



AACC Practical Recommendations for Implementing and Interpreting SARS-CoV-2 Emergency Use Authorization and Laboratory-Developed Test Serologic Testing in Clinical Laboratories

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BACKGROUND: The clinical laboratory continues to play a critical role in managing the coronavirus pandemic. Numerous US Food and Drug Administration emergency use authorization (EUA) and laboratory-developed test (LDT) serologic assays have become available. The performance characteristics of these assays and their clinical utility continue to be defined in real time during this pandemic. The AACC convened a panel of experts from clinical chemistry, microbiology, and immunology laboratories; the in vitro diagnostics industry; and regulatory agencies to provide practical recommendations for implementation and interpretation of these serologic tests in clinical laboratories.

CONTENT: The currently available EUA serologic tests and platforms, information on assay design, antibody classes including neutralizing antibodies, and the humoral immune responses to SARS-CoV-2 are discussed. Verification and validation of EUA and LDT assays are described, along with a quality management approach. Four indications for serologic testing are outlined. Recommendations for result interpretation, reporting comments, and the role of orthogonal testing are also presented.

SUMMARY: This document aims to provide a comprehensive reference for laboratory professionals and health-care workers to appropriately implement SARS-CoV-2 serologic assays in the clinical laboratory and to interpret test results during this pandemic. Given the more frequent occurrence of outbreaks associated with either vector-borne or respiratory pathogens, this document will be a useful resource in planning for similar scenarios in the future.

Introduction

Coronavirus disease 2019 (COVID-19), caused by severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) has resulted in millions of deaths worldwide and continues to spread at the time of this publication (1–3).

The Secretary of Health and Human Services issued a public health emergency declaration for SARS-CoV-2 on January 31, 2020, which allowed the US Food and Drug Administration (FDA) to grant emergency use authorization (EUA) of unapproved medical products or devices. Although the FDA immediately required EUA for SARS-CoV-2 molecular tests, EUA was not required for serologic assays until May 4, 2020. As of January 8, 2021, >200 SARS-CoV-2 serologic tests are available, of which 64 have obtained an EUA (4).

Questions have arisen about how to best utilize and interpret these tests as a result of the limited FDA review process for EUA approval, numerous available tests, varied performance characteristics, and incomplete understanding of the humoral immune response in COVID-19. Interim guidelines were published by several professional organizations (5–8), but no guidance to date provides comprehensive and practical recommendations for the selection, validation, implementation, and quality management of EUA or laboratory-developed test (LDT) serologic tests. To provide assistance on these topics, a panel of clinical diagnostic laboratory

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and industry experts from the AACC reviewed the recommendation document.

This document provides the most up-to-date understanding of host immune responses to SARS-CoV-2, the associated antibody kinetics, and the currently available EUA assays. Clinical utility and limitations are discussed to help laboratories select appropriate tests for their purposes and targeted population needs. The processes and considerations to verify or validate either EUA or LDT serologic tests in a clinical setting are described. In addition, quality management, test interpretation, and orthogonal testing strategies are outlined.

SARS-CoV-2 and the Humoral Immune Response

ANTIGENIC TARGETS

SARS-CoV-2 encodes 4 structural proteins—spike (S), envelope, membrane, and nucleocapsid (N)—of which the S and N proteins are most commonly used for SARS-CoV-2 serologic assays (9–11). The S protein is divided into S1 and S2 subunits; S1 contains the receptor binding domain (RBD), which binds the human angiotensin-converting enzyme 2 receptor, mediating host cell entry; S2 facilitates fusion of the viral and host membranes (11). Distal regions of the S protein (S1, RBD) are the least conserved among members of β -CoV (e.g., SARS-CoV, Middle East respiratory syndrome CoV) and are likely to induce a SARS-CoV-2–specific antibody response. Overall, the SARS-CoV-2 S protein shares 76% homology with SARS-CoV-1 and only about 30% homology with seasonal β -CoVs (e.g., OC43 and HKU1) (12).

The N protein is the most abundantly expressed immunodominant protein among CoVs, functioning to stabilize viral RNA (12–14, 15). It is highly conserved between SARS-CoV-2 and SARS-CoV, with approximately 90% identity (14), but shares only 33% identity with seasonal β -CoVs (12).

ANTIBODY CLASSES

Commercial SARS-CoV-2 serologic assays are available for detection of total antibodies, specific antibody subclasses (IgG, IgM, or IgA), or neutralizing antibodies (nAbs) using qualitative or semiquantitative methods. No clear evidence supports the clinical utility of stand-alone IgM testing (16). IgA-based assays have been reported to suffer from lower specificity compared with IgG-based assays (12) and currently are not recommended for use by either the CDC or the Infectious Diseases Society of America (6, 16). Detection of total antibodies may enhance sensitivity (15, 17–19).

The antibody response to a virus can be split into 2 broad categories—binding and neutralizing. Although binding antibodies inactivate the virus through

mechanisms such as complement activation or opsonization, nAbs inhibit by binding to regions of the virus that directly interact with host cell receptors, effectively blocking viral entry and inhibiting replication. Unlike the detection of binding antibodies, the detection of nAbs requires functional assays. The gold standard is the plaque reduction neutralization test, which is technically challenging to perform, requires live viral and cellular culture, has prolonged turnaround time (days to weeks), and, for SARS-CoV-2, requires biosafety level 3 facilities.

