limitation of study		<ul> <li>a. Control samples were limited in number from patients with frequent respiratory disorders</li> </ul>	b. Antibody response studies in asymptomatic or mild	individuals were not performed c. Participants were not tested daily to accurately	determine the true period of seroconversion																											

idble 1. continued	luea		9					
Citation	Product name/source	Туре	sample size	Sensitivity (%)	Specificity (%)	PPV (%)	(%) NPV	Limitation of study
Jääskeläinen et al. <sup>b 53</sup>	2019-nCoV IgG/IgM rapid test cassette (Acro Biotech, California, USA)	Total	123	56.1 (23/41)	74.39 (61/82)	52.27 (23/44)	77.22 (61/79)	a. Low PPVs for Acro Biotech IgG/IgM rapid test due to low SARS-CoV-2 seroprevalence
		IgG	123	56.1	74.39	52.27	77.22	
		IgM	123	(23/41) 46.34	(61/82) 69.51	(23/44) 43.18	(61/79) 72.15	
		1		(19/41)	(57/82)	(19/44)	(57/79)	
	SARS-CoV-2 IgG/IgM rapid test (Xiamen Biotime, Fujian, China)	Total	112	81.25 (26/32)	97.5 (78/80)	92.86 (26/28)	92.86 (78/84)	
		IgG	112	71.88	97.5	92 (23/25)	89.66	
		IgM	112	81.25 (26/32)	88.75 (71/80)	81.25 (26/35)	92.21 (71/77)	
Kohmer et al. <sup>c 54</sup>	FasStep (COVID-19 IgG/IgM) rapid test cassettes (COV-W32M, Assure Tech (Hangzhou) Co., Ltd, China)	Total	29	93.75 (15/16)	100.0 (13/13)	100.0 (15/15)	92.86 (13/14)	a. Small sample size
		IgG	29	93.75	100.0	100.0	92.86	
		MoI	79	(15/16)	(13/13)	(15/15)	(13/14)	
		<u>.</u>	7	(10/16)	(13/13)	(10/10)	(13/19)	
Montesinos et al.º 5	2019-n-CoV IgG/IgM rapid test cassette (LabOn Time) (LabOn Time, Bio Marketing Diagnostics, or Akiva, Israel)	Total	200	71.88 (92/128)	100.0 (72/72)	100.0 (92/92)	66.67	a. The reference standard used for the comparative study of the serological kits
		IgG	200	67.19 (86/128)	100.0 (72/72)	100.0 (86/86)	63.16 (72/114)	<ul> <li>b. Poor diagnostic performance based on the sensitivity of IgM and IgG for LabOn and Quickzen, respectively</li> </ul>
		IgM	200	48.44	100.0	100.0	52.17 (72/138)	
	Novel coronavirus (2019-n-CoV) antibody 1gG/1gM assay (colloidal gold) (Avioq, Biotech, Shandona, China)	Total	200	(88/128)	95.83 (69/72)	96.7 (88/91)	(69/109)	
	ì	IgG	200	68.75	95.83	96.7	63.3	
		Σ	000	(88/128)	(69/72)	(88/91)	(69/109)	
		1	000	(88/128)	(69/72)	(88/91)	(69/109)	
	QuickZen COVID-19 IgM/IgG kit (QuickZen) (ZenTech, Angleur, Belgium)	Total	200	71.09 (91/128)	100.0 (72/72)	100.0 (91/91)	66.06 (72/109)	
		IgG	200	49.22	100.0	100.0	52.55	
		IgM	200	(03/120) 68.75 (88/128)	100.0	(85/85) 100.0 (88/88)	(72/137) 64.29 (72/112)	
				(0.21	(7,17,1)	(00,00)	(77177)	

Limitation of study	a. Presence of false-positives due to cross-reactivity of non-specific immunoglobulins, which reflects past exposure to other seasonal viral infections of the coronavirus group	b. Small sample size, which did not encourage strong confidence intervals around the diagnostic performance of the LFIA kits.	c. The kits could not distinguish the immunoglobulins																											on the contract of the contrac	or because the performance of these has a based on the PCR-reference standard, the determination of the actual prevalence of the viral infection is limited and cannot reveal the actual status of participants with viral load values that are below the PCR detection limit
NPV (%)	80.0	NAb	NA <sup>b</sup>	85.71 (90/105)	NAb	NA <sup>b</sup>	(58/70)	NAb	NAb	81.94	(59/72) NA <sup>b</sup>	Z Z Z Z Z	82.86	(58/70)	Z Z	84.79	(59/70)	NA <sup>b</sup>	NAb	85.07	(27/67)	Z Z	NA <sup>5</sup>	(60/74)	NAb	NAb	88.46	(138/156)	a A V V V	0 17	(40/47)
(%) Add	100.0 (18/18)	NAb	NAb	95.83 (23/24)	NAb	NA <sup>b</sup>	(21/23)	NAb	NAb	96.15	(72/76) NA <sup>b</sup>	ς α X	90.48	(19/21)	AZ Z	95.74	(20/21)	NA <sup>b</sup>	NAb	88.46	(23/26)	a S	NAN O	(18/18)	NAp	NAb	84.62	(22/26)	a A A	000	(36/36)
Specificity (%)	100.0 (60/60)	NAb	NA <sup>b</sup>	98.9 (90/91)	NAb	NA <sup>b</sup> 96 67	(58/60)	NAb	NAb	98.33	(59/60) NA <sup>b</sup>	Y S	29.96	(58/60)	Z Z	98 33	(59/60)	NA <sup>b</sup>	NAb	95.0	(21/60)	a S	NA <sup>2</sup>	(09/09)	NAb	NAb	97.18	(138/142)	N N	000	(07/07)
Sensitivity (%)	54.55 (18/33)	N de	NA <sup>b</sup>	60.53 (23/38)	NAb	NA <sup>b</sup>	(21/33)	NAb	NAb	65.79	(25/38) NA <sup>b</sup>	Z Z	61.29	(19/31)	A Z	64.52	(70/31)	NA <sup>b</sup>	NΑδ	69.70	(23/33)	A ?	NA" 56.25	(18/32)	NAb	NAb	55.0	(22/40)	a Z V V	77	(36/43)
Sample	93	93	93	129	129	129	2	93	93	86	80	86	91	Ç	91	91	1	91	91	93		93	93 02	70	92	92	182		182 182	0	0
Туре	Total	IgG	IgM	Total	IgG	IgM Ig+oT	וסכם	IgG	IgM	Total	ריים	IaM	Total	(	lgo Mel	Total	Ď.	IaG	MpI	Total		. IgG	Total	Ď.	IgG	MgI	Total		IgG IaM	) to	
nued Product name/source	56 RDT 1		6 1	RD1 2		DDT 2				RDT 4			RDT 5			RDT 6	2			RDT 7			DDT 8	2			RDT 9				matographic CARD
<b>Table 1.</b> Continued	Adams et al.ª 56																														et al. <sup>a</sup> 57

_	3
2	שַׁ
+	=
2	= 5
Ċ	ز.
7	4
4	ธั
F	2

Limitation of study																	a. Study location was restricted to a healthcare	center, which produced data that needs to be	reinforced using a multicenter study	b. No consideration of the study participants with a	(100/136) range of clinical manifestations so as to generate non-biased data	c. Validation of just a kit		d. Poor diagnostic performance for IgM based on	ivity
NPV (%)	35.11	(40/47)	70.18	(40/57)	90.91	(40/44)	90.91	(40/44)	38.89	(40/42)	93.02	(40/43)	93.02	(40/43)	35.11	(40/47)	75.76 a. Study	(100/132) cente		73.53 b. No co	100/136) range non-b	60.61 c. Valida	(100/165)	d. Poor c	sensitivity
IN (%) Adc				(26/26)												(36/36)	100.0	~			(24/24)		(25/25) (		
I. –	100.0	(40/40)	100.0	3)(40/40)	100.0	(40/40)	100.0	(40/40)	100.0	(40/40)	100.0	(40/40)	100.0	(40/40)	100.0	(40/40)	100.0	(100/100)	0	100.0	(100/100)	100.0	(100/100)		
Sensitivity Specificity (%)	83.72	(36/43)		60.47(26/43)(40/40)	90.70	(39/43)	90.70	(39/43)	88.37	(38/43)	93.02	(40/43)	93.02	(40/43)	83.72	(36/43)	94.49	(28/90)	(	0.09	(24/90)	27.78	(25/90)		
Sample	83		83		83		83		83		83		83		83		190		()	190		190			
Type	IgG		MgI		Total		IgG		IgM		Total		IgG		MgI	ı	Total		(	IgG		IgM			
Product name/source					SARS-CoV-2	immunochromatographic CARD 2					SARS-CoV-2 immunofluorescence	CARD 3					AllTest COV-19 IgG/IgM kit (AllTest	Biotech, Hangzhou, China)							
Citation																	Pérez-García	et al.ª 58							

Abbreviations: CEFA, cyclic enhanced fluorescence assay; CLIA, chemiluminescence immunoassay; LFIA, lateral flow immunoassay; MNT, microneutralization test; NAª, not applicable; NPV, negative predictive value; POCT, point of care test; POS, postonset of symptoms; PPV, positive predictive value; PRNT, plaque reduction neutralization test.

