CareStart™
COVID-19 IgM/IgG

Rapid Diagnostic Test for the Detection of SARS-CoV-2 IgM/IgG Ab

Package Insert
(Instructions for Use)

Revision number: D
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Intended Use

The CareStart™ COVID-19 IgM/IgG is an immunochromatographic lateral flow assay intended for the qualitative detection and differentiation of Immunoglobulin M (IgM) and Immunoglobulin G (IgG) antibodies to SARS-CoV-2 in human serum, plasma (sodium citrate, sodium heparin, or dipotassium EDTA), venous whole blood (sodium citrate, sodium heparin, or dipotassium EDTA) and fingerstick whole blood. The CareStart™ COVID-19 IgM/IgG is intended for use as an aid in identifying individuals with an adaptive immune response to SARS-CoV-2, indicating recent or prior infection. At this time, it is unknown for how long antibodies persist following infection and if the presence of antibodies confers protective immunity. The CareStart™ COVID-19 IgM/IgG should not be used to diagnose or exclude acute SARS-CoV-2 infection.

Testing of serum, plasma and venous whole blood specimens is limited to laboratories certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA), 42 U.S.C 263a, that meet requirements to perform high or moderate complexity tests.

Testing of fingerstick whole blood specimens is limited to laboratories certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA), 42 U.S.C. §263a, that meet the requirements to perform high, moderate or waived complexity tests. Testing of fingerstick whole blood specimens is authorized for use at the Point of Care (POC), i.e., in patient care settings operating under a CLIA Certificate of Waiver, Certificate of Compliance, or Certificate of Accreditation.

Results are for the detection of SARS-CoV-2 antibodies. IgM and IgG antibodies to SARS-CoV-2 are generally detectable in the blood several days after initial infection, although the duration of time antibodies are present post-infection is not well characterized. Individuals may have detectable virus present for several weeks following seroconversion.

Laboratories within the United States and its territories are required to report all results to the appropriate public health authorities.

The sensitivity of CareStart™ COVID-19 IgM/IgG early after infection is unknown. Negative results do not preclude acute SARS-CoV-2 infection. If acute infection is suspected, direct testing for SARS-CoV-2 is necessary.

False-positive results for CareStart™ COVID-19 IgM/IgG may occur due to cross-reactivity from pre-existing antibodies or other possible causes. Due to the risk of false-positive results, confirmation of positive results should be considered using a second, different SARS-CoV-2 IgG or IgM assay.
The CareStart™ COVID-19 IgM/IgG is only for use under the Food and Drug Administration’s Emergency Use Authorization.

Summary and Explanation of the Test

Coronavirus disease 2019 (COVID-19) is a respiratory illness that can spread from person to person. This antibody test is intended for use as an aid in identifying individuals with an adaptive immune response to SARS-CoV-2, the virus that causes COVID-19, indicating recent or prior infection, by detecting antibodies to SARS-CoV-2 in human blood specimens. Although not everyone infected will develop an antibody response, appropriately validated serology tests, when used broadly, can be useful in understanding how many people have developed an adaptive immune response to the virus and how far the pandemic has progressed. At this time, it is unknown for how long antibodies persist following infection and if the presence of antibodies confers protective immunity.

Principles of the Test

The CareStart™ COVID-19 IgM/IgG test is an immunochromatographic assay for the detection and differentiation of SARS-CoV-2 IgM and/or IgG antibodies in human blood specimens. Control antibody, anti-human IgG, and streptavidin (test line for IgM) are immobilized onto a nitrocellulose membrane to form three distinct lines, the control line, the IgG test line, and the IgM test line. The nitrocellulose membrane is attached onto a plastic backing card and combined with the other reagents and pads to construct a test strip. The test strip is encased inside a plastic device. Blood samples, including human serum, plasma (sodium citrate, sodium heparin, or dipotassium EDTA), venous whole blood (sodium citrate, sodium heparin, or dipotassium EDTA), and fingerstick whole blood, are added to the sample well of the test device to initiate a test. The sample specimens migrate sequentially through filter pad, conjugate pad, nitrocellulose membrane, and absorbent pad. SARS-CoV-2 antibodies in sample specimens interact with the recombinant SARS-CoV-2 antigen (SARS-CoV-2 nucleocapsid and spike protein S1 RBD) that is conjugated to colloidal gold nanobeads and biotin-conjugated anti-human antibodies to form an immune complex while it migrates through the conjugate pad. SARS-CoV-2 IgM antibodies react with the gold-conjugated SARS-CoV-2 antigen and biotin-conjugated anti-human IgM. SARS-CoV-2 IgG antibodies only react with the gold-conjugated SARS-CoV-2 antigen. The immune complexes migrate through the nitrocellulose membrane and bind to each respective test line. The IgM immune complexes bind to the streptavidin region (IgM test line, “M”) on the membrane to generate a purple-colored line to indicate a positive IgM result. The IgG immune complexes bind to the anti-human IgG region (IgG test line, “G”) on the membrane to generate a purple-colored line to indicate a positive IgG result. The gold-conjugated chicken IgY migrates through the membrane and binds to the control antibody (anti-chicken IgY) in the control region to generate a red-colored line (control line, “C”).
The test results should be interpreted 10 minutes after the addition of buffer to the sample well. The test results should not be interpreted after 15 minutes. The color intensity in the test region will vary. Any faint-colored line(s) in the test region(s) should be considered as positive.

The presence of two lines marked by “C” and “G” indicates a SARS-CoV-2 IgG positive result. The presence of two lines marked by “C” and “M”, indicates a SARS-CoV-2 IgM positive result. The presence of three lines “C”, “G,” and “M”, indicates positive results for both SARS-CoV-2 IgG and IgM. The appearance of only the control line “C” indicates negative. If the control line does not appear, regardless of the presence of “G” or “M” test lines, the test result is not valid. With an invalid result, it is recommended to repeat the test using a new, unopened device following the instructions.