To overcome these challenges, alternative methods have been developed, including pseudovirus-based live cell neutralization assays or blockade-of-binding immunoassays. Pseudovirus neutralization assays can be performed at biosafety level 2 (20), although these assays are still complex, associated with significant analytical variability, and challenging to support in most clinical laboratories. Blockade-of-binding immunoassays can be performed in a 96-well format and can be automated on different immunoassay processing platforms for high-throughput analysis. The nAb assays have played an important role in the development and assessment of SARS-CoV-2 vaccines and in research studies probing the host immune response to infection (21). Given the challenges associated with assay maintenance, lack of standardization, and the currently unknown correlation of nAb titers with protective immunity, the role of nAb assays in the clinical laboratory will likely be limited.

ANTIBODY KINETICS

Understanding the kinetics of the antibody response to SARS-CoV-2 is a prerequisite for test selection and accurate result interpretation. The current understanding of the kinetics of the antibody responses against SARS-CoV-2 are depicted in Fig. 1. Of note, antibody kinetics in specific subpopulations, including immunosuppressed patients, cancer patients, and other subgroups, may differ and continue to be studied.

Unlike viral RNA and antigens, detection of antibodies during the incubation phase is unlikely. Multiple published studies demonstrate that most individuals develop an IgM, IgA, or IgG response within 7–14 days of symptom onset, with >90% of individuals seropositive after 3 weeks (22, 23). IgM and IgA peak and decline earlier than IgG, often within weeks of symptom onset (24–27). IgG antibodies correlate with disease severity, decline at varying rates, and may be detectable for months following infection (28–35). Notably, approximately 4%–10% of the population with confirmed SARS-CoV-2 infection may have either an undetectable or delayed antibody response (36). Regarding antibody longevity, some studies indicate that up to 40% of confirmed individuals become IgG seronegative by the early convalescent phase (37), whereas others have

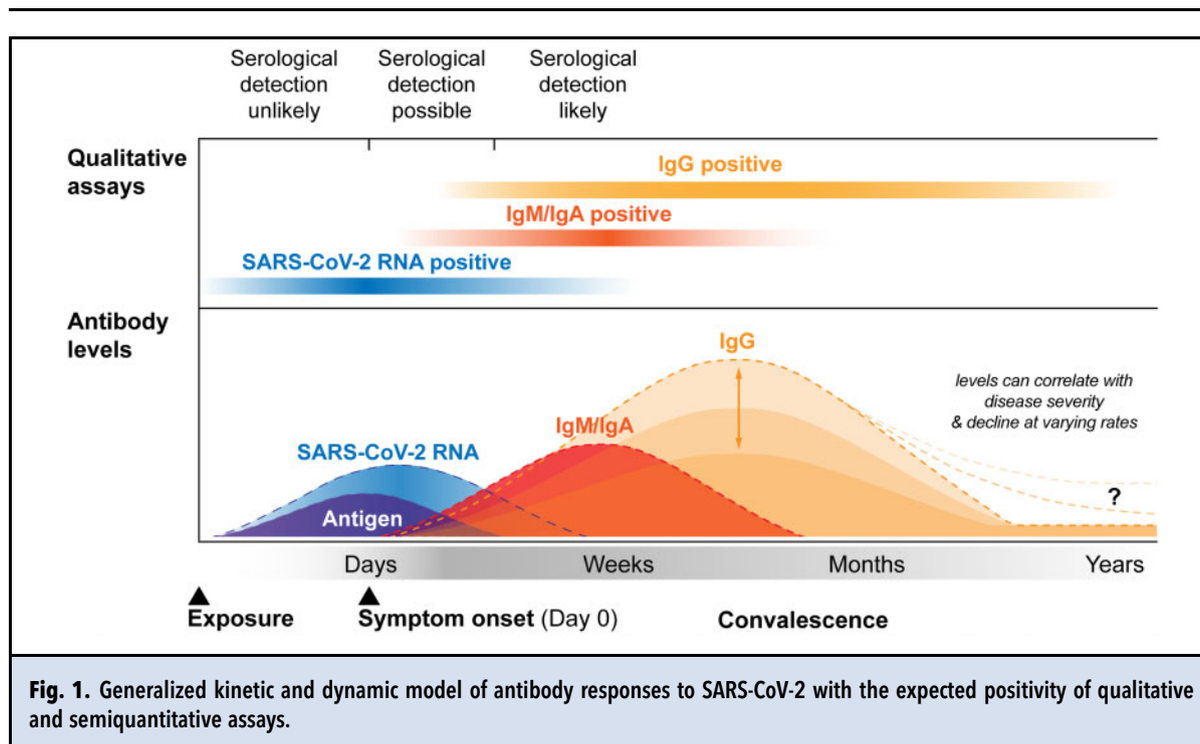


Fig. 1. Generalized kinetic and dynamic model of antibody responses to SARS-CoV-2 with the expected positivity of qualitative and semiquantitative assays.

demonstrated that antibodies decline but remain detectable for months after infection (36, 38, 39, 41). Given these inconsistencies, the precise kinetics of the SARS-CoV-2 antibody response requires further elucidation.

EUA Serologic Tests

ASSAY DESIGNS

Several assay formats for detection of SARS-CoV-2 antibodies have received EUA. Lateral flow assays utilize immunochromatographic chemistry to detect antibodies, usually at the point of care. Manual or semiautomated 96-well ELISAs are also available, as are chemiluminescent immunoassays/chemiluminescent microparticle immunoassays for fully automated, high-throughput platforms. These methods are illustrated in Fig. 2.

CHARACTERISTICS OF EUA SEROLOGIC TESTS

Sixty-four assays received EUA from the FDA as of January 8, 2021. Selected examples are listed in online Supplemental Table 1. The majority detect IgG, followed by IgM or IgG, total antibody, and IgM only. All EUA assays use serum; some accept plasma and, less frequently, whole blood or dried blood spots (42). Currently, serologic testing is not recommended for other sample types such as saliva and cerebrospinal fluid. The most frequent antigen targeted in these assays is the RBD, followed by S (including full S, S1, and S2), and

N. Currently, only one assay uses all 3 antigens. Most current EUA assays are qualitative, with a few being semiquantitative.