 $^{\rm o}$  diagnostic performance performed with reference to RT-PCR.  $^{\rm b}$  diagnostic performance performed with reference to a microneutralization test (MNT).  $^{\rm c}$  Diagnostic performance performed with reference to plaque-reduction neutralization test (PRNT).

All computed values were PSO.
 All products with ≥95% each for sensitivity, specificity, PPV and NPV may be used for epidemiological purposes. Furthermore, performance characteristics ≥95% values reported from acute COVID-19 samples could be considered for clinical use (in conjunction with clinical presentations of patients).

	Product name/		Sample	Sensitivity	Specificity			
Citation	source	Type	size	(%)	(%)	PPV (%)	(%) AAN	Limitation of study
Van Elslande et al.ª <sup>52</sup>	Euroimmun	Total	256	NA <sup>b</sup>	NA <sup>b</sup>	NAb	NAb	<ul> <li>Samples used to determine both specificity and sensitivity were challenging</li> </ul>
		IgG	256	55.56 (85/153)	96.12 (99/103)	95.51 (85/89)	56.28 (99/167)	b. Diagnostic performance data both for total antibody and IgM were not made available
		IgM	256	NA <sup>b</sup>	NA <sup>b</sup>	NAb	NAb	
Zhao et al.ª 17	COVID-19 ELISA kit (Beijing Wantai Biological Pharmacy Enterprise Co. Ltd)	Total	386	93.06 (161/173)	99.06 (211/213)	98.77 (161/163)	94.62 (211/223)	<ul> <li>Sampling was for upper respiratory tract instead of lower respiratory tract with higher sensitivity for RNA tests</li> </ul>
	-	IgG	386	64.74 (112/173)	98.98 (195/197)	98.25 (112/114)	76.17 (195/256)	<ul> <li>b. No evaluation of the persistence of antibodies as sampling was performed during the acute phase of the participants</li> </ul>
		IgM	370	82.66 (143/173)	98.59 (210/213)	97.95 (143/146)	87.5 (210/240)	c. Cross-reactivity studies were not performed for the serological kits
Xiang et al.ª 59	Sandwich ELISA kit (Livzon Inc, Zhuhai, China, lot numbers 20200308 [IgM] and 20200308 [IaG])	Total	126	83.33 (55/66)	100 (60/60)	100 (55/55)	84.51 (60/71)	a. Small sample sizes were used to determine the seropositive rate of IgG
		IgG	126	83.33 (55/66)	95.0 (57/60)	94.83 (55/58)	83.82 (57/68)	b. Unreliable for testing within the window period of infection due to misdiagnosis; retesting was recommended for those with early serone antive immunopholylins
		MgI	126	77.27 (51/66)	100 (60/60)	100 (51/51)	80.0 (60/75)	

Table 2. Continued	nued							
Citation	Product name/ source	Туре	Sample size	Sensitivity (%)	Specificity (%)	PPV (%)	(%) NPV	Limitation of study
Jääskeläinen et al. <sup>b 53</sup>	Anti-SARS-CoV-2 IgA and IgG EIA (Euroimmun, Lu <sup>-</sup> beck,	Total	123	87.8 (36/41)	86.59 (71/82)	76.6 (36/47)	93.42 (71/76)	a. No extensive investigation on prozone phenomenon capable of causing false-neartive results
		IgG	123	70.73 (29/41)	86.59 (71/82)	72.5 (29/40)	85.54 (71/83)	b. IgA detection is not useful for screening purposes but can only be applied for follow-up investigations in patients with proven COVID-19
		IgA	123	87.8 (36/41)	68.29 (56/82)	58.06 (36/62)	91.8 (56/61)	
Jääskeläinen et al.ª 60	Anti-SARS-CoV-2 IgA and IgG EIA (Euroimmun, Lu¨beck, Germany)	Total	40	92.86 (13/14)	92.31 (24/26)	86.67 (13/15)	96.0 (24/25)	a. Small sample size
		IgG	40	92.86	92.31 (24/26)	86.67	96.0	
		IgA	40	78.57 (11/14)	73.08 (19/26)	(11/18)	86.36 (19/22)	
Geurtsvan Kessel et al. <sup>c 43</sup>	Wantai SARS-CoV-2 total Ig and IgM ELISA (Beijing Wantai Biological Pharmacy	Total	226	98.68 (75/76)	99.33 (149/150)	98.68 (75/76)	99.33 (149/150)	a. Not entirely adequate for population screening during an early phase of the pandemic
		IqG	226	NAb	NAb	NAb	NAb	
		IgM	226	89.47	98.67	97.14	94.87	
	Anti-SARS-CoV-2 Ing and InA	Total	237	(9//89)	(148/150) 99 38	(0//89)	(148/156) 98 77	
	ELISA assay (EUROIMMUN Medizinische Labordiagnostika AG)		) N	(74/76)	(160/161)	(74/75)	(160/162)	
	,	IgG	237	81.58	99.38	98.41	91.95	
		IgA	237	97.37	93.79 (151/161)	88.10 (74/84)	98.69 (151/153)	

	>	ased on											ot detect les of n of	e ion of was not urther		
	Limitation of study	a. Diagnostic performance based on sensitivity was very poor			a. Small sample size								a. Both ELISA assays could not detect immunoglobulins in samples of participants with mild form of	b. The study on the protective mechanism and the duration of immune response, which was not performed in detail, was further	pesodold	
	(%) ANN	53.33 (16/30)	53.33 (16/30)	53.33 (16/30)	95.65 (22/23)	95.65	75.86	(22/29)	(20/20)	100.0	(20/20)	80.0 (20/25)	NAb	60.61 (20/33)	NAb	NAb
	(%) Add	100.0 (12/12)	100.0 (12/12)	100.0 (12/12)	93.75 (15/16)	93.75	90.91	(10/11)	94.12	94.12	(16/17)	92.31 (12/13)	NAb	100.0 (32/32)	NAb	NAb
	Specificity (%)	100.0 (16/16)	100.0 (16/16)	100.0 (16/16)	95.65 (22/23)	95.65	95.65	(22/23)	95.24	95.24	(20/21)	95.24 (20/21)	N A <sup>b</sup>	100.0 (20/20)	NAb	g N
	Sensitivity (%)	46.15 (12/26)	46.15 (12/26)	46.15 (12/26)	93.75 (15/16)	93.75	58.82	(10/17)	(16/16)	100.0	(16/16)	70.59 (12/17)	NA <sup>b</sup>	71.11 (32/45)	NAb	g V V
	Sample size	42	45	42	04	40	40	Ċ	SS SS	38		38	92	65	65	08
	Туре	Total	IgG	IgA	Total	IgG	IgA	H	lotal	IgG	1	IgA	Total	IgG	AgI	Total
pı	Product name/ source	EUROIMMUN anti-SARS-CoV-2 IgA and IgG ELISA test			Euroimmun SARS-CoV-2 IgG ELISA (Euroimmun, Lübeck, Germany)				Vircell COVID-19 ELISA IGG (Vircell Spain S.L.U., Granada, Spain)	-			Anti-SARS-CoV-2 ELISA IgG (S1 protein-based) (Euroimmun, Lübeck,			Virotech SARS-CoV-2 IgG ELISA (N protein-based) (Virotech Diagnostics GmbH Riisseisheim, Germany)
Table 2. Continued	Citation	Müller et al. <sup>d 61</sup>			Kohmer et al. <sup>c 54</sup>								Kohmer et al.° 62			