Reagents and Materials Provided

<table>
<thead>
<tr>
<th>Contents Name</th>
<th>Quantity (in a kit)</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Test device</td>
<td>25 each</td>
<td>Foil pouched test device containing one test strip which is encased on plastic device cassette.</td>
</tr>
<tr>
<td>Assay buffer</td>
<td>1 each</td>
<td>Na₂CO₃ &lt; 0.1% sodium azide as a preservative.</td>
</tr>
<tr>
<td>Blood transfer pipette</td>
<td>25 each</td>
<td>For blood transfer.</td>
</tr>
<tr>
<td>Alcohol swab</td>
<td>25 each</td>
<td>Use to clean a collection site prior to sampling the fingerstick whole blood.</td>
</tr>
<tr>
<td>Sterile safety lancet</td>
<td>25 each</td>
<td>Single-use lancets intended for sampling fingerstick whole blood.</td>
</tr>
<tr>
<td>Package insert</td>
<td>1 each</td>
<td>Instructions for use</td>
</tr>
<tr>
<td>Quick Reference Instructions (QRI)</td>
<td>1 each</td>
<td>Quick reference instructions</td>
</tr>
</tbody>
</table>

*Materials required but not supplied
- Timer
- Pair of gloves

Optional materials:
- 20 µl micropipette

Warnings and Precautions

- For prescription and in vitro diagnostic use only. For use under an Emergency Use Authorization Only.
- This product has not been FDA cleared or approved; but has been authorized for emergency use by FDA under an EUA for use by authorized laboratories.
- This product has been authorized only for detecting the presence of IgM and IgG antibodies to SARS-CoV-2, not for any other viruses or pathogens.
- The emergency use of this product is only authorized for the duration of the declaration that circumstances exist justifying the authorization of emergency use of in vitro diagnostics for detection and/or diagnosis of COVID-19 under Section 564(b)(1) of the Federal Food, Drug and Cosmetic Act, 21 U.S.C. § 360bbb-3(b)(1), unless the declaration is terminated or authorization is revoked sooner.
• Testing of serum, plasma, and venous whole blood specimens is limited to laboratories certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA), 42 U.S.C. 263a, that meet requirements to perform high or moderate complexity tests.
• Testing of fingerstick whole blood specimens is limited to laboratories certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA), 42 U.S.C. §263a, that meet the requirements to perform high, moderate or waived complexity tests. Testing of fingerstick whole blood specimens is authorized for use at the Point of Care (POC), i.e., in patient care settings operating under a CLIA Certificate of Waiver, Certificate of Compliance, or Certificate of Accreditation.
• As with all diagnostic tests, all results must be interpreted together with other clinical information available to the physician.
• Immediately use after opening the test device in the pouch.
• Immediately add the assay buffer to the test device after the specimen is applied.
• In order to obtain accurate results, the test must follow this package insert.
• Do not interpret the test result before 10 minutes and after 15 minutes following the addition of buffer to the sample well.
• Do not use if the test device package is damaged.
• Do not use the kit contents beyond the expiration date.
• Do not eat, drink, or smoke in the area where the specimens and kit contents are handled.
• Use appropriate precautions in the collection, handling, storage, and disposal of patient samples and used kit contents.
• Dispose of used contents as biohazardous wastes in accordance with federal, state, and local requirements.
• Nitrile or latex gloves should be worn when performing this test.
• If the assay buffer contacts the skin or eye, flush with copious amounts of water.
• Handle all specimens as though they contain infectious agents.
• Adding additional blood sample volume to the sample well may cause false positive or invalid results.
• Observe established precautions against microbiological hazards throughout the procedure and follow the standard procedures for proper disposal of specimens.
• Reagents contain sodium azide, which is harmful if inhaled, swallowed, or exposed to skin. Contact with acids produces a very toxic gas. If there is contact with skin, wash immediately with plenty of water. Sodium azide may react with lead and copper plumbing to form highly explosive metal azides. On disposal, flush with a large volume of water to prevent azide build-up.
• Do not interchange kit contents from different lots.
• Do not re-use any contents in the kit as they are single-use only.

Storage and Stability

• Store the test kit as packaged between 1 ~ 30°C.
• The reagents and materials in the CareStart™ COVID-19 IgM/IgG are stable until the expiration date printed on the outer packaging. Do not use beyond the expiration date.
• The test device must remain in the sealed pouch until use.
• Do not freeze any contents of the kit.
Quality Control

**Internal Quality Control:** The CareStart™ COVID-19 IgM/IgG contains a built-in internal procedural control in the test device. A red-colored line appearing in the control region “C” is designed as an internal procedural control. The appearance of the procedural control line indicates that sufficient flow has occurred, and the functional integrity of the test device has been maintained. If the procedural control line does not develop in 10 minutes, the test result is considered invalid, and retesting with a new device is recommended. If the internal procedural control line is still absent in the retest, please contact the manufacturer or distributor.

**External Control:** It is recommended to follow the laboratory regulations or quality control procedures to perform external controls in CareStart™ COVID-19 IgM/IgG. Controls are available through Access Bio under catalog number: SCLM-02571 or SCLM-10071.

**NOTE:** The external controls are available for separate purchase.

- Positive External Control: Mixture of human chimeric SARS-CoV-2 IgM and IgG spike S1 antibodies in heat-inactivated SARS-CoV-2 antibody-negative confirmed serum.
- Negative External control: Heat inactivated SARS-CoV-2 antibody-negative confirmed serum.