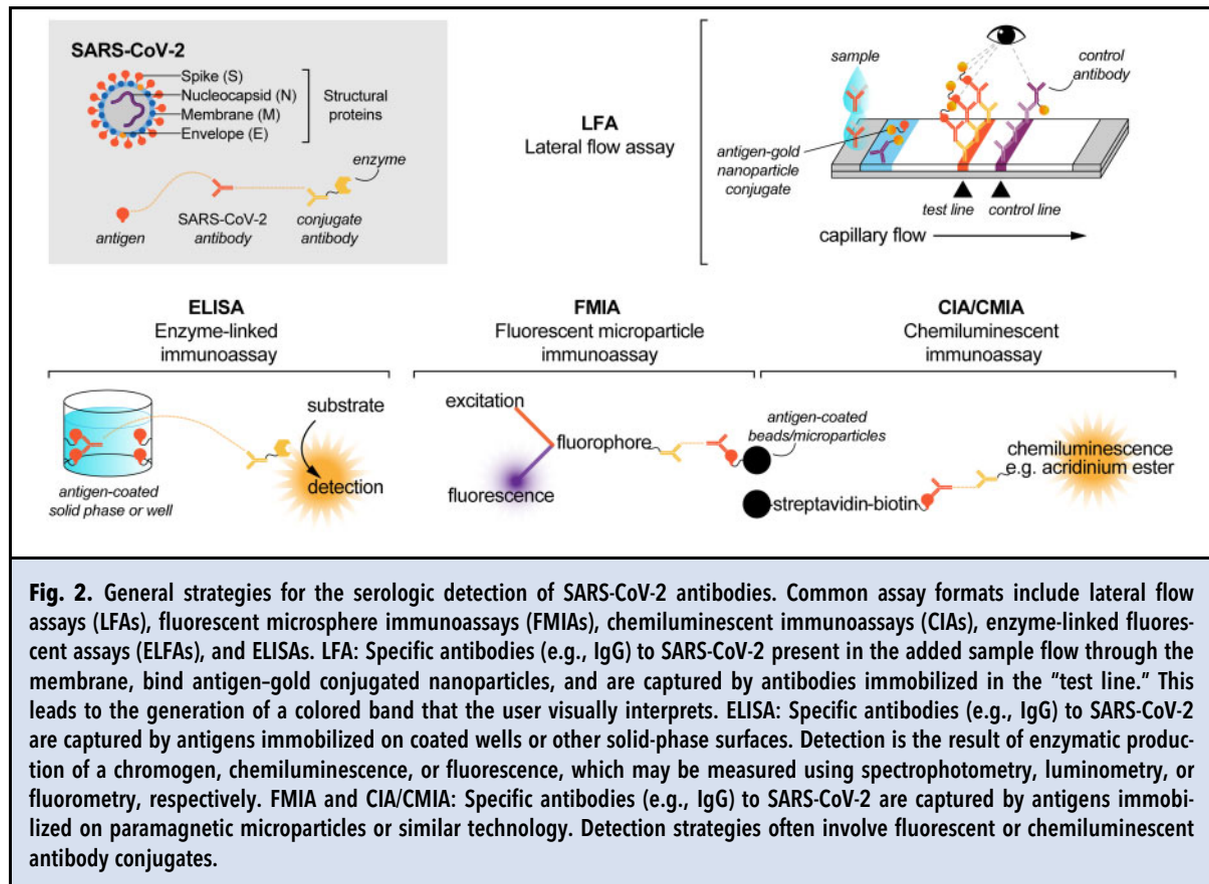
Assessing the relative performance characteristics of each EUA assay is complicated because the approach, the sample size, the sample collection time, and the disease prevalence in the population tested by each manufacturer vary widely. Clinical laboratory professionals should take these variables into consideration when evaluating assay performance.

Utility and Limitations of SARS-CoV-2 Serology

SARS-CoV-2 serologic testing is not recommended as the primary approach for diagnosis of SARS-CoV-2 infection. However, it can be used for supportive diagnosis of COVID-19, manufacture of convalescent plasma (CP), epidemiologic and seroprevalence studies, and vaccine response and efficacy studies (5, 6, 16) (Table 1).

SUPPORTING DIAGNOSIS OF COVID-19

Serologic testing may be helpful to diagnose COVID-19 in symptomatic patients who present later in disease (e.g., >9–14 days after symptom onset) and who test negative by molecular assay, with optimal assay sensitivity occurring at least 2–3 weeks after symptom onset (43, 44). Total antibody or IgG testing may be more



useful for evaluating patients who present later in the disease course (18, 44–48).

Serologic testing, alongside reverse transcriptase PCR (RT-PCR), has been recommended to support the diagnosis of multisystem inflammatory syndrome in children, including for hospitalized individuals <21 years of age who present with fever, inflammation, and multisystem organ involvement following exclusion of other potential diagnoses (45, 49–52). Serologic testing should precede intravenous immunoglobulin or blood product administration because these therapies may influence serologic results.

CP DONOR IDENTIFICATION AND MANUFACTURING

Identification of potential CP donors for COVID-19 CP therapy, which has received FDA EUA, is a recognized application of serologic testing. The FDA continues to refine donor eligibility criteria, identify serologic assays for the manufacture of COVID-19 CP units, and define acceptable antibody thresholds. Originally, the FDA recommended that the qualitative Ortho Clinical Diagnostics SARS-CoV-2 IgG chemiluminescent immunoassay be used in the manufacturing of CP, with signal/cutoff (S/CO) threshold values ≥ 12 considered

“high titer” and preferred for infusion (53). Given that most currently available serologic assays are qualitative, there are limited mechanisms for distinguishing donors with high vs low titers (53). Recently, the FDA updated its COVID-19 CP EUA to include 9 serologic assays for manufacture of CP, including 2 semiquantitative assays (54).