	Limitation of study			Prolonged average sampling collection period was 12 d, which could influence the diagnostic performance of assays	Based on the use of gRT-PCR as reference protocol for the study, there is a possibility of missing positive races whose rescriptions viral land is	lower than the detection limit for PCR						All assays indicated moderate cross-reactivity with samples from participants for other communicable and non-communicable disorders			
	Limitat			<ul> <li>a. Prolonged average sampling collection period was 12 d, which could influence the diagnostic performance of assays</li> </ul>	b. Based on the use of dXT-PCR as reference protocol for the study is a possibility of missing positive	lower than the						<ul> <li>a. All assays indicated moderate cross-reactivity with samples f participants for other commun and non-communicable disord</li> </ul>			
	NPV (%)	70.0 (35/50)	NAb	82.88 (92/111)	77.31 (92/119)	00 0	(92/111)	76.19	(96/126)	76.19 (96/126)	71.11 (96/135)	98.02 (346/353)	97.30 (324/333)	95.69	(346/353)
	PPV (%)	100.0 (30/30)	NAb	95.83	95.45 (84/88)	0 0 0 0	95.65 (92/96)	100.0	(81/81)	100.0 (81/81)	100.0 (72/72)	93.91 (108/115)	78.52 (106/135)	70.63	93.91 (108/115)
Specificity	(%)	100.0 (35/35)	NAb	95.83 (92/96)	95.83 (92/96)	00 00	92.63 (92/96)	100.0	(96/96)	100.0 (96/96)	100.0 (96/96)	98.02 (346/353)	91.78 (324/353)	88.10	(346/353) (346/353)
ity	(%)	66.67	NAb	82.88 (92/111)	75.68 (84/111)	0	(92/111)	72.97	(81/111)	72.97 (81/111)	64.86 (72/111)	93.91 (108/115)	92.17 (106/115)	87.83	93.91 (108/115)
Sample	size	80	80	207	207	700	/07	207		207	207	468	468	894	468
	Туре	IgG	IgA	Total	IgG	< [	IgA	Total		IgG	IgM	Total	IgG	IgM	IgA
Product name/	source			Euroimmun anti-SARS CoV-2 ELISA IgG and IgA assays (Euroimmun, Luebeck, Germany)	·			VIDAS anti-SARS CoV-2 (ELFA)	(BioMérieux, Marcy-l'Etoile, France)			Anti-SARS-CoV-2 IgG, IgM and IgA ELISA tests (ENZY-WELL SARS-CoV-2 ELISA, DIESSE Diagnostica Senese S.p.a.)			
	Citation			Wolff et al.ª 63								Francesca et al. <sup>e 64</sup>			

D
Ū
Š
Ή.
$\subseteq$
0
$\cup$
7
ø
回
.0

		, <del>'</del>									
	Limitation of study	a. The retrospective nature of the study, which involved no fresh samples, could adversely affect the accuracy of results			<ul> <li>a. No detailed data to investigate immunoglobulin-positivity as a correlate of protective immunity.</li> </ul>	b. No further studies to confirm the lack of evidence to establish the	relationship between severity of the disorder and antibody titers		a. Small sample size	<ul> <li>b. Prolonged average sampling collection period, which could affect the diagnostic performance of the kit</li> </ul>	
	(%) ANN	75.9 (63/83)	59.17 (71/120)	74.7 (62/83)	NA <sup>b</sup>	89.29 (50/56)		NAb	NAb	75.68 (84/111)	84.44 (76/90)
	(%) Add	92.31 (108/117)	98.75 (79/80)	91.45 (107/117)	NAb	100.0 (34/34)		NA <sup>b</sup>	NA <sup>b</sup>	96.49 (55/57)	87.18 (68/78)
	Specificity (%)	87.5 (63/72)	98.61 (71/72)	86.11 (62/72)	NAb	100.0 (50/50)		NAb	NAb	97.67 (84/86)	88.37 (76/86)
11.00	Sensitivity (%)	84.38 (108/128)	61.72 (79/128)	83.59 (107/128)	NAb	85.0 (34/40)		NA <sup>b</sup>	NA <sup>b</sup>	67.07 (55/82)	82.93 (68/82)
3	size	200	200	200	06	06		06	168	168	168
	Туре	Total	IgG	IgA	Total	IgG		IgM	Total	IgG	IgA
/ compart to the col	Product name/ source	Euroimmun anti-SARS-CoV-2 ELISA IgG and IgA assays (Euroimmun, Luebeck, Germany)			In-house ELISA recombinant SARS-CoV-2 trimeric spike protein				EUROIMMUN anti-SARS-CoV-2 assay	,	
	Citation	Montesinos et al. <sup>a 55</sup>			Adams et al.ª <sup>56</sup>				Beavis et al.ª 65		

Abbreviations: ELFA, enzyme linked fluorescence assay; IFA, immunofluorescence assay; IFT, immunofluorescence test; MNT, microneutralization assay; NA<sup>b</sup>, not available; NPV, negative predictive value; NT, neutralization test; POS, postonset of symptoms; PPV, positive predictive value; PRNT, plaque-reduction neutralization assay.

<sup>&</sup>lt;sup>a</sup> Diagnostic performance performed with reference to RT-PCR.

<sup>&</sup>lt;sup>b</sup> Diagnostic performance performed with reference to microneutralization test (MNT).
<sup>c</sup> Diagnostic performance performed with reference to plaque-reduction neutralization test (PRNT).
<sup>d</sup> Diagnostic performance performed with reference to NT and IFT.

e Diagnostic performance performed with reference to IFA.

Note:

All computed values were POS.
 All products with ≥95% each for sensitivity, specificity, PPV and NPV may be used for epidemiological purposes. Furthermore, performance characteristics ≥95% values reported from acute COVID-19 samples could be considered for clinical use (in conjunction with clinical presentations of patients).

Citation	Product name/source	Туре	Sample Size	Sensitivity (%)	Specificity (%)	(%) /dd	(%) ANN	Limitation of study
Jin et al.ª 23	CLIA test kit Shenzhen YHLO Biotech Co., Ltd	Total	76	88.37 (38/43)	100 (33/33)	100 (38/38)	86.84 (33/38)	a. Sample size used was small as just 43 lab-confirmed COVID-19 participants and 33
	(China)	IgG	92	88.37	90.91	92.68	85.71	apparently healthy participants b. The period to viral molecular detection and
				(38/43)	(30/33)	(38/41)	(30/35)	to serological investigation was not constant and was based on clinical judgment
		IgM	92	48.84 (21/43)	100 (33/33)	100 (38/38)	86.84	c. The value of serological investigation in participants with severe cases requires
								assessment as those enrolled into the study had mild to moderate COVID-19 cases d. The average period from clinical hallmark onset to serological investigation was long due to the late availability of testing kits e. There was scarse follow-up data on participants who were discharged
Geurtsvan Kessel et al. <sup>c 43</sup>	DiaSorin Liaison XL	Total	122	73.58 (39/53)	98.55	97.5 (39/40)	82.93 (68/82)	a. Lack of sensitivity at the early phase of symptom onset
Müller et al. <sup>d 61</sup>	LIAISON SARS-CoV-2 S1/S2 IgG CLIA test (DiaSorin S1/S2 IgG)	Total	45	NA <sup>b</sup>	NAb	NA <sup>b</sup>	NAb	a. Small size used for the study
	,	IgG	45	61.54 (16/26)	(11/16)	76.19 (16/21)	52.38 (11/21)	b. All test kits missed a great proportion of neutralizing antibody
	SARS-CoV-2 IgG CMIA from Abbott detecting Anti-nucleocapsid IgG antibodies (Abbott N	IgM Total	47 47	o g V V V V	NA <sup>b</sup>	N N Ad D	NA <sup>o</sup>	

Sa Product name/source Type S
Sample Sensitivity Size (%)
Specificity (%)
1 (%) Add
NPV (%) Limitation of study

Specificity (%) NPV (%)		$(33/34)$ $(34/35)$ $(33/44)$ $NA^{b}$ $NA^{b}$	NAb			100.0 100.0 76.09	NA <sup>b</sup>	NA	100.0	(31/31) (40/40) (31/36) NA <sup>b</sup> NA <sup>b</sup> NA <sup>b</sup>	NA <sup>b</sup> a. Poor diagnostic performance based on sensitivity; liaison rapid test kit revealed no adeauacy for clinical use	77.78 80.65	(75/79) (14/18) (75/93)	NAb	95.12 89.19 90.7
Sensitivity (%)	75.56	(34/45) NA <sup>b</sup>	AAN q			75.56	(Ct/tC) NA <sup>b</sup>	NAb	88.89	(40/45) NA <sup>b</sup>	NAb	43.75	(14/32)	NA <sup>b</sup>	80.49
Sample Size	79	79	08			80	80	92	92	92	111	111	, ,	123	123
Туре	IgG	MpI	Total			IgG	IgM	Total	IgG	IgM	Total	IgG	Z	Total	IgG
Product name/source Type	. IgG	X CI	LIAISON XL SARS-CoV-2 Total	S1/S2 IgG CLIA test (DiaSorin S1 and S2	protein-based) (DiaSorin Deutschland GmbH, Dietzenbach, Germany)		MpI	Vircell VIRCLIA automation Total system IgG MONOTEST (CLIA) (S1 and N protein-based) (Vircell Spain S.L.U., Granada, Snain)		MgI	IAISON SARS-CoV-2 IgG (CLIA) (DiaSorin, Saluaaia, Italy)		V	Architect SARS-CoV-2 IgG Total CMIA assay (Abbott, Illinois, USA)	
Citation Pl			LIAI	S. (D	ā Ğ G			Viro Sp. Sp.	, i		LIAI Jääskeläinen (C et al. <sup>b 53</sup> Sc			Arch CN III	