**Specimen Type**

Acceptable specimen types for testing with the CareStart™ COVID-19 IgM/IgG are human serum, plasma (sodium citrate, sodium heparin, or dipotassium EDTA), venous whole blood (sodium citrate, sodium heparin, or dipotassium EDTA), and fingerstick whole blood. Proper specimen collection methods must be followed. Inadequate specimen collection and/or improper specimen handling may yield false results; therefore, specimen collection requires specific training and guidance due to the importance of specimen quality to obtain accurate test results.

**Specimen Collection and Handling Procedures**

**Procedural Notes**

1. The CareStart™ COVID-19 IgM/IgG can be performed using human serum, plasma (sodium citrate, sodium heparin, or dipotassium EDTA), venous whole blood (sodium citrate, sodium heparin, or dipotassium EDTA), and fingerstick whole blood.
2. Allow test devices, reagents, and specimens to equilibrate up to room temperature (15~30°C) prior to testing.
3. Remove the CareStart™ COVID-19 IgM/IgG test device from its foil pouch immediately before testing.

Fingerstick whole blood
1. Collect the specimen wearing safety gloves to avoid contact and contamination.
2. Use only the provided blood lancet, alcohol swab, and blood transfer pipette for human fingerstick whole blood specimen collection.
3. Process the fingerstick sample immediately after collection.
4. Testing should be performed immediately after specimen collection.

1. Clean the fingertip to be pierced with an alcohol swab.

2. Squeeze the end of the fingertip and pierce the cleaned area using a blood lancet. Properly discard the blood lancet.

3. Press the top part of the provided blood transfer pipette.

4. Touch the blood using the pipette tip while still pressing the pipette.
5. Fill the pipette with the blood sample up to the blue marked line by releasing slowly.

**Venous Whole Blood:**

Draw venous whole blood following the general laboratory procedures by a trained operator. Collect the blood sample in a commercially available blood collection tube containing anticoagulants including sodium citrate, sodium heparin, or dipotassium EDTA. Swirl the tube gently as needed. It is recommended to test whole blood specimens immediately after blood collection.

**Serum:**

Collect venous whole blood into a container NOT containing anticoagulants. Wait for the blood clot and separate the serum by centrifugation.

**Plasma:**

Collect venous whole blood into a container containing anticoagulants (sodium citrate, sodium heparin, or dipotassium EDTA). Separate the plasma by centrifugation.

- Use only the provided blood transfer pipette or micropipette for sample loading to the test device.
- Separate serum or plasma from blood as soon as possible to avoid hemolysis. Use only clear, non-hemolyzed specimens.
- Testing should be performed immediately after specimen collection. Do not leave the serum and plasma specimens at room temperature beyond 8 hours. Serum and plasma specimens may be stored at 2-8°C for up to 48 hours. For long term storage, serum and plasma specimens should be kept below -20°C for up to one month. It is recommended to test whole blood specimens immediately after blood collection.
- Bring specimens to room temperature prior to testing. Frozen specimens must be completely thawed and mixed well prior to testing. Specimens cannot be frozen and thawed more than once.
• If specimens are to be shipped, they should be packed in compliance with local regulations covering the transportation of etiologic agents.
1. Place a device on a clean, flat surface after removing it from the pouch. Write the patient's ID on the device if required.

2. Transfer serum, plasma sample, venous whole blood, or fingerstick whole blood:
   a) using a provided blood transfer pipette:
   Press the top part of the provided blood transfer pipette and touch the sample by the pipette tip while pressing the pipette. Releasing the press slowly to fill the pipette with sample up to the blue marked line (approximately 10 µl). Add the blood sample to the sample well “S” of the test device by pressing the top part of the blood transfer pipette.

   ![Image showing blood transfer pipette and sample well]

   **NOTE:** Excessive blood specimen may cause false positive or invalid test results.

   b) using a micropipette:
   Transfer 10 µl of the venous whole blood, serum, or plasma sample to the sample well “S” of the test device using a micropipette.

3. Open the cap and invert the assay buffer bottle and hold vertically above the sample well. Squeeze the bottle gently to add one (1) drop of the assay buffer solution to the sample well “S” immediately after sample loading.

4. Start a timer. Read the result at 10 minutes. The test results should not be read earlier than 10 minutes. Test results should not be read after 15 minutes.
Interpretation of Results

**NOTE:** The test results should be read and interpreted not earlier than 10 minutes after the sample application and the reading and interpretation of the results should not exceed 15 minutes. The test results should not be interpreted using any instruments.

**IgM Positive:** Two distinct lines appear.

One red-colored line next to “C” and one purple-colored line next to “M” indicates a SARS-CoV-2 IgM positive result.

**IgG Positive:** Two distinct lines appear.

One red-colored line next to “C” and one purple-colored line next to “G” indicates a SARS-CoV-2 IgG positive result.

**IgM/IgG Positive:** Three distinct lines appear.

One red-colored line next to “C”, one purple-colored line next to “M”, and one purple-colored line next to “G” indicates a SARS-CoV-2 IgM and IgG positive result.

**Result with faint colored line(s):**

The color intensity in the test region will vary. Any faint colored line(s) in the test region(s) should be considered as positive.

**Negative:**

Only one line next to “C” indicates a negative result.
**Invalid:** No control line appears.

If the control line “C” is not visible, the result is invalid. Re-run the test using a new test device. If the same invalid result persists, contact the manufacturer or distributor before continuing to test samples.

**Expected Results of the External Controls**

**Positive Control:** three distinct lines appear.

One red-colored line next to “C”, one purple-colored line next to “M”, and one purple-colored line next to “G” indicates SARS-CoV-2 IgM and IgG positive result.

**Negative Control:** one distinct line appears.

Only one red-colored line next to “C” indicates a negative result.