EPIDEMIOLOGIC AND SEROPREVALENCE STUDIES

Determination of seroprevalence is important to characterize the epidemiology of COVID-19 in the community and to support public health efforts (36, 40, 41, 55). However, serologic assays have limitations that may lead to an underestimate of true seroprevalence. First, most commercial assays were developed with patients who were symptomatic with moderate to severe disease. It is unknown whether the cutoffs based on these populations will detect antibodies in cases of asymptomatic or mild disease. Second, a small proportion of the population may never develop detectable antibodies following infection. Third, the accuracy of this approach depends on the prevalence of the disease in the community, as the positive predictive value (PPV) may be low in regions with little disease, even if using a highly specific assay (27, 56).

Table 1. Recommended use of serologic testing and limitations.

| | Description |
|-----------------|--|
| Recommended use | <ul style="list-style-type: none"> • Serologic testing may be offered as an approach to support diagnosis of COVID-19 illness in symptomatic patients and late-phase negative molecular testing or for patients presenting with late complications such as multisystem inflammatory syndrome in children. • Serologic testing can help identify people who may have been infected with or have recovered from the SARS-CoV-2 infection. • Serologic testing can be used to screen potential CP donors and in the manufacture of CP. • Serologic testing can be used for epidemiology and seroprevalence studies. • Serologic testing can be used for vaccine response and efficacy studies. |
| Limitations | <ul style="list-style-type: none"> • False positive results may occur. • Negative results do not preclude acute SARS-CoV-2 infection or viral shedding. • Serologic tests may not differentiate between natural infection and vaccine response. • The durability and kinetics of the humoral immune response continue to be elucidated. <p>Serologic results should not be used for:</p> <ul style="list-style-type: none"> • Determining individual protective immunity. • Return to work decisions. • Grouping individuals in congregate settings. • Assessment of convalescent plasma recipients. • Use of personal protective equipment. • Placement of high-risk job functions. |

VACCINE RESPONSE AND EFFICACY

Available and developing vaccines range from inactivated or live platforms to more novel DNA- or RNA-based preparations, such as the 2 recently authorized vaccines in the United States: Moderna and Pfizer/BioNTech (57). Because the primary target of neutralizing antibodies is the S protein, the majority of the vaccines target the S protein (58–61). Vaccine trials have assessed vaccine efficacy by using end point outcome measures such as prevention of moderate or severe disease due to SARS-CoV-2 infection in the placebo vs vaccinated populations. Vaccine trials have also used several different approaches to assess vaccine response, including binding antibody ELISAs, and both plaque-reduction neutralization tests and pseudovirus-based neutralization assays to determine that the majority of vaccinated individuals developed a robust antibody response, including neutralizing antibodies (13, 14, 57, 59–62). To date, only one assay has received an EUA for detection of nAbs. It is important to note that although a detectable antibody response in a vaccinated individual (including immunosuppressed persons) indicates that an antibody response has developed in response to vaccination, no threshold on any assay is

indicative of vaccine efficacy. Consequently, at this point in time, even semiquantitative or quantitative assays against S protein that can quantify the magnitude of the antibody response to vaccines should not be used to determine vaccine efficacy and protective immunity. This guidance is true not only for binding antibody ELISAs but also for neutralization assays. Currently, there are no recommendations from any professional societies in the United States for monitoring or assessing vaccine response in any population, including immunosuppressed individuals.

Because the S/RBD protein is primarily used for vaccines, the availability of antibody assays that detect N- vs S-specific antibodies may also be useful to distinguish between naturally infected vs vaccinated individuals, but further studies are needed to understand the merits and limitations of this approach.

Performance Verification of EUA Assays

Verification studies for nonwaived EUA assays are the same as those for FDA-approved or cleared assays. However, waived EUA tests should be verified in a manner similar to moderately complex, nonwaived tests.

Further resources for detailed method verification protocols are available through the CLSI (online [Supplemental Table 2](#)).

REGULATORY AND ACCREDITATION REQUIREMENTS

Clinical laboratories in the United States are required by CLIA to verify assay performance of unmodified, FDA-approved or cleared, and EUA assays and must adhere to manufacturer instructions. Several accreditation organizations are available; laboratories should refer to the specific requirements. The College of American Pathology (CAP) is used as an example to discuss some specific requirements for EUA verification:

1. Ensure testing personnel are properly trained and qualified based on test complexity authorized by the FDA;
2. Perform testing as outlined in the EUA without modification;
 - a. Any deviation from instructions for use will render the assay an LDT, which needs to be validated (see Validation of LDTs).
3. Verify test method performance following the CAP's All Common Checklist:
 - a. COM.40300—The laboratory must assess analytical accuracy, analytical precision, and reportable range (as appropriate).
 - b. COM.40475—Laboratory director must sign the laboratory's written assay assessment.
 - c. COM.40500—Laboratory understands analytical interferences for each test and has a plan of action when present.
4. Update the laboratory's activity menu.

Next, we consolidate and expand on prior recommendations to provide a systematic approach for EUA assay verification (63–65).

SAMPLE COLLECTION

Sample type, target population (e.g., symptomatic, asymptomatic, ambulatory, hospitalized, pediatric, pregnant patients), and number of positive samples may be difficult to discern early on in a public health emergency. The following strategies are recommended.

Positive samples.