							a-r				ıts		
	Limitation of study	a. Low detection rate at early stage of COVID-19 infection					a. The criteria for evaluating the period of illness onset were retrieved from medical archives and may include imprecisions due to subjectivity in the lack of objective determination of symptoms and periods	b. Low diagnostic performance based on sensitivity		a. Variation in the period between sampling and symptom onsets	b. Late-stage enrolment of study participants	c. None of the test group of participants provided a negative status sample	a. No further study on the relation between antibody levels and protective immune response
	(%) AdN	82.05 (96/117)	NA <sup>b</sup>	NAb	74.02 (94/127)	NAb	61.54 (72/117)	54.96 (72/131)	58.06 (72/124)	75.86 (44/58)	75.86 (44/58)	71.93 (41/57)	95.24 (40/42)
	(%) Add	100.0	NA <sup>b</sup>	NAb	97.5 (78/80)	NAb	100.0	100.0	100.0 (74/74)	100.0 (47/47)	100.0 (47/47)	93.75 (45/48)	100.0 (41/41)
	Specificity (%)	100.0 (96/96)	NA <sup>b</sup>	NAb	97.92 (94/96)	NAb	100.0 (72/72)	100.0 (72/72)	100.0 (72/72)	100.0 (44/44)	100.0 (44/44)	93.18 (41/44)	(100.0 (40/40)
	Sensitivity (%)	81.08 (90/111)	NA <sup>b</sup>	NAb	70.27 (78/111)	NAb	63.11 (77/122)	53.17 (67/126)	58.73 (74/126)	67.21 (47/61)	67.21 (47/61)	73.77 (45/61)	95.35 (41/43)
	Sample Size	207	207	207	207	207	194	198	198	105	105	105	83
	Туре	Total	IgG IgM	Total	IgG	IgM	Total	IgG	IgM	Total	IgG	IgM	Total
inued	Product name/source	Elecsys anti-SARS CoV-2 IgM/IgG assay (Roche Diagnostics, Vilvoorde, Belqium)		Liaison SARS-CoV-2 IgG kit (CLIA) (Diasorin, Saluggia, Italy)			Maglumi 2019-n-Cov IgG and IgM (CLIA)			SARS-CoV-2 antibodies IgM and IgG at cuff-off values 10.0 AU/mL respectively for CLIA kits (Shenzhen YHLO Biotech Co, Ltd, China)			CLIA
Table 3. Continued	Citation	Wolff et al. <sup>a</sup> 63					Montesinos et al.ª 55			Infantino et al.ª 66			Nuccetelli et al. <sup>a 57</sup>

Citation	Product name/source	Туре	Sample Size	Sensitivity (%)	Specificity (%)	(%) Add	(%) NPV	Limitation of study
		IgG	83	95.35 (41/43)	(100.0 (40/40)	100.0 (41/41)	95.24 (40/42)	
		IgM	83	83.72 (36/43)	95.0 (38/40)	94.74 (36/38)	84.44	
Ma et al.ª 67	CLIA RBD-specific anti-SARS-CoV-2 IgA, IgM and IgG kit	Total	669	94.44 (204/216)	90.48 (437/483)	81.6 (204/250)	97.33	a. Irregular and prolonged average sampling duration, which could influence the accuracy of the assay
		IgG	669	96.76 (209/216)	99.79 (482/483)	99.52 (209/210)	98.57 (482/489)	b. No evaluation of the relationship between immunoglobulin levels and severity of the disorder
		IgM	669	96.76 (209/216)	92.34 (446/483)	84.96 (209/246)	98.45 (446/453)	
ian et al.º 68	CLIA test kit Shenzhen YHLO Biotech Co., Ltd (China)	Total	2113	NAb	NAb	Q V	NA <sup>b</sup>	a. Insufficient data on sensitivity for convalescent samples due to limited period after the development of SARS-CoV-2 IgM/IgG assays and access to limited participant demographics
		IgG	2113	95.68 (531/555)	98.07 (1528/1558)	94.65 (531/561)	98.45 (1528/1552)	
		IgM	2113	84.68 (470/555)	98.14 (1529/1558)		94.44 (1529/1614)	
Suhandynata et al.º 69	Diazyme DZ-LITE 2019-nCoV IgG (CLIA) Assay Kit (cat. # 130219015M)/ IgM (CLIA) assay kit (cat. # 130219016M)	Total	289	100.0 (54/5	94.74 100.0 (54/54) 98.72 (232/235§54/57)	94.74 35§54/57)	100.0 (232/232)	a. 50 of the 54 SARS-CoV-2-confirmed participants were hospitalized and were more likely have acute phase infection compared with other average participants that were infected with COVID-19 b. A participant had a medical history of common variable IgG immunodeficiency, which can adversely affect the diagnostic performance of the kit based on sensitivity as the participant was wrongly categorized as false-negative for IgG despite the positive status as revealed using PCR

Sample Sensitivity Specificity Size (%) (%) PPV (%) NPV (%) Limitation of study	c. Insufficient serologic data on SARS-CoV-2 participants with less severe symptoms who	99.14 96.23	_	96.57 97.96	(48/54) (234/235) (48/49) (234/240)
Sample Si Size		289		289	
Туре		IgG		IgM	
Product name/source					
Citation					

Abbreviations: CLIA, chemiluminescence immunoassay; IFT, immunofluorescence test; NA<sup>b</sup>, not available; NPV, negative predictive value; NT, neutralization test; POS, postonset of symptoms; PPV, positive predictive value; PRNT, plaque-reduction neutralization assay; RBD, receptor-binding domain.

<sup>a</sup> diagnostic performance performed with reference to RT-PCR.

<sup>b</sup> diagnostic performance done with reference to microneutralization assay (MNT)

<sup>c</sup> diagnostic performance performed with reference to PRNT.

<sup>d</sup> diagnostic performance performed with reference to NT and IFT.

All computed values were POS.
 All products with ≥95% each for Sensitivity, Specificity, PPV and NPV may be used for epidemiological purposes. Furthermore, performance characteristics ≥95% values reported from acute COVID-19 samples could be considered for clinical use (in conjunction with clinical presentations of patients).

needed to guide laboratories, public health agencies and governments in the appropriate and responsible deployment of POCT, and serological assays more broadly. Currently, the WHO recommends the use of POCT immunodiagnostic assays in research settings only, and not for clinical decision-making until further evidence is available. If Ideally, validation of serological assays, including POCT, should be performed against a serum panel that includes samples from (1) patients at acute and convalescent stages of infection (to assess sensitivity) and (2) patients with other human coronavirus infections (to assess specificity).

Also, serological tests are relevant to fully characterize the SARS-CoV-2-specific antibody response. Differences in the profile of the antibody response across patients might reveal important aspects of the pathogenesis of COVID-19, explaining the great differences observed in the general population. Indeed, the correlation with disease severity and clinical characteristics is poorly understood. Old age and comorbidities seem to increase the risk for a poor outcome of the disease; however, increasing cases of young people who experience severe illness, requiring hospitalization for assistance by mechanical ventilation, may pose questions about the leading factors of disease progression.<sup>32</sup>

Some challenges are posed by the potential cross-reactivity with other human coronaviruses, due to their high homology at the genetic level. The evidence related to this aspect are still controversial; however, SARS-CoV-specific antibodies are undetectable in the sera of patients 6 y after infection. This observation excludes the presence of cross-reactivity in the sera of COVID-19 patients and might make researchers confident about the specificity of these antibodies.<sup>32</sup> Moreover, it would be interesting to understand whether the differences in the progression of the disease might be related to the level of the immune response.

# Accuracy and applications of COVID-19 serological assays

Although various reports of reputable serology assays are incredibly encouraging, product end-users must be pragmatic regarding their accuracy and applicability for COVID-19 clinical and epidemiological use. The unsustainability of RT-PCR tests for the COVID-19 laboratory response in some countries has necessitated a search for alternative assays with high sensitivity and specificity with a short turnaround time from preanalytical (sample collection) to postanalytical phases (availability of test results),<sup>32</sup> thus enabling prompt and large-scale testing for COVID-19. While none of the antibody-based serological assays have been approved by the WHO, a number of them have been approved for clinical and epidemiological use in some countries.<sup>32</sup> Antibody testing might have a useful role in clinically diagnosing COVID-19 patients with late presentations, prolonged symptoms and those with negative results from RT-PCR tests. Furthermore, these tests could be used to monitor the quality and duration of humoral immune response in COVID-19 patients and vaccination. Epidemiologically, SARS-CoV-2 antibody tests can be used for seroprevalence studies in public health research and to inform decisions about returning to work following asymptomatic SARS-CoV-2 infection.