**NOTE:** If the test result for either the Negative Control or the Positive Control is not as expected, the test should be repeated using a new Test Device. If the test result for any of the controls is not as expected upon retesting, contact Technical Support.

**Limitations**

1. Use of the CareStart™ COVID-19 IgM/IgG is limited to laboratory personnel who have been trained. Not for home use.

2. The test is limited to the qualitative detection of anti-COVID-19 antibody levels in human serum and ACD plasma samples and does not indicate the quantity of the antibodies. The intensity of the test line does not necessarily correlate to the SARS-CoV-2 antibody titer in the specimen.

3. The test results should be interpreted 10 minutes after starting the test. The test results should not be interpreted after 15 minutes.

4. This test can only be used for the analysis of human serum, plasma (sodium citrate, sodium heparin, or dipotassium EDTA), venous whole blood (sodium citrate, sodium heparin, or dipotassium EDTA), and fingerstick whole blood samples.

5. Negative results do not preclude SARS-CoV2 infection and should not be used as the sole basis for patient management decisions. IgM antibodies may not be detected in the first several days of infection; the sensitivity of the Rapid COVID-19 IgM/IgG Combo Test Kit early
after an infection is unknown. False-positive results for IgM and IgG antibodies may occur due to cross-reactivity from pre-existing antibodies or other possible causes.

6. A negative result can occur if the quantity of antibodies for the SARS-CoV-2 virus present in the specimen is below the detection limit of the assay, or if the virus has undergone minor amino acid mutation(s) in the epitope recognized by the antibody used in the test.

7. The test may have lower sensitivity for IgG and IgM detection in symptomatic individuals prior to 15 days since symptom onset.

8. Direct testing with a molecular diagnostic test should be performed to evaluate for acute SARS-CoV-2 infection in symptomatic individuals.

9. Results from antibody testing should not be used to diagnose or exclude acute SARS-CoV-2 infection or to determine infection status.

10. It is unknown for how long antibodies persist following infection and if the presence of antibodies confers protective immunity.

11. Positive results may not indicate previous SARS-CoV-2 infection. Consider other information including clinical history and local disease prevalence, in assessing the need for an alternative serology test to confirm an adaptive immune response. Positive results may be due to past or present infection with non-SARS-CoV-2 coronavirus strains, such as coronavirus HKU1, NL63, OC43, or 229E.

12. This test will only indicate the presence of SARS-CoV-2 IgM and/or IgG antibodies in the specimen.

13. The detection of SARS-CoV-2 IgM/IgG antibodies is dependent upon proper specimen collection, handling, storage, and preparation. Failure to observe proper procedures in any one of these steps can lead to incorrect results.

14. This device has been evaluated for use with human specimen material only.

15. This test cannot rule out diseases caused by other bacterial or viral pathogens.

16. This device should not be used for the screening of donated blood.

17. The performance of this device has not been established in individuals that have received a COVID-19 vaccine. The clinical significance of a positive or negative antibody result following COVID-19 vaccination has not been established, and the result from this assay should not be interpreted as an indication or degree of protection from infection after vaccination.

18. The performance of this test was established based on the evaluation of a limited number of clinical specimens (serum and plasma) collected from March 27, 2016 to May 21, 2020 from three sites in the US (California and Illinois). The clinical performance characteristics of the CareStart™ COVID-19 IgM/IgG test for POC testing was evaluated in a multi-site prospective study in the U.S (California and New Mexico). The clinical performance has not been established in all circulating variants but is anticipated to be reflective of the prevalent variants in circulation at the time and location of the clinical evaluation. Performance at the time of testing may vary depending on the variants circulating, including newly emerging strains of SARS-CoV-2 and their prevalence, which change over time.
Conditions of Authorization for Laboratories

The CareStart™ COVID-19 IgM/IgG Letter of Authorization, along with the authorized Fact Sheet for Healthcare Providers, the authorized Fact Sheet for Recipients, and other authorized labeling are available on the FDA website:


Authorized Laboratories using the CareStart™ COVID-19 IgM/IgG must adhere to the Conditions of Authorization indicated in the Letter of Authorization are listed below:

1. Authorized laboratories using the CareStart™ COVID-19 IgM/IgG must include with test result reports, all authorized Fact Sheets. Under exigent circumstances, other appropriate methods for disseminating these Fact Sheets may be used, which may include mass media.
2. Authorized laboratories must use the CareStart™ COVID-19 IgM/IgG as outlined in the authorized labeling. Deviations from the authorized procedures, including the authorized clinical specimen types, authorized control materials, authorized other ancillary reagents and authorized materials required to use the CareStart™ COVID-19 IgM/IgG are not permitted.
3. Authorized laboratories that receive the CareStart™ COVID-19 IgM/IgG must notify the relevant public health authorities of their intent to run the CareStart™ COVID-19 IgM/IgG prior to initiating testing.
4. Authorized laboratories using the CareStart™ COVID-19 IgM/IgG must have a process in place for reporting test results to healthcare providers and relevant public health authorities, as appropriate.
5. Authorized laboratories must collect information on the performance of the CareStart™ COVID-19 IgM/IgG and report to Division of Microbiology Devices (DMD)/Office of Health Technology7 (OHT7)-(Office of In Vitro Diagnostics and Radiological Health (OIR)/Office of Product Evaluation and Quality (OPEQ)/Center for Devices and Radiological Health (CDRH) (via email: CDRH-EUA-Reporting@fda.hhs.gov) and Access Bio, Inc. Technical Support (via email: TShelp@accessbio.net) any suspected occurrence of false positive or false negative results and significant deviations from the established performance characteristics of the assay of which they become aware.
6. All laboratory personnel using the CareStart™ COVID-19 IgM/IgG must be appropriately trained in immunochromatographic techniques and use appropriate laboratory and personal protective equipment when handling this kit, and use the CareStart™ COVID-19 IgM/IgG in accordance with the authorized labeling. All laboratory personnel using the assay must also be trained in and be familiar with the interpretation of results of the CareStart™ COVID-19 IgM/IgG.
7. Access Bio, Inc., authorized distributors, and authorized laboratories using the CareStart™ COVID-19 IgM/IgG must ensure that any records associated with this EUA are maintained
until otherwise notified by FDA. Such records will be made available to FDA for inspection upon request.