1. Residual, unmodified patient samples (preferred) may be collected after testing positive on a comparative EUA assay. Comparator assays should be matched to sample matrix, antibody class(es), and antigenic targets for optimal evaluation. If a comparator EUA assay is unavailable, samples collected from patients confirmed by RT-PCR can be used, with knowledge of

days after symptom onset or first RT-PCR-positive result.

2. Residual, positive samples with an increased S/CO may be mixed, at different ratios, with one or more confirmed negative patient samples to generate a range of S/CO-positive samples.
3. Commercially verified materials (e.g., positive QC, patient, or pooled patient samples) may be used in an emergent situation if residual, positive patient samples are not available. However, judicious selection of third-party materials must be performed to mitigate possible matrix effects.
4. Sample and antibody stabilities should be considered. Laboratories should use manufacturers' package inserts as a guide and can also validate alternative stability time frames. Limited studies are available in CP settings (66–68).

Negative samples.

1. Residual, unmodified, prepandemic patient samples were collected and properly stored.
2. Residual, unmodified patient samples (matched to sample matrix, antibody class(es), and antigenic targets) were collected after testing negative on a comparative EUA assay.
3. Commercially verified materials may be used if the other samples are limited or not available.

ACCURACY

Accuracy is verified by assessing the result concordance with either another EUA assay or clinical correlate, reflecting assay clinical sensitivity and specificity. The following recommendations are for accuracy assessments.

Single analyte or total analyte. A minimum of 10 negative and 10 positive samples per sample type should be used. For total antibody tests, it is optimal to use known positive patient samples from each antibody class.

Multiple differentiated analytes. Accuracy verification must be demonstrated with known positive samples for each antibody class that could be reported. Combinations of a minimum of 20 samples (e.g., IgM–/IgG+, IgM+/IgG–, IgM+/IgG+, IgM–/IgG–) should be used to assess class-specific positive and negative agreement and to verify clinical specificity. For SARS-CoV-2, it may be challenging to identify IgM+/IgG– samples given concurrent seroconversion.

PRECISION

The reproducibility and repeatability of an EUA assay around the positive cutoff must be verified. For

qualitative assays, a positive and negative sample can be used, with the positive sample near the cutoff. Semiquantitative assays should be evaluated as quantitative assays, and samples should span low, mid, and high S/CO, with at least one sample near the cutoff. The intra- and interday precision experiments should test both positive and negative samples over 10 replicates on the same day or over 10 runs on a minimum of 5 days and over multiple shifts, respectively. Precision for single-use lateral flow assays should be assessed for interday only over 5 days with multiple testing operators.

REPORTABLE RANGE

For semiquantitative or quantitative serologic assays with EUA, the reportable range must be verified. Verification should be done by using nondiluted, known standards of anti-SARS-CoV-2 antibodies, such as the recently available standard from the World Health Organization (69) or, if unavailable, an alternative calibrator lot or patient samples that span the analytic measuring range. Future standardization of quantitative assays to a single international standard will be essential for accurate assessment of antibody levels once a protective immunity threshold is established.

Validation of LDTs

An in vitro diagnostic test that is designed and used in a clinical setting by a single laboratory is considered an LDT. Clinical laboratories authorized to perform high-complexity testing under CLIA must perform thorough LDT validation studies before patient testing. This section will discuss minimum validation requirements of LDTs that go beyond those necessary for EUA assay verification with respect to sensitivity and specificity, the establishment of assay result cutoffs, class specificity, and carryover.

REGULATORY AND ACCREDITATION REQUIREMENTS

Typically, following the declaration of a public health emergency by the US Department of Health and Human Services, any clinical test used to diagnose that condition, regardless of type (i.e., molecular or serologic or antigen), requires EUA. On August 19, 2020, the EUA requirement for COVID-19 laboratory assays was removed to ease regulatory burdens placed on high-complexity CLIA laboratories capable of developing LDTs (70). The CAP and other accreditation organizations provide specific requirements for clinical laboratories that must be used in the implementation of LDTs (71).

SAMPLE COLLECTION

Generally, LDTs require additional samples to establish assay performance compared with EUA assays that

require verification only. The FDA recommends at least 30 positive samples and 75 antibody-negative (or pre-COVID-19) samples in their guidance for EUA applications (4). In situations where an assay using 75 negative specimens does not demonstrate >95% specificity, or if 75 specimens are not available, the FDA recommends specific cross-reactivity studies with samples known to be positive for a variety of potentially cross-reactive antibodies or those directed against other respiratory pathogens. It is also our recommendation to collect at least 30 positive (with known days from symptom onset) and 75 negative samples (ideally 100–200).

ANALYTIC SENSITIVITY AND SPECIFICITY

Sensitivity. Assay sensitivity can be evaluated with well-characterized RT-PCR-positive samples, ideally with chart data that indicate the days from a patient's symptom onset. In this way, assay sensitivity as a function of time can be assessed. It is also valuable to compare performance with another EUA assay, if available.

Specificity, cross-reactivity, and interfering substances. Samples from patients with known acute respiratory infections should be included for any LDT assay assessing the serologic response to SARS-CoV-2. Ideally, these would include samples from patients infected with one of the circulating seasonal human CoVs (NL63, OC43, HKU1, 229E), although data indicating that these CoVs are not a source of significant cross-reactivity on SARS-CoV-2 serologic tests have begun to accumulate. A disclaimer to the effect that cross-reactivity cannot be ruled out should be included if such samples were not evaluated in the validation (72, 73). In addition, samples from patients diagnosed with other infectious and autoimmune conditions known to give false-positive results in immunoassays (e.g., syphilis, Lyme disease, cytomegalovirus, rheumatoid arthritis, etc.) should be included.

Investigation of other interfering substances is another core component for determining an assay's analytic specificity (e.g., hemoglobin, lipids, bilirubin). For laboratories validating an LDT, it is necessary to investigate potential interferences based on assay design and to devise interference validation studies.