This could offer an opportunity for clinical diagnosis and interruption of transmission through targeted isolation of the most infectious cases and their close contacts.<sup>32</sup> SARS-CoV-2 an-

tibody testing has been shown to have good clinical applications, given the varied symptoms of COVID-19 and reported cases of false-negative results of RT-PCR tests when respiratory swabs are collected >5 d PSO as their sensitivity begins to decline.<sup>32</sup>

Considering this, many researchers are now conducting an independent performance evaluation of these antibody-based assays. For instance, a study referred to as the 'COVID-19 Testing Project' was conducted by the University of California, Massachusetts General Hospital, the Chan Zuckerberg Biohub and the University of California.<sup>33</sup> This study evaluated 10 lateral flow assays and 2 ELISAs to assess performance characteristics for anti-SARS-CoV-2<sup>33</sup> on plasma/serum of 80 symptomatic COVID-19 patients with RT-PCR positive results, 52 non-SARS-COV-2 patients' respiratory viral infections (SARS-CoV-2 RT-PCR negative) and 108 archived sera of blood donors collected in 2018 or earlier.<sup>33</sup>

The assessment found that the assays of the products had varying sensitivities that increased over time, increasing from about 81% to 100% at >20 d PSO.<sup>33</sup> Based on this, it was inferred that anti-SARS-CoV-2 tests were important for longitudinal studies because a negative result may indicate an actively infected person who has not developed a detectable level of antibodies to the virus. Conversely, the proportion of false-positive samples reported from the non-COVID-19 group ranged from 0% to 16%. The detection agreement indices of the lateral flow assays and ELISAs ranged from 75% to 94%.<sup>33</sup>

In another evaluation study of SARS-CoV-2 antibody-based tests by the Chinese company Innovita, anti-SARS-CoV-2 antibodies were found in 83% of COVID-19-confirmed patients with an assay specificity of 96%.<sup>34</sup> After FDA authorization, these tests were anticipated for at-home use.<sup>34</sup> Despite the merits of serological devices, limitations abound due to issues of misdiagnosis following indications of significant false-negative and false-positive results observed during the evaluation of these kits and devices during quality checks.

These rapid test kits have been observed to be unsuitable for testing patients with  $\leq$ 14 d PSO. To augment these lapses, several studies recommend combining both the serological and RT-PCR-based protocols to provide a more accurate diagnosis of COVID-19 instead of only using the molecular testing approach, which introduces a myriad of strenuous demands on diagnostic and healthcare delivery establishments and regulatory bodies, as well as material, financial and human resources meant to sustain testing capacity. 17,19,20,35 Also, there are several studies that have been conducted by diagnostic industries and independent researchers aiming to evaluate the performance characteristics of various anti-SARS-CoV-2 test protocols, some of which have reported promising results.<sup>35-42</sup> It is worth noting that the clinical use of SARS-CoV-2 antibody tests should be on products that evaluated and reported the performance characteristics (especially sensitivity and specificity) during the acute phase of COVID-19.

# Performance characteristics of COVID-19 serological assays

The most promising (best) LFA on total SARS-CoV-2 antibody test had a sensitivity, specificity, PPV and NPV of 100%, 100%, 100% and 100% at days 4, 5, 4 and 5, respectively, while the worst had a

		Antibody assessment	Detection time	Mean time of	Peak period	Period of
Citation	Product name/source	type	range (d)	detection (d)	(p)	decline (d)
Van Elslande et al. <sup>52</sup>	Clungene COVID-19 IgG/IgM rapid test	Total	9-9	5	17-18	NAb
;		IqG	2-6	7	17-18	NAb
		MpI	2-6	2	17-18	NAb
	OrientGene COVID-19 IaG/IaM rapid test	Total	2-6	2	17-18	NAb
		IgG	2-6	7	17-18	NAb
		MpI	2-6	2	17-18	NAb
	VivaDiaq COVID-19 IqG/IqM rapid test	Total	2-6	- 10	17-18	NAb
		IgG	2-6	7	17-18	NA <sup>b</sup>
		MgI	2-6	2	17-18	NA <sup>b</sup>
	StrongStrep COVID-19 IgG/IgM rapid test	Total	2-6	2	17-18	NA <sup>b</sup>
		IgG	2-6	7	17-18	NAb
		MgI	2-6	2	17-18	NAb
	Dynammiker COVID-19 IgG/IgM rapid test	Total	2-6	2	17-18	NAb
		IgG	2-6	7	17-18	NAb
		MpI	2-6	2	17-18	NAb
	Multi-G COVID-19 IqG/IqM rapid test	Total	2-6	2	17-18	NAb
		IqG	2-6	7	17-18	NAb
		MpI	2-6	2	17-18	NAb
	Prima COVID-19 IqG/IqM rapid test	Total	2–6	2	17-18	NAb
		IqG	2-6	7	17-18	NAb
		IgM	9-9	2	17-18	NAb
Montesinos et al. <sup>55</sup>	2019-n-CoV IgG/IgM rapid test cassette (Laboon Time) (Labon Time, Bio Marketina Diagnostics, or Akiva, Israel)	Total	2-0	4	>15	>15
		IgG	2-0	9	>15	>15
		MpI	2-0	4	>15	>15
	Novel coronavirus (2019-n-CoV) antibody IgG/IgM assay (colloidal gold) (Avioq, Bio-Tech, Shandong, China)	Total	2-0	4	>15	>15
		IgG	2-0	9	>15	>15
		MpI	2-0	4	>15	>15
	QuickZen COVID-19 IgM/IgG kit (ZenTech, Angleur, Belgium)	Total	0-7	4	>15	>15
		IqG	2-0	7	>15	>15
		IgM	2-0	4	>15	>15
Pérez-García	AllTest COV-19 IgG/IgM kit (AllTest Biotech, Hangzhou, China)	Total	NAb	NAb	NAb	NAb
et al.38		001	7	c	21 26	96
		ביים ביים	) ·	n (	00-10	0
						C L

Abbreviations:  $\ensuremath{\mathsf{NA}^{\mathsf{b}}}$  , not available; POS, postonset of symptoms. Note: All computed days were POS.

Citation	Product name/source	Antibody assessment type	Detection time range (d)	Mean time of detection (d)	Peak period (d)	Period of decline (d)
Van Elslande et al. <sup>52</sup>	Euroimmun	Total	NAb	NAb	NAb	NAb
		IgG	9-5	9	17-18	NAb
		IgM	NAb	NA <sup>b</sup>	NAb	NAb
Zhao et al. <sup>17</sup>	COVID-19 ELISA kit (Beijing Wantai Biological Pharmacy Enterprise Co. Ltd)	Total	7	7	14-25	>35
		IgG	14	14	25	>35
		IgM	7	4	14	21
Xiang et al. <sup>59</sup>	Sandwich ELISA kit (Livzon Inc., Zhuhai, China, lot numbers 20200308 [IgM] and 20200308 [IgG])	Total	4	4	24	31
		IgG	4	4	24	28
		IgM	4	4	18	28
Padoan et al. <sup>70</sup>	COVID-19 IgG/IgA ELISA kit (Euroimmun Medizinische Laboradiagnostika, Luebeck, Germany)	Total	NA <sup>b</sup>	NAb	NA <sup>b</sup>	NAb
		IgG	NA <sup>b</sup>	NA <sup>b</sup>	NAb	NA <sup>b</sup>
		IgA	4	4	18	34
Jääskeläinen et al. <sup>60</sup>	Anti-SARS-CoV-2 IgA and IgG EIA (Euroimmun, Lu¨beck, Germanv)	Total	11	11	NA <sup>b</sup>	NA <sup>b</sup>
		IgG	12	12	NAb	NAb
		IgA	11	11	NAb	NAb
Okba et al. <sup>29</sup>	Anti-SARS-CoV-2 IgG and IgA ELISA (EUROIMMUN Medizinische Labordiagnostika AG)	Total	72	2	13-21	>21
		IgG	2	2	13-21	>21
		IgA	2	2	11-15	20
Montesinos et al. <sup>55</sup>	Euroimmun anti-SARS-CoV-2 ELISA IgG and IgA assays (Euroimmun, Luebeck, Germany)	Total	2-0	2	>15	>15
		IgG	2-0	7	>15	>15
		IgA	0-7	2	>15	>15

Citation	Product name/source	Antibody assessment type	Detection time range (d)	Mean time of detection (d)	Peak period (d)	Period of decline (d)
Sun et al.71	In-house ELISA produced using N protein (residue 1–419) from baculovirus insect cells (cat. # 40588-V08B, Sino biological, Beijing, China) and S protein (residue 16–685) from HEK293 cells (cat. #40591-V08H, Sino biological, Beijing, China)	Total	7-0	2	8-15	> 22
		IgG	7-0	7	8-15	>22
		IgM	2-0	2	8-15	22
Beavis	EUROIMMUN anti-SARS-CoV-2 assay	Total	NAb	NAb	NAb	NAb
ם מו		IgG	0-2	2	19-49	>50
		IgA	0-2	2	19-49	>50