- The letter of authorization refers to “authorized laboratories” as the following:
  Testing of serum, plasma and venous whole blood specimens is limited to laboratories certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA), 42 U.S.C 263a, that meet requirements to perform high or moderate complexity tests.
  Testing of fingerstick whole blood specimens is limited to laboratories certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA), 42 U.S.C. §263a, that meet the requirements to perform high, moderate or waived complexity tests. Testing of fingerstick whole blood specimens is authorized for use at the Point of Care (POC), i.e., in patient care settings operating under a CLIA Certificate of Waiver, Certificate of Compliance, or Certificate of Accreditation.

Performance Characteristics

Clinical Agreement

Study I: Retrospective Study – Serum and Plasma

The clinical performance of CareStart™ COVID-19 IgM/IgG was evaluated using retrospectively collected SARS-CoV-2 serum and plasma samples at 3 sites by 10 operators in the U.S. A total of 246 samples, 211 plasma (47 positives and 164 pre-COVID) and 35 serum (17 positives and 18 negatives), collected from the U.S were tested in this study. A total of 164 pre-COVID samples were collected before December 2019 in the U.S. All the collected positive serum and plasma samples were confirmed by FDA authorized SARS-CoV-2 RT-PCR tests as comparators.

All the negative and positive samples were tested in a blinded fashion. Each sample was assigned with a unique subject identification code during collection and randomized prior to the testing. The expected results of the samples were completely blinded to the operators in this study. All the samples were tested according to the CareStart™ COVID-19 IgM/IgG testing procedures.

A total of 246 samples were considered evaluable in this study.

CareStart™ COVID-19 IgM/IgG Performance against the Comparator Methods – Serum and Plasma

For IgG antibody detection, the positive percent agreement (PPA) of CareStart™ COVID-19 IgM/IgG was 96.88% (62/64) (95% CI of 89.30 – 99.13%). For IgM antibody detection, the PPA was 89.06% (57/64) (95% CI of 79.10 - 94.60%). The overall NPA (either IgG positive or IgM positive counted as positive) was 98.90% (180/182) (95% CI 96.08 – 99.70%).

IgG results stratified by days post-onset of symptoms – serum and plasma
**IgM results stratified by days post-onset of symptoms – serum and plasma**

<table>
<thead>
<tr>
<th>Days from Symptom Onset</th>
<th>Total number of samples</th>
<th>Non-reactive</th>
<th>Reactive</th>
<th>PPA</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 - 7</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>8 - 14</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>100% (1/1)</td>
<td>20.65 – 100%</td>
</tr>
<tr>
<td>≥ 15</td>
<td>62</td>
<td>2</td>
<td>60</td>
<td>96.77% (60/62)</td>
<td>88.98 – 99.11%</td>
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<tr>
<td>Unknown</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>100% (1/1)</td>
<td>20.65 - 100%</td>
</tr>
</tbody>
</table>

**Study II: Independent Clinical Agreement Validation**

The CareStart™ COVID-19 IgM/IgG from Access Bio was tested on Jun 2, 2020, at the Frederick National Laboratory for Cancer Research (FNLCR) sponsored by the National Cancer Institute (NCI). The test was validated against a panel of previously frozen samples consisting of 30 SARS-CoV-2 antibody-positive serum samples and 80 antibody-negative serum and plasma samples. Each of the 30 antibody-positive samples was confirmed with a nucleic acid amplification test (NAAT) and both IgM and IgG antibodies were confirmed to be present in all 30 samples. The presence of antibodies in the samples was confirmed by several orthogonal methods prior to testing with the CareStart™ COVID-19 IgM/IgG. The presence of IgM and IgG antibodies specifically was confirmed by one or more comparator methods. Antibody-positive samples were selected at different antibody titers.

All antibody-negative samples were collected prior to 2020 and include: i) Seventy (70) samples selected without regard to clinical status, “Negatives” and ii) Ten (10) samples selected from banked serum from HIV+ patients, “HIV+”. Testing was performed by one operator using one lot of the CareStart™ COVID-19 IgM/IgG. Confidence intervals for sensitivity and specificity were calculated per a score method described in CLSI EP12-A2 (2008).

For the evaluation of cross-reactivity with HIV+, it was determined whether an increased false-positive rate among antibody-negative samples with HIV was statistically higher than the false positive rate among antibody-negative samples without HIV (for this, a confidence interval for the difference in false-positive rates was calculated per a score method described by Altman). The results and data analysis are shown in the tables below:
### CareStart™ COVID-19 IgM/IgG