ESTABLISHING ASSAY CUTOFFS

Cutoffs for a qualitative or semiquantitative LDT can be established with a limit of blank studies using known negative samples tested repeatedly over several runs (e.g., 20 known negative samples tested by multiple operators on 5 separate runs). The mean optical density (or equivalent readout) and standard deviations from the mean should be calculated, with the assay threshold determined as the mean readout plus 3 to 5 times the standard deviations. Further refinement of cutoffs can

be performed using ROC analysis to optimize sensitivity and specificity. Alternatively, if risk assessment dictates an overriding concern, then cutoffs can be set accordingly (e.g., for 100% specificity).

Assays that report quantitative results, as well as those that indicate neutralization levels, are less commonly used in clinical laboratories and require additional layers of validation. Once a cutoff is established, it is also recommended that the cutoff be verified as required by the respective accreditation agencies.

ANTIBODY CLASS SPECIFICITY

If a claim about antibody class specificity is made for an LDT, it must be validated. Methods for this process include the use of a detection or capture antibody with a known class specificity or class-specific antibody depletion of the sample. As an alternative, the FDA recommends treating samples with dithiothreitol (74), which effectively removes IgM-class antibodies.

CARRYOVER

Clinical laboratories should perform an assessment to verify that a positive result was not due to positivity from a nearby high-titer positive sample (e.g., for probe-based instruments). This is commonly performed by alternating testing of a negative sample before and after a positive sample with a high index or S/CO value. If carryover cannot be eliminated from the assay, it is recommended to assess the impact on accuracy of a positive and a negative result. Carryover should not exceed 20% of the lower limit of quantitation, according to the FDA, and additional details are available through CLSI EP10-A3-AMD (online [Supplemental Table 2](#)) and CAP.

Other Considerations for EUA Assay Implementation

QUALITY MANAGEMENT

QC must be identified, verified, and implemented for routine SARS-CoV-2 testing based on test complexity and manufacturer instructions. A minimum of 2 levels of QC (positive and negative) should be included with each run of the specified assay. For qualitative and semi-quantitative assays, a negative QC and a positive QC near the cutoff must be run at least daily. For SARS-CoV-2 serologic assays, controls may be provided as part of the assay kit or may need to be sourced separately. For the latter, laboratories can purchase separate controls provided by the assay manufacturer or third-party vendors or use pooled patient samples. Use of assay calibrator material to create assay controls is discouraged, but if needed, the calibrator material must be from a different kit lot. QC material should also match

the analyte detected by the specific assay and patient matrix.

Typically, 20 QC data points on separate days are used to determine the target control mean and SD to establish the range. For vendor material with assigned QC ranges, the laboratory should verify the product. QC performance should be monitored in real time to identify shifts and trends.

Laboratories should participate in proficiency testing using either vendor products or an alternative assessment program. Finally, because EUA assays were not extensively evaluated, laboratories may implement a more rigorous quality management system until assay reliability is established. This approach may include analysis of additional QC material, performing additional lot-to-lot comparisons, and identifying a partner laboratory for more frequent sample exchanges than the biannual proficiency testing requirement.

PREANALYTICAL CONSIDERATIONS

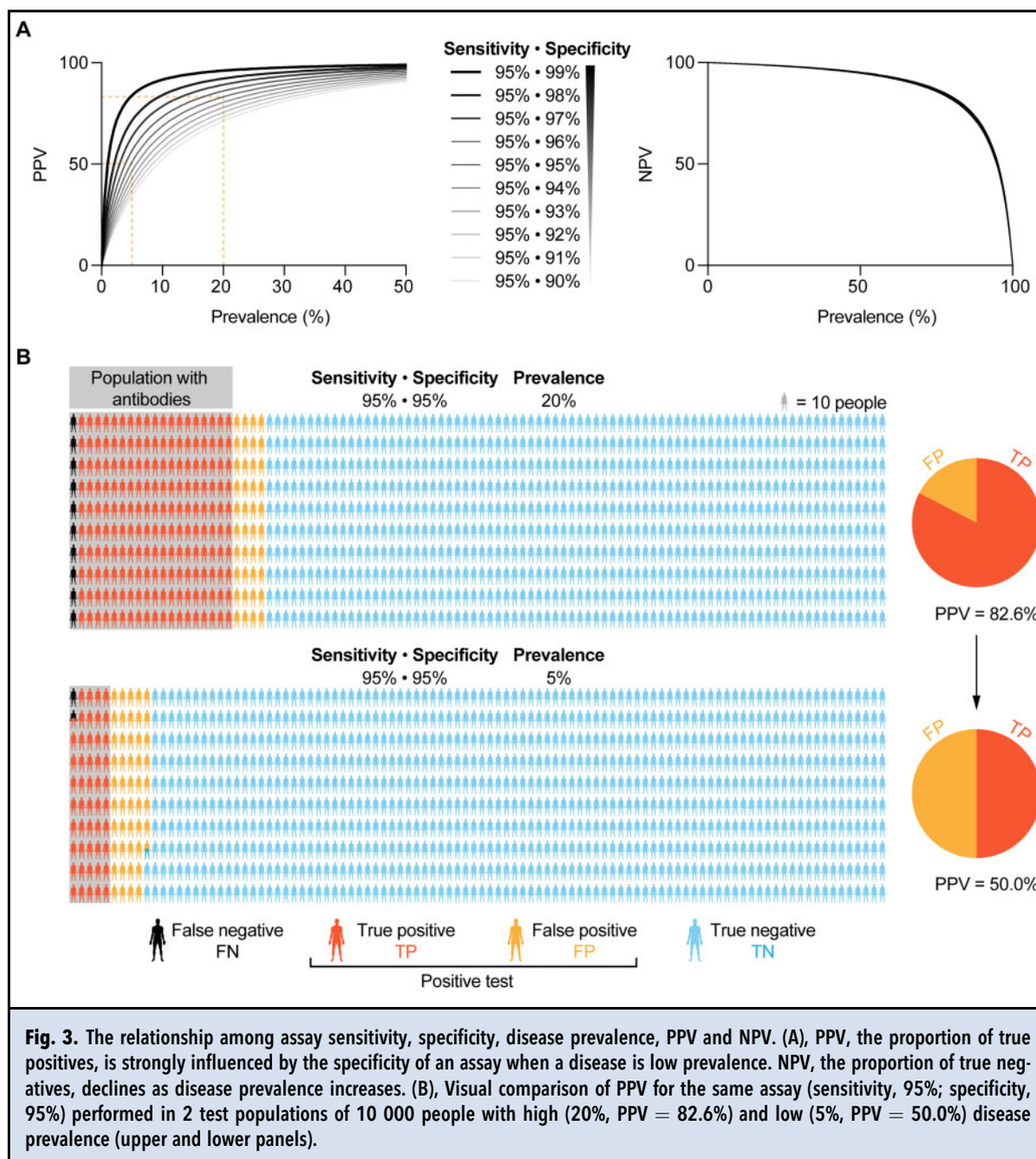
Preanalytical variables should be noted for SARS-CoV-2 antibody tests and thoroughly reviewed to determine possible limitations. These include sampling time and sample stability, as well as potential endogenous and exogenous interferences (e.g., hemoglobin). Time of sample collection is important when selecting positive samples for verification studies. To verify test performance at the optimal reported sensitivity for most current EUA assays, samples collected ≥ 14 days after symptom onset with PCR positivity should be used for verification. The limitation of test performance in patients tested < 14 days before symptom onset with PCR positivity should be clearly stated. For LDTs, clinical sensitivity relative to days from symptom onset needs to be determined during validation (see Validation of LDTs).