Citation	Product name/source	Antibody assessment type	Detection time range (d)	Mean time of detection (d)	Peak period (d)	!Period of decline (d)
Long et al. <sup>72</sup>	MCLIA kits (Bioscience Co.; approved by the China National Medical Products Administration)	Total	2-4	2	11-13	>23
		IgG	2-4	4	11-13	>23
		IgM	2-4	2	11-13	>23
Jin et al. <sup>23</sup>	CLIA test kit Shenzhen YHLO Biotech Co., Ltd (China)	Total	1–5	1	16-20	>32
		IgG	1–5	2	16-20	>32
		IgM	1–5		16-20	21–25
Padoan et al 70	CLIA assay (MAGLUMI 2000 Plus)	Total	NAb	NAb	NAb	NAb
;		IqG	NAb	NAb	NAb	NAb
		IgM	4	4	12	34
Padoan et al. <sup>73</sup>	MAGLUMI 2000 Plus 2019-nCov IgM and IgG assays (Snibe, Shenzhen, China)	Total	<5	\ \ \	26-30	>30
		IgG	<5>	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	26-30	>30
		IgM	< 5	< 5	12-13	18-19
Wolff et al. <sup>63</sup>	Elecsys anti-SARS CoV-2 IgM/IgG assay (Roche Diagnostics, Vilvoorde, Belgium)	Total	4	4	11	>24
		IgG	NA <sup>b</sup>	NA <sup>b</sup>	NAb	NAb
		MgI	NAb	NA <sup>b</sup>	NAb	NA <sup>b</sup>
	Liaison SARS-CoV-2 IgG kit (CLIA) (Diasorin, Saluggia, Italy)	Total	NA <sup>b</sup>	NA <sup>b</sup>	NAb	NA <sup>b</sup>
		IgG	4	4	11-13	>24
		IgM	NA <sup>b</sup>	NA <sup>b</sup>	$NA^b$	NAb
Hou et al. <sup>74</sup>	Anti-SARS-CoV-2 CLIA-YHLO kit	Total				
		IgG	c	c	48	>48
		IgM	3	3	30	48
Montesinos et al. <sup>55</sup>	Maglumi 2019-n-Cov IgG and IgM (CLIA)	Total	2-0	4	>15	>15
		IgG	2-0	7	>15	>15
		IgM	7-0	4	>15	>15
Ma et al. <sup>67</sup>	CLIA RBD-specific anti-SARS-CoV-2 IgA, IgM, and IgG kit	Total	NA <sup>b</sup>	NA <sup>b</sup>	NAb	NAb
		IgG	4-10	10	16-41	>41
		IgM	4-10	7	11-30	31-41
		\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	/,_10	7	11_20	71 75

Citation	Product name/source	Antibody assessment type	Detection time range (d)	Mean time of detection (d)	Peak period (d)	!Period of decline (d)
Qian et al. <sup>68</sup>	CLIA test kit Shenzhen YHLO Biotech Co., Ltd (China)	Total	NAb	NAb	NAb	NAb
		IgG	9	9	20	>35
		IgM	9	9	20	35
Suhandynata et al. <sup>69</sup>	Diazyme DZ-LITE 2019-nCoV IgG (CLIA) assay kit (cat. # 130219015M)/IgM (CLIA) assay kit (cat. # 130219016M)	Total	NAb	NA <sup>b</sup>	NAb	NA <sup>b</sup>
		IgG	2-6	9	8-22	>24
		MgI	9-0	m	8-9	14-22

Abbreviations: CLIA, chemiluminescence immunoassay; NA<sup>b</sup>, not available; POS, postonset of symptoms; RBD, receptor-binding domain. Note: 1. All computed values were POS.

sensitivity, specificity, PPV and NPV of 35.95%, 63.6%, 33.3% and 26.2% at days 1, 3, 1 and 2, respectively. The most promising LFA with the best anti-SARS-CoV-2 IgM test had a sensitivity, specificity, PPV and NPV of 95.8%, 100%, 100% and 98.4% at days 5, 6, 6 and 5, respectively, while the least had a sensitivity, specificity, PPV and NPV of 15.7%, 36.4%, 43.2% and 17.4% at days 1, 3, 1 and 3, respectively. The most promising (best) LFA on anti-SARS-CoV-2 IgG test had a sensitivity, specificity, PPV and NPV of 100%, 100%, 100% and 100% at days 8, 9, 8 and 10, respectively, while the least had a sensitivity, specificity, PPV and NPV of 13.2%, 59.9%, 65.1% and 25.0% at days 1, 2, 3 and 2, respectively (Table 1).

The most promising (best) ELISA on total SARS-CoV-2 antibody test had a sensitivity, specificity, PPV and NPV of 93.9%, 100%, 100% and 100% at days 3, 5, 4 and 3, respectively, while the least had a sensitivity, specificity, PPV and NPV of 46.1%, 86.6%, 76.6% and 55.3% at days 1, 3, 2 and 1, respectively. The most promising ELISA with best anti-SARS-CoV-2 IgM test had a sensitivity, specificity, PPV and NPV of 89.5%, 100%, 100% and 95.7% at days 4, 6, 4 and 5, respectively, while the least had a sensitivity, specificity, PPV and NPV of 64.9%, 88.1%, 70.6% and 80.0% at days 1, 3, 2 and 3, respectively. The most promising (best) ELISA on anti-SARS-CoV-2 IaG test had a sensitivity, specificity, PPV and NPV of 100%, 100%, 100% and 100% at days 8, 10, 8 and 9, respectively, while the least had a sensitivity, specificity, PPV and NPV of 46.1%, 86.6%, 72.5% and 56.2% at days 5, 7, 6 and 7, respectively. The most promising (best) ELISA on anti-SARS-CoV-2 IgA test had a sensitivity, specificity, PPV and NPV of 97.4%, 100%, 100% and 98.0% at days 4, 5, 6 and 5, respectively, while the least had a sensitivity, specificity, PPV and NPV of 46.1%, 68.3%, 58.1% and 53.3% at days 14, 13, 14 and 13, respectively (Table 2).

The most promising (best) CLIA on total SARS-CoV-2 antibody test had a sensitivity, specificity, PPV and NPV of 100%, 100%, 100% and 100% at days 2, 3, 2 and 3, respectively, while the least had a sensitivity, specificity, PPV and NPV of 58.7%, 92.3%, 81.6% and 61.5% at days 1, 2, 1 and 1, respectively. The most promising (best) CLIA on anti-SARS-CoV-2 IgM test had a sensitivity, specificity, PPV and NPV of 96.8%, 100%, 100% and 98.5% at days 1, 3, 3 and 2, respectively, while the least had a sensitivity, specificity, PPV and NPV of 63.1%, 90.5%, 84.9% and 58.1% at days 12, 10, 9 and 11, respectively. The most promising (test) CLIA on anti-SARS-CoV-2 IgG test had a sensitivity, specificity, PPV and NPV of 95.7%, 100%, 100% and 98.7% at days 7, 9, 8 and 7, respectively, while the least had a sensitivity, specificity, PPV and NPV of 43.8.1%, 68.7%, 76.1% and 54.9% at days 1, 3, 2 and 1, respectively (Table 3).

The detection, peak and decline periods of blood anti-SARS-CoV-2 IgM, IgG and total antibodies for POCT, ELISA and CLIA vary widely. The most promising of these assays for POCT detected anti-SARS-CoV-2 at day 3 POS in 21.1% (n=19) and peaked on the 15th day in 93.3% (n=21)<sup>58</sup> of COVID-19 patients; ELISA products detected anti-SARS-CoV-2 IgM and IgG at days 2 and 6 in 34.1% (n=38)<sup>71</sup> and in 46.7% (n=15)<sup>52</sup> COVID-19 patients, respectively, and peaked on the eighth day in 92.1% (n=38)<sup>71</sup> of COVID-19 patients. The most promising CLIA product detected anti-SARS-CoV-2 IgM and IgG at days 1 and 4 in 33.3% (n=6)<sup>23</sup> and 60.0% (n=35),<sup>63</sup> respectively, and peaked on the 30th day in 97.8% (n=87)<sup>73</sup> of COVID-19 patients (Tables 4-6).

#### Conclusions

Given the varied performance characteristics of all the sero-logical assays, there is a need to continuously improve their detection thresholds, as well as to monitor and re-evaluate their performances to ensure their significance and applicability for COVID-19 clinical and epidemiological purposes. When found satisfactory, their use will be imperative for scaling up COVID-19 testing in the face of the economic downturn in resource-limited countries. It is recommended that public institutions, private firms, researchers and healthcare policymakers consider the development and evaluation of alternative means of reducing the current PCR test turnaround time and improving the test capacity of serological tests to secure improved epidemiological data for SARS-COV-2.

**Authors' contributions:** INA, LDR and AUE conceptualized and planned the study. INA, AUE, HAS, LU, SM, AUA, HAA, LDR, DA, SH, YU, HYM, AMG, JON, HMC, CCO, OSA, LO, CO, CNE, PEG, LOA and BOPM conducted the literature search and compilation of data. INA, AUE, HAS, LU, SM, AUA, HAA and LDR performed the data curation and statistical analysis. All the authors participated in writing the draft, revised and final versions of the manuscript. All the authors read and approved the final version of the manuscript for intellectual content before submission. INA is responsible for the overall content as guarantor.

**Acknowledgements:** The authors greatly appreciate Gabriel Ilerioluwa Oke for proofreading and copyediting parts of the manuscript.

Funding: None received.

Competing interests: None.

**Ethical approval:** Not applicable (this is a review article).

**Data availability:** This study being a review article, the data presented in the results section and in discussing our main findings are well referenced. However, raw data will be made available on request through the corresponding author (I.N. Abdullahi).