Rapid Diagnostic Test for Detection of SARS-CoV-2 IgM/IgG Ab

**Comparator Method**

<table>
<thead>
<tr>
<th>Antibody Positive</th>
<th>Collected pre-2020</th>
<th>Antibody Negative</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>CareStart™ COVID-19 IgM/IgG</strong></td>
<td><strong>IgM+, IgG+</strong></td>
<td><strong>IgM+, IgG-</strong></td>
</tr>
<tr>
<td>IgM+, IgG+</td>
<td>27</td>
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<tr>
<td>IgM+, IgG-</td>
<td></td>
<td>1</td>
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<td>IgM-, IgG+</td>
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<tr>
<td><strong>Total</strong></td>
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<table>
<thead>
<tr>
<th>Measure</th>
<th>Estimate</th>
<th>Confidence Interval</th>
</tr>
</thead>
<tbody>
<tr>
<td>IgM Sensitivity</td>
<td>90.0% (27/30)</td>
<td>74.4%; 96.5%</td>
</tr>
<tr>
<td>IgM Specificity</td>
<td>98.8% (79/80)</td>
<td>93.3%; 99.8%</td>
</tr>
<tr>
<td>IgG Sensitivity</td>
<td>100% (30/30)</td>
<td>88.7%; 100%</td>
</tr>
<tr>
<td>IgG Specificity</td>
<td>98.8% (79/80)</td>
<td>93.3%; 99.8%</td>
</tr>
<tr>
<td>Combined Sensitivity</td>
<td>100% (30/30)</td>
<td>88.7%; 100%</td>
</tr>
<tr>
<td>Combined Specificity</td>
<td>97.5% (78/80)</td>
<td>91.3%; 99.3%</td>
</tr>
<tr>
<td>Combined PPV for prevalence = 5.0%</td>
<td>67.8%</td>
<td>35%; 88.4%</td>
</tr>
<tr>
<td>Combined NPV for prevalence = 5.0%</td>
<td>100%</td>
<td>99.4%; 100%</td>
</tr>
<tr>
<td>Cross-reactivity with HIV+</td>
<td>0.0% (0/10), not detected</td>
<td>-</td>
</tr>
</tbody>
</table>

Important limitations of the study:

1. Samples were not randomly selected, and sensitivity and specificity estimates may not be indicative of the real-world performance of the device.
2. These results are based on serum and plasma samples only and may not be indicative of performance with other sample types, such as whole blood, including finger stick blood.
3. Information about anticoagulants used is not known.
4. The number of samples in the panel is a minimally viable sample size that still provides reasonable estimates and confidence intervals for test performance, and the samples used may not be representative of the antibody profile observed in patient populations.

**Study III: Prospective Fingerstick Study (POC)**

The clinical performance characteristics of the CareStart™ COVID-19 IgM/IgG test was evaluated in a multi-site prospective study in the U.S. against an FDA authorized RT-PCR molecular assay as a comparator method. A total of two (2) Point-of-Care investigational sites in the U.S. (California and New Mexico) participated in the study. To be enrolled in the study, the positive patients had to be confirmed as positive by an FDA authorized RT-PCR and exhibit signs/symptoms of the COVID-19-like illness within 5 to 28 days prior to testing with the devices.
under evaluation. The negative subjects had to be confirmed as negative by an FDA authorized RT-PCR and currently present no signs or onset of symptoms and no previous infection history.

Testing was performed by eight (8) operators with no laboratory experience and who were representative of the intended users. Operators only used the QRI for the test without any training provided and completed an ease-of-use questionnaire indicating favorable responses to all questions asked after completing the study.

The clinical study tested 75 samples (32 were collected from RT-PCR positive subjects and 43 were collected from RT-PCR negative subjects). Of the 32 RT-PCR positive clinical samples, date of symptom onset was known for 26 samples and unknown for 6 samples. Results were assessed stratified by (A) the days post symptom onset and (B) the days post RT-PCR testing in comparison to multiple FDA authorized assays as comparators:

IgG and IgM PPA results stratified by days post symptoms onset – fingerstick

<table>
<thead>
<tr>
<th>Days post symptom onset</th>
<th>No. of RT-PCR Positive</th>
<th>IgG Reactive</th>
<th>IgG PPA</th>
<th>95% CI</th>
<th>IgM Reactive</th>
<th>IgM PPA</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 – 7 days</td>
<td>1</td>
<td>1</td>
<td>100%</td>
<td>20.65 – 100%</td>
<td>1</td>
<td>100%</td>
<td>20.65 – 100%</td>
</tr>
<tr>
<td>8 – 14 days</td>
<td>12</td>
<td>12</td>
<td>100%</td>
<td>75.76 – 100%</td>
<td>12</td>
<td>100%</td>
<td>75.76 – 100%</td>
</tr>
<tr>
<td>≥15 days</td>
<td>13</td>
<td>13</td>
<td>100%</td>
<td>77.19 – 100%</td>
<td>13</td>
<td>100%</td>
<td>77.19 – 100%</td>
</tr>
<tr>
<td>Total</td>
<td>26</td>
<td>26</td>
<td>100%</td>
<td>87.13 – 100%</td>
<td>26</td>
<td>100%</td>
<td>87.13 – 100%</td>
</tr>
</tbody>
</table>

IgG and IgM PPA results stratified by days post RT-PCR result – fingerstick

<table>
<thead>
<tr>
<th>Days post RT-PCR positive</th>
<th>No. of RT-PCR Positive</th>
<th>IgG Reactive</th>
<th>IgG PPA</th>
<th>95% CI</th>
<th>IgM Reactive</th>
<th>IgM PPA</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>0 – 7 days</td>
<td>11</td>
<td>11</td>
<td>100%</td>
<td>74.12 – 100%</td>
<td>11</td>
<td>100%</td>
<td>74.12 – 100%</td>
</tr>
<tr>
<td>8 – 14 days</td>
<td>11</td>
<td>11</td>
<td>100%</td>
<td>74.12 – 100%</td>
<td>11</td>
<td>100%</td>
<td>74.12 – 100%</td>
</tr>
<tr>
<td>≥15 days</td>
<td>10</td>
<td>10</td>
<td>100%</td>
<td>72.25 – 100%</td>
<td>10</td>
<td>100%</td>
<td>72.25 – 100%</td>
</tr>
<tr>
<td>Total</td>
<td>32</td>
<td>32</td>
<td>100%</td>
<td>89.28 – 100%</td>
<td>32</td>
<td>100%</td>
<td>89.28 – 100%</td>
</tr>
</tbody>
</table>

The overall NPA of the CareStart COVID-19 IgG/IgM Rapid Test Device for IgG/IgM in fingerstick whole blood samples is 100% (43/43) (95% CI 91.80 – 100.0%).