If an assay is performed with several sample types, including dried blood spots, laboratories should define and specify collection device, transportation, and preanalytical requirements before patient testing.

Interpretation of Serologic Test Results

The majority of SARS-CoV-2 serologic assays are qualitative in design, and generally, positive results indicate recent or prior SARS-CoV-2 infection. Negative results indicate that SARS-CoV-2 antibodies were not present or were below defined detection limits.

Negative results cannot rule out active or prior infection. Results should be interpreted in the context of antibody class(es) detected (see Supporting Diagnosis of COVID-19) and antigenic target(s), time of sample collection (see Antibody Kinetics), disease severity, and assay analytical performance characteristics (see Analytic Sensitivity and Specificity). Understandings of clinical sensitivity, clinical specificity, and disease prevalence are



also key considerations for interpretation of serologic test results.

IMPACT OF CLINICAL SENSITIVITY, SPECIFICITY, AND DISEASE PREVALENCE

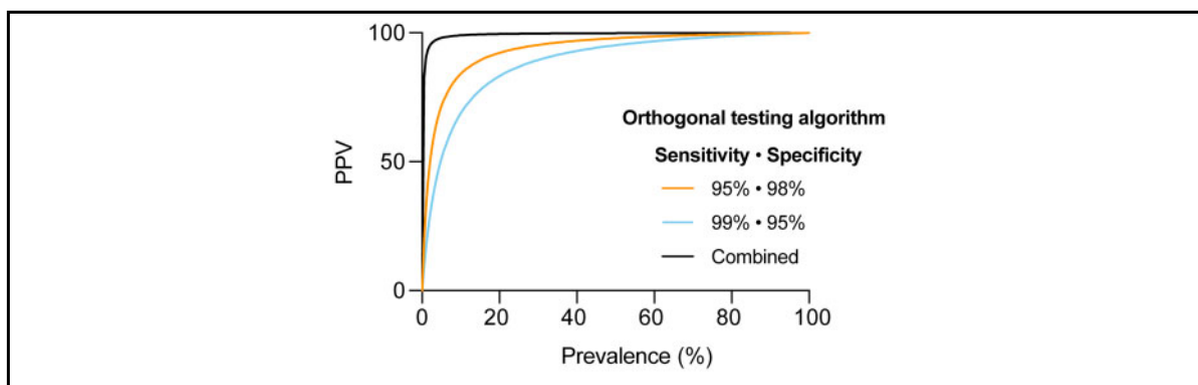
To minimize potential false positives and to be of clinical value, the CDC and the Infectious Diseases Society of America have suggested using tests with clinical sensitivity and specificity of $\geq 99.5\%$.

PPV and negative predictive value (NPV) depend on disease prevalence in the target population and on assay clinical sensitivity and specificity. They indicate the percentage of probability that a positive (or negative) test result will correctly identify individuals with (or without) antibodies in a given population. An assay with 95% sensitivity and 90%–99% specificity was used to illustrate this relationship (Fig. 3A). The PPV increases as specificity increases. Using these sensitivity

Table 2. Relationship of PPV and NPV with sensitivity and specificity in OTA given 2% population disease prevalence.

| | Test 1 sensitivity | Test 1 specificity | Test 2 sensitivity | Test 2 specificity | % initial positive (test 1) | % discordant | Test 1 PPV | T1 + T2 PPV ^a | T1 + T2 NPV ^b |
|-----------|--------------------|--------------------|--------------------|--------------------|-----------------------------|--------------|------------|--------------------------|--------------------------|
| OTA 1 (%) | 95 | 98 | 99 | 95 | 3.9 | 1.9 | 49.2 | 95.0 | 99.9 |
| OTA 2 (%) | 99 | 95 | 95 | 98 | 6.9 | 4.9 | 28.8 | 95.0 | 100.0 |

^aT1, test 1 with sensitivity of 95% and specificity of 98%; T2, test 2 with sensitivity of 99% and specificity of 95%.
^bAn online calculator from the FDA (<https://www.fda.gov/media/137612/download>) is a helpful tool to assess the combined PPV/NPV.