## References

- 1 Decaro N, Lorusso A. Novel human coronavirus (SARS-CoV-2): A lesson from animal coronaviruses. Vet Microbiol. 2020;244:108693. doi: 10.1016/j.vetmic.2020.108693.
- 2 Worldometers.info. Situation Update Worldwide, as of 2 July 2020. Delaware, USA: Dove, 2020. Available at https://www.worldometers.info/coronavirus/#countries [accessed 20 July 2020].
- 3 Padula WV. Why only test symptomatic patients? Consider random screening for COVID-19. Appl Health Econ Health Policy. 2020;18(3):333-4.
- 4 Hellewell J, Abbott S, Gimma A, et al. Feasibility of controlling COVID-19 outbreaks by isolation of cases and contacts. Lancet Global Health. 2020;8(4):e488–96.
- 5 Ueda M, Martins R, Hendrie PC, et al. Managing cancer care during the COVID-19 pandemic: agility and collaboration toward a common goal. J Natl Compr Canc Netw. 2020;18(4):366–9.

- 6 Ranney ML, Griffeth V, Jha AK. Critical supply shortages the need for ventilators and personal protective equipment during the Covid-19 pandemic. N Engl J Med. 2020;382(18):e41.
- 7 Villaraza, Angangco. FDA issuances on evaluation, approval and use of COVID-19 test kits. Available at https://www.lexology.com/library/detail.aspx?g=7689832c-83ce-4683-a5ea-4630eee5788f [accessed 1 June 2020].
- 8 Rashid ZZ, Othman SN, Samat MNA,, et al.. Diagnostic performance of COVID-19 serology assays. Malaysian J Pathol. 2020;42(1):13–21.
- 9 Cui J, Li F, Shi Z-L. Origin and evolution of pathogenic coronaviruses. Nat Rev Microbiol. 2018;17(3):181–92.
- 10 Chan JF-W, Kok K-H, Zhu Z, et al. Genomic characterization of the 2019 novel human-pathogenic coronavirus isolated from a patient with atypical pneumonia after visiting Wuhan. Emerg Microbes Infect. 2020;9(1):221–36.
- 11 Wu A, Peng Y, Huang B, et al. Genome composition and divergence of the novel coronavirus (2019-nCoV) originating in China. Cell Host Microbe. 2020;27(3):325–8.
- 12 Ji W, Wang W, Zhao X, Zai J, Li X. Cross-species transmission of the newly identified coronavirus 2019-nCoV. J Med Virol. 2020;92(4):433-40
- 13 World Health Organization. 2020 Global Surveillance for human infection with coronavirus disease (COVID-2019), interim guidance, Geneva. Available at https://www.who.int/publicationsdetail/global-surveillance-for-humaninfection-with-novel-coronavirus-(2019-ncov) [accessed 4 April 2020].
- 14 Happi C, Ihekweazu C, Oluniyi PE,, et al.. SARS-CoV-2 genomes from Nigeria reveal community transmission, multiple virus lineages and spike protein mutation associated with higher transmission and pathogenicity. Virological. 2020. http://virological.org/t/sars-cov-2-genomes-from-nigeria-reveal-community-transmission-multiple-virus-lineages-and-spike-protein-mutation-associated-with-higher-transmission-and-pathogenicity/494. [accessed 2 June 2020].
- 15 Xiao AT, Gao C, Zhang S. Profile of specific antibodies to SARS-CoV-2: The first report. J Infect. 2020;81(1):147–78.
- 16 Morrison A, Li Y, Loshak H. Serological tests for COVID-19. Ottawa: CADTH; 2020 ( CADTH Horizon Scan; 188:13) .
- 17 Zhao J, Yuan Q, Wang H, et al. Antibody Responses to SARS-CoV-2 in Patients With Novel Coronavirus Disease 2019. Clin Infect Dis. 2020;71(16):2027–34. doi: 10.1093/cid/ciaa344.
- 18 Long QX, Liu BZ, Deng HJ, et al. Antibody responses to SARS-CoV-2 in patients with COVID-19. Nat Med. 2020;26(6):845–8.
- 19 Li Z, Yi Y, Luo X, et al. Development and clinical application of a rapid IgM-IgG combined antibody test for SARS-CoV-2 infection diagnosis. J Med Virol. 2020;92(9):1518–24.
- 20 Hoffman T, Nissen K, Krambrich J, et al. Evaluation of a COVID-19 IgM and IgG rapid test; an efficient tool for assessment of past exposure to SARS-CoV-2. Infect Ecol Epidemiol. 2020;10(1):1754538.
- 21 Mo HY, Xu J, Ren XL, et al. Evaluation by indirect immunofluorescent assay and enzyme linked immunosorbent assay of the dynamic changes of serum antibody responses against severe acute respiratory syndrome coronavirus. Chin Med J. 2005;118(6):446–50.
- 22 Cao W-C, Liu W, Zhang P-H, Zhang F, Richardus JH. Disappearance of antibodies to SARS-associated coronavirus after recovery. N Engl J Med. 2007;357(11):1162-3.
- 23 Jin Y, Wang M, Zuo Z, et al. Diagnostic value and dynamic variance of serum antibody in coronavirus disease 2019. Int J Infect Dis. 2020;94:49–52.

- 24 Guo L, Ren L, Yang S, et al. Profiling early humoral response to diagnose novel coronavirus disease (COVID-19). Clin Infect Dis. 2020;71(15):778–85. doi: 10.1093/cid/ciag310.
- 25 Cerutti A. The regulation of IgA class switching. Nat Rev Immunol. 2008;8(6):421–34.
- 26 Szomolanyi-Tsuda E, Welsh RM. T-cell-independent antiviral antibody responses. Curr Opin Immunol. 1998;10(4):431–5.
- 27 Liu W, Liu L, Kou G, et al. Evaluation of nucleocapsid and spike protein-based enzyme-linked immunosorbent assays for detecting antibodies against SARS-CoV-2. J Clin Microbiol. 2020;58(6): e00461-20.
- 28 Amanat F, Stadlbauer D, Strohmeier S, et al. A serological assay to detect SARS-CoV-2 seroconversion in humans. Nat Med. 2020;26(7):1033–6.
- 29 Okba NMA, Müller MA, Li W, et al. Severe acute respiratory syndrome coronavirus 2-specific antibody responses in coronavirus disease patients. Emerg Infect Dis. 2020;26(7):1478–88.
- 30 Zhang W, Du R-H, Li B, et al. Molecular and serological investigation of 2019-nCoV infected patients: implication of multiple shedding routes. Emerg Microb Infect. 2020;9:386–9.
- 31 Zeng H, Xu C, Fan J, et al. Antibodies in Infants Born to Mothers With COVID-19 Pneumonia. JAMA. 2020;323(18):1848–9. doi:10.1001/jama.2020.4861.
- 32 Weinstein MC, Freedberg KA, Hyle EP, Paltiel AD. Waiting for certainty on Covid-19 antibody tests at what cost? N Engl J Med. 2020;383(6):e37.
- 33 Whitman JD, Hiatt J, Mowery CT, et al. Test performance evaluation of SARS-CoV-2 serological assays. medRxiv. [Preprint]2020 May 17. doi: 10.1101/2020.04.25.20074856.
- 34 Mandavilli A. Coronavirus Antibody Tests: Can You Trust the Results? The New York Times, 2020. Retrieved from: https://www.nytimes.com/2020/04/24/health/coronavirus-antibody-tests.html.
- 35 Lassaunière R, Frische A, Harboe ZB, et al. Evaluation of nine commercial SARS-CoV-2 immunoassays. medRxiv. 2020. doi: https://doi.org/10.1101/2020.04.09.20056325.
- 36 Nicol T, Lefeuvre C, Serri O, et al. Assessment of SARS-CoV-2 serological tests for the diagnosis of COVID-19 through the evaluation of three immunoassays: Two automated immunoassays (Euroimmun and Abbott) and one rapid lateral flow immunoassay (NG Biotech). J Clin Virol. 2020;129:104511.
- 37 Lin D, Liu L, Zhang M, et al. Evaluations of the serological test in the diagnosis of 2019 novel coronavirus (SARS-CoV-2) infections during the COVID-19 outbreak. Eur J Clin Microbiol Infect Dis. 2020;39(12):2271-7. doi: 10.1007/s10096-020-03978-6.
- 38 Chew KL, Tan SS, Saw S, et al. Clinical evaluation of serological IgG antibody response on the Abbott Architect for established SARS-CoV-2 infection. Clin Microbiol Infect. 2020;26(9):1256.e9–11.
- 39 Yang HS, Racine-Brzostek SE, Lee WT, et al. SARS-CoV-2 antibody characterization in emergency department, hospitalized and convalescent patients by two semi-quantitative immunoassays. Clin Chim Acta. 2020;509:117–25.
- 40 Nagura-Ikeda M, Imai K, Tabata S, et al. Clinical Evaluation of Self-Collected Saliva by Quantitative Reverse Transcription-PCR (RT-qPCR), Direct RT-qPCR, Reverse Transcription-Loop-Mediated Isothermal Amplification, and a Rapid Antigen Test To Diagnose COVID-19. J Clin Microbiol. 2020;58(9):e01438-20. doi: 10.1128/JCM.01438-20.
- 41 National SARS-CoV-2 Serology Assay Evaluation Group. Performance characteristics of five immunoassays for SARS-CoV-2: a head-to-head