Cross-Reactivity (Exclusivity)

The cross-reactivity study was conducted by testing a total of 87 plasma or serum samples, including 14 non-SARS-CoV-2 pathogens and one (1) autoantibody. All the plasma and serum
samples tested as negative showed no cross-reactivity and resulted in 100% agreement between CareStart™ COVID-19 IgM/IgG test result and the expected result as presented in the table below:

<table>
<thead>
<tr>
<th>Samples</th>
<th>Sample number and type</th>
<th>Test results (# of positive / # of replicate)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>IgM Results</td>
</tr>
<tr>
<td>Anti-Influenza A</td>
<td>5 plasma</td>
<td>0/5</td>
</tr>
<tr>
<td>Anti-Influenza B</td>
<td>5 plasma</td>
<td>0/5</td>
</tr>
<tr>
<td>Anti-HCV</td>
<td>8 plasma, 2 serum</td>
<td>0/10</td>
</tr>
<tr>
<td>Anti-HBV</td>
<td>5 serum</td>
<td>0/5</td>
</tr>
<tr>
<td>Anti-229E (alpha coronavirus)</td>
<td>3 plasma</td>
<td>0/3</td>
</tr>
<tr>
<td>Anti-NL63 (alpha coronavirus)</td>
<td>7 plasma</td>
<td>0/7</td>
</tr>
<tr>
<td>Anti-OC43 (beta coronavirus)</td>
<td>9 plasma</td>
<td>0/9</td>
</tr>
<tr>
<td>Anti-HKU1 (beta coronavirus)</td>
<td>5 plasma</td>
<td>0/5</td>
</tr>
<tr>
<td>Antinuclear antibodies (ANA)</td>
<td>5 serum</td>
<td>0/5</td>
</tr>
<tr>
<td>Anti-respiratory syncytial virus</td>
<td>1 plasma, 5 serum</td>
<td>0/6</td>
</tr>
<tr>
<td>Anti-HIV</td>
<td>10 plasma</td>
<td>0/10</td>
</tr>
<tr>
<td>Anti-Dengue virus</td>
<td>5 plasma</td>
<td>0/5</td>
</tr>
<tr>
<td>Anti-T. pallidum (Syphilis)</td>
<td>5 serum</td>
<td>0/5</td>
</tr>
<tr>
<td>Anti-Rhinovirus</td>
<td>2 plasma</td>
<td>0/2</td>
</tr>
<tr>
<td>Anti-Hepatitis B</td>
<td>5 plasma</td>
<td>0/5</td>
</tr>
</tbody>
</table>

**Interfering Substances**

To assess substances with the potential to interfere with the performance of the CareStart™ COVID-19 IgM/IgG, SARS-CoV-2 IgM, IgG positive, and negative samples were tested with the addition of potentially interfering substances. The CareStart™ COVID-19 IgM/IgG test performance was not affected by any of the eight potentially interfering substances tested.

- Acetaminophen
- Acetylsalicylic acid
- Albendazole
- Chloroquine diphosphate
- HAMA
- Hemoglobin
- Ibuprofen
- Rifampicin

The interfering effects of biotin concentrations ranging between 10 ng/ml and 100 µg/ml were tested in a separate study. Biotin concentrations up to 2.5 µg/ml did not lead to false results. Biotin concentrations >5 µg/ml can cause false-negative IgM results with the CareStart™ COVID-19 IgM/IgG. None of the IgG positive samples tested produced false negative in all biotin concentrations tested.

**Class Specificity**
The CareStart™ COVID-19 IgM/IgG was evaluated to determine that the assay accurately detects each SARS-CoV-2 IgM and IgG antibody class on its corresponding test lines. A total of five (5) IgM and IgG positive serum samples were treated with DTT to determine the class specificity of the test. All samples treated with DTT showed no visible IgM line with the CareStart™ COVID-19 IgM/IgG, whereas the IgG results were not affected by DTT treatment. IgM and IgG results after DTT treatment showed 100% agreement to the expected results.

Matrix Equivalency

The matrix equivalency study was performed by spiking SARS-CoV-2 IgM/IgG positive sample into negative sample matrices for serum, venous whole blood, and plasma using different anticoagulants. All testing matrices were collected from the same donor and a total of five donors were evaluated in CareStart™ COVID-19 IgM/IgG. The venous whole blood samples were collected from each individual in four different containers to prepare serum (no anticoagulant), and three matrices each of venous whole blood (sodium citrate, sodium heparin, and dipotassium EDTA) and plasma (sodium citrate, sodium heparin, and dipotassium EDTA). To prepare positive sample panels, each sample matrix was spiked with SARS-CoV-2 IgM/IgG positive serum sample at low positive and moderate positive levels and randomized for testing. The samples were tested in duplicate with the CareStart™ COVID-19 IgM/IgG. All the test results of seven different matrices from the individual showed 100% agreement to the expected results.