**Fig. 4.** Effect of an OTA on PPV. The performance of 2 tests with different characteristics (test 1: sensitivity, 95%; specificity, 98%; test 2: sensitivity, 99%; specificity, 95%) and their combined performance are shown. Regardless of the order in which the tests are performed, sequential testing can increase PPV in testing populations with low disease prevalence.

and specificity values, the PPV increases very rapidly with increased disease prevalence until it plateaus at $\geq 20\%$ prevalence. The NPV, however, changes minimally with different levels of assay specificity and drops markedly when disease prevalence increases.

A test that has 95% sensitivity and 95% specificity within a population of 20% or 5% antibody prevalence (2000 or 500 individuals have antibodies, respectively, assuming a population of 10,000) is used as an example to show the impact of disease prevalence on PPV and NPV (Fig. 3B). In a population with 20% prevalence, the test would correctly detect 1900 of the 2000 positive individuals, resulting in 400 false-positive results. With 5% prevalence, the test would correctly identify 475 positive individuals, resulting in 475 false-positive results. The PPVs are 82% and 50%, respectively, for these 2 populations.

RESULTS REPORTING

Clear and concise comments are needed to aid interpretation of serologic test results. It is advisable for clinical

laboratories to include a statement that diagnosis of COVID-19 should be performed using molecular tests. According to FDA EUA requirements, reports should include the assay name, and clinicians and patients should have access to the respective assay fact sheets.

The qualitative nature of most EUA tests prohibits reporting of actual S/CO values because (a) the assay-specific values are not standardized and do not indicate an actual concentration of SARS-CoV-2 antibodies; (b) no S/CO cutoffs are currently available to correlate with protective immunity, and (c) different dynamic ranges are available for current platforms. Some examples of results comments are provided in online Supplemental Table 3.

Orthogonal Testing

If a desired PPV cannot be achieved using a single assay, the CDC recommends use of an orthogonal testing algorithm (OTA), a 2-step testing strategy in which all initially positive results are tested with a second

independent serologic test (5). Studies on the effectiveness of this approach are still scarce (34, 75).

OTA TEST SELECTION

Both tests should ideally have high sensitivity (>90%, ideally > 95%). The test with higher specificity should be selected as the first-line test to minimize the number of discordant results while retaining optimal PPV. OTA may use different methods but same antigenic target, same methods and same antigenic target, but different domains or methods that detect antibodies against different antigenic targets (34). OTAs incorporating IgM or IgA serologic tests are not recommended because there is a higher likelihood of discordant results.

The relationship of PPV/NPV and discordant rate in different OTA designs is illustrated in Table 2 with a 2% disease prevalence. If evaluated by a single test (test 1) with specificity of 98%, approximately 50% false positive results would be expected. Adding a sequential test (test 2) with specificity of 95% will, however, result in PPV >90% (Fig. 4). OTA1 and OTA2 represent different designs from test 1 and test 2. Both OTA1 and OTA2 have the same combined PPV of 95%. However, OTA1, which uses the highest specificity test first, results in a lower discordant rate without affecting combined PPV/NPV.

OTA RESULT REPORTING AND INTERPRETATION

If the initial result is negative, the second test is not needed and a negative report is issued; if both tests are positive, a positive report is issued. The interpretive challenge arises when the first result is positive and the second is negative, granting discordant or indeterminate results (online Supplemental Table 4). In this case, OTA results should be interpreted in the context of disease prevalence, sensitivity and specificity of each test, assay methodology, and antigenic targets. If different antigenic targets are used, a discordant result may be attributed to (a) initial false positive, (b) early recovery and/or differences in antibody kinetics, (c) skewed immune response toward one antigen, or (d) waning immunity. To rule out the contribution of differences in antibody kinetics, retesting in 2–4 weeks may be attempted.

Conclusions

Although not recommended as first-line testing for diagnosis of SARS-CoV-2, serologic testing can play important functions in the management of SARS-CoV-2 infection and the pandemic. It is AACC's position that clinical laboratories should only use EUA assays or LDTs that have been developed and properly verified or validated, respectively, in the clinical laboratory. Laboratorians should recognize the utility and limitations of serologic tests and carefully select and implement

EUA or LDT assays and interpret test results. The authors have provided expert opinions and practical recommendations based on the current research on these topics for the clinical laboratory community.

Many studies are underway to gain deeper and broader understanding of SARS-CoV-2, and the tests used to detect and manage the infection will continue to improve. Clinical laboratory professionals, in collaboration with their clinical colleagues, will continue to play an indispensable role in reviewing the evolving scientific literature and adjusting testing strategies to best serve patient and public health needs during this pandemic.

Supplemental Material

Supplemental material is available at *Clinical Chemistry* online.

Nonstandard Abbreviations: COVID-19, coronavirus disease 2019; SARS-CoV-2, severe acute respiratory syndrome coronavirus 2; FDA, US Food and Drug Administration; EUA, emergency use authorization; LDT, laboratory-developed test; S, spike; N, nucleocapsid; RBD, receptor binding domain; nAb, neutralizing antibody; CP, convalescent plasma; RT-PCR, reverse transcriptase PCR; S/CO, signal/cutoff; PPR, positive predictive value; CAP, College of American Pathology; NPV, negative predictive value; OTA, orthogonal testing algorithm.

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