- benchmark comparison. Lancet Infect Dis. 2020;20(12):1390-1400. doi: 10.1016/S1473-3099(20)30634-4.
- 42 Theel ES, Harring J, Hilgart H, Granger D. Performance characteristics of four high-throughput immunoassays for detection of IgG antibodies against SARS-CoV-2. J Clin Microb. 2020;58:e01243–20. doi: 10.1128/JCM.01243-20.
- 43 GeurtsvanKessel CH, OKBA NMA, Igloi Z, et al. Towards the next phase: evaluation of serological assays for diagnostics and exposure assessment. medRxiv 2020.04.23.20077156. doi: https://doi.org/10.1101/2020.04.23.20077156.
- 44 Cassaniti I, Novazzi F, Giardina F, et al. Members of the San Matteo Pavia COVID-19 Task Force. Performance of VivaDiag COVID-19 IgM/IgG Rapid Test is inadequate for diagnosis of COVID-19 in acute patients referring to emergency room department. J Med Virol. 2020;92(10):1724–7. doi: 10.1002/jmv.25800.
- 45 Porte L, Legarraga P, Vollrath V, et al. Evaluation of novel antigenbased rapid detection test for the diagnosis of SARS-CoV-2 in respiratory samples. Int J Infect Dis. 2020;99:328–33.
- 46 Pan Y, Li X, Yang G, et al. Serological immunochromatographic approach in diagnosis with SARS-CoV-2 infected COVID-19 patients. J Infect. 2020;81(1):e28–32. doi: 10.1016/j.jinf.2020.03.051.
- 47 Pallett S, Denny S, Patel A, et al. Point-of-care serological assays for SARS-CoV-2 in a UK hospital population: potential for enhanced case. Research Square Preprint. 2020. doi: 10.21203/rs.3.rs-28006/v1.
- 48 Spicuzza L, Montineri A, Manuele R, et al. Reliability and usefulness of a rapid IgM-IgG antibody test for the diagnosis of SARS-CoV-2 infection: A preliminary report. J Infect. 2020;81:e53-4.
- 49 Wu J-L, Tseng W-P, Lin C-H, et al. Four point-of-care lateral flow immunoassays for diagnosis of COVID-19 and for assessing dynamics of antibody responses to SARS-CoV-2. J Infect. 2020;81: 435–47
- 50 Green K, Graziadio S, Turner P,, et al.. Molecular and antibody point-of-care tests to support the screening, diagnosis and monitoring of COVID-19. The Centre for Evidence-based Medicine. 2020. https://www.cebm.net/covid-19/molecular-and-antibody-point-of-care-tests-to-support-the-screening-diagnosis-and-monitoring-of-covid-19/
- 51 Mlcochova P, Collier D, Ritchie AV, et al. Combined point of care SARS-CoV-2 nucleic acid and antibody testing in suspected moderate to severe COVID-19 disease. Cell Reports Medicine. 2020;1(6): 100099.
- 52 Van Elslande J, Houben E, Depypere M, et al. Diagnostic performance of seven rapid IgG/IgM antibody tests and the Euroimmun IgA/IgG ELISA in COVID-19 patients. Clin Microbiol Infect. 2020;26:1082-7.
- 53 Jääskeläinen A, Kuivanen S, Kekäläinen E, et al. Performance of six SARS-CoV-2 immunoassays in comparison with microneutralisation. J Clin Virol. 2020;129:104512.
- 54 Kohmer N, Westhaus S, Rühl C, Ciesek S, Rabenau HF. Clinical performance of different SARS-CoV-2 IgG antibody tests. J Med Virol. 2020;92(10):2243–7. doi: 10.1002/jmv.26145.
- 55 Montesinos I, Gruson D, Kabamba B, et al. Evaluation of two automated and three rapid lateral flow immunoassays for the detection of anti-SARS-CoV-2 antibodies. J Clin Virol. 2020;128:104413.
- 56 Adams ER, Ainsworth M, Anand R, et al. Antibody testing for COVID-19: a report from the national COVID scientific advisory panel. medRxiv. 2020.04.15.20066407. doi: https:// doi.org/10.1101/2020.04.15.20066407.
- 57 Nuccetelli M, Pieri M, Grelli S, et al. SARS-CoV-2 infection serology: a useful tool to overcome lockdown? Cell Death Discov. 2020; 6:38.

- 58 Pérez-García F, Pérez-Tanoira R, Romanyk J, Arroyo T, Gómez-Herruz P, Cuadros-Gonzàlez J. Alltest rapid lateral flow immunoassays is reliable in diagnosing SARS-CoV-2 infection from 14 days after symptom onset: A prospective single-center study. J Clin Virol. 2020;129:104473.
- 59 Xiang F, Wang X, He X, et al. Antibody detection and dynamic characteristics in patients with coronavirus disease 2019. Clini Infect Dis. 2020;71:1930–4.
- 60 Jääskeläinen AJ, Kekäläinen E, Kallio-Kokko H, et al. Evaluation of commercial and automated SARS-CoV-2 IgG and IgA ELISAs using coronavirus disease (COVID-19) patient samples. Eurosurveillance. 2020;25(18):2000603 doi: 10.2807/1560-7917.ES.2020.25.18.2000603.
- 61 Müller L, Ostermann PN, Walker A, et al. Sensitivity of commercial Anti-SARS-CoV-2 serological assays in a high-prevalence setting. medRxiv. 2020.06.11.20128686. doi: https://doi.org/10.1101/2020.06.11.20128686.
- 62 Kohmer N, Westhaus S, Rühl C, Ciesek S, Rabenau HF. Brief clinical evaluation of six high-throughput SARS-CoV-2 IgG antibody assays. J Clin Virol. 2020;129:104480.
- 63 Wolff F, Dahma H, Duterme C, et al. Monitoring antibody response following SARS-CoV-2 infection: Diagnostic efficiency of four automated immunoassays. Diagn Microbiol Infect Dis. 2020;98:115140.
- 64 Francesca C, Alessandra B, Daniele L, et al. Evaluation of ELISA tests for the qualitative determination of IgG, IgM and IgA to SARSCOV-2. medRxiv preprint. 2020. doi: https://doi.org/10.1101/2020.05.24.20111682.
- 65 Beavis KG, Matushek SM, Abeleda PF, et al. Evaluation of the EUROIM-MUN Anti-SARS-CoV-2 ELISA Assay for detection of IgA and IgG anti-bodies. J Clin Virol. 2020;129:104468.
- 66 Infantino M, Grossi V, Lari B, et al. Diagnostic accuracy of an automated chemiluminescent immunoassay for anti-SARS-CoV-2 IgM and IgG antibodies: an Italian experience. J Med Virol. 2020;92: 1671–5.
- 67 Ma H, Zeng W, He H, et al. COVID-19 diagnosis and study of serum SARS-CoV-2 specific IgA, IgM and IgG 2 by a quantitative and sensitive immunoassay. medRxiv preprint. 2020. doi: https://doi.org/10.1101/2020.04.17.20064907.
- 68 Qian C, Zhou M, Cheng F, et al. Development and multicenter performance evaluation of fully automated SARS-CoV-2 IgM and IgG immunoassays. Clin Chem Lab Med. 2020;58(9):1601–7. doi: 10.1515/cclm-2020-0548.
- 69 Suhandynata RT, Hoffman MA, Keiner MJ, McLawhon RW, Reed SL, Fitzgerald RL. Longitudinal monitoring of SARS-CoV-2 IgM and IgG seropositivity to detect COVID-19. J App Lab Med. 2020;5:908–20.
- 70 Padoan A, Sciacovelli L, Basso D, et al. IgA-Ab response to spike glycoprotein of SARS-CoV-2 in patients with COVID-19: A longitudinal study. Clinica Chimica Acta. 2020;507:164–6.
- 71 Sun B, Feng Y, Mo X, et al. Kinetics of SARS-CoV-2 specific IgM and IgG responses in COVID-19 patients. Emerg Microb Infect. 2020;9(1): 940–8.
- 72 Long Q-X, Liu B-Z, Deng H-J, et al. Antibody responses to SARS-CoV-2 in patients with COVID-19. Nat Med. 2020;26(6):845–8.
- 73 Padoan A, Cosma C, Sciacovelli L, Faggian D, Plebani M. Analytical performances of a chemiluminescence immunoassay for SARS-CoV-2 IgM/IgG and antibody kinetics. Clin Chem Lab Med. 2020;58(7): 1081–8.
- 74 Hou H, Wang T, Zhang B, et al. Detection of IgM and IgG antibodies in patients with coronavirus disease 2019. Clin Transl Immunol. 2020;9(5):e1136.