<table>
<thead>
<tr>
<th>Sample type</th>
<th>Test results (# of positive / # of replicate (% agreement))</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>LP results</td>
</tr>
<tr>
<td></td>
<td>IgM</td>
</tr>
<tr>
<td>Venous whole blood</td>
<td></td>
</tr>
<tr>
<td>(EDTA)</td>
<td>10/10</td>
</tr>
<tr>
<td></td>
<td>(100%)</td>
</tr>
<tr>
<td>Venous whole blood</td>
<td></td>
</tr>
<tr>
<td>(Heparin)</td>
<td>10/10</td>
</tr>
<tr>
<td></td>
<td>(100%)</td>
</tr>
<tr>
<td>Venous whole blood</td>
<td></td>
</tr>
<tr>
<td>(Citrate)</td>
<td>10/10</td>
</tr>
<tr>
<td></td>
<td>(100%)</td>
</tr>
<tr>
<td>Plasma</td>
<td></td>
</tr>
<tr>
<td>(EDTA)</td>
<td>10/10</td>
</tr>
<tr>
<td></td>
<td>(100%)</td>
</tr>
<tr>
<td>Plasma</td>
<td></td>
</tr>
<tr>
<td>(Heparin)</td>
<td>10/10</td>
</tr>
<tr>
<td></td>
<td>(100%)</td>
</tr>
<tr>
<td>Plasma</td>
<td></td>
</tr>
<tr>
<td>(Citrate)</td>
<td>10/10</td>
</tr>
<tr>
<td></td>
<td>(100%)</td>
</tr>
<tr>
<td>Serum</td>
<td>10/10</td>
</tr>
<tr>
<td></td>
<td>(100%)</td>
</tr>
</tbody>
</table>

Technical Support
For questions, or to report a problem, please call Technical Support at +1-888-898-1270 (Available Hours: Mon. to Fri.: 8 a.m. – 5 p.m.) or TShelp@accessbio.net (24/7 available).
## Description of Symbols

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>![IVD]</td>
<td><em>In vitro</em> diagnostic medical device. Indicates a medical device that is intended to be used as an <em>in vitro</em> diagnostic medical device.</td>
</tr>
<tr>
<td>![i]</td>
<td>Consult instructions for use. Indicates the need for the user to consult the instructions for use.</td>
</tr>
<tr>
<td>![Manufacturer]</td>
<td>Manufacturer. Indicates the medical device manufacturer.</td>
</tr>
<tr>
<td>![LOT]</td>
<td>Batch code. Indicates the manufacturer's batch code so that the batch or lot can be identified.</td>
</tr>
<tr>
<td>![Do not re-use]</td>
<td>Do not re-use. Indicates a medical device that is intended for one use, or uses on a single patient during a single procedure.</td>
</tr>
<tr>
<td>![Use by date]</td>
<td>Use by date. Indicates the date after which the medical device is not to be used.</td>
</tr>
<tr>
<td>![Prescription-only]</td>
<td>Prescription-only.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>![Catalog number]</td>
<td>Catalog number. Indicates the manufacturer's catalog number so that the medical device can be identified.</td>
</tr>
<tr>
<td>![Caution]</td>
<td>Caution. Indicates the need for the user to consult the instructions for use for important cautionary information such as warnings and precautions that cannot, for a variety of reasons, be presented on the medical device itself.</td>
</tr>
<tr>
<td>![Date of manufacture]</td>
<td>Date of manufacture. Indicates the date when the medical device was manufactured.</td>
</tr>
<tr>
<td>![Temperature limit]</td>
<td>Temperature limit. Indicates the temperature limits to which the medical device can be safely exposed.</td>
</tr>
<tr>
<td>![Do not use if the package is damaged]</td>
<td>Do not use if the package is damaged. Indicates a medical device that should not be used if the package has been damaged or opened.</td>
</tr>
<tr>
<td>![Contains sufficient for &lt;n&gt; tests]</td>
<td>Contains sufficient for &lt;n&gt; tests. Indicates the total number of IVD tests that can be performed with the IVD.</td>
</tr>
</tbody>
</table>

**Manufactured by:**
Access Bio, Inc.
65 Clyde Road, Suite A.
Somerset, NJ 08873, USA
Tel: 732-873-4040
Fax: 732-873-4043
Email: info@accessbio.net
Website: [www.accessbio.net](http://www.accessbio.net)

**Technical Support in the U.S.:**
Tel: +1-888-898-1270 (Toll Free)
Email: TShelp@accessbio.net

**Manufactured for:**
Intrivo Diagnostics, Inc.
2021 Santa Monica Blvd, #11
Santa Monica, CA 90404, USA
Tel: 888-965-0301
Fax: 888-965-0302
Email: info@intrivo.com
Website: [www.intrivo.com](http://www.intrivo.com)

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Revision number: D
Effective date: 2021-6-15
**Quick Reference Instructions for CareStart™ COVID-19 IgM/IgG**

**Rapid Test for the Detection of SARS-CoV-2 IgM/IgG Antibody**

For use under Emergency Use Authorization Only

For Prescription Use only

For In Vitro Diagnostic Use Only

**Important Note:**
- Test should not be used beyond its expiration date.
- Read the complete Quick Reference Instructions before performing the test for fingerstick whole blood.
- Due to the risk of false positive results, confirmation of positive results should be considered using second, different IgG and IgM assay.
- Refer to the package insert for specimen collection and handling for venous whole blood, serum, or plasma.
- Allow test devices, reagents and specimens to equilibrate to room temperature prior to testing.
- Remove the CareStart™ COVID-19 IgM/IgG test device from its foil pouch immediately before specimen collection.

**SPECIMEN COLLECTION AND HANDLING FOR FINGERSTICK WHOLE BLOOD**

**Materials required:**
- Test device
- Assay buffer
- Blood transfer pipette
- Alcohol swab
- Sterile safety lancet
- Package insert
- Quick Reference Instructions (QRI)

**Materials required but not supplied:**
- Timer
- Pair of gloves

1. Clean the fingertip to be pierced with an alcohol swab.
2. Squeeze the end of the fingertip and pierce the cleaned area using a blood lancet. Properly discard the blood lancet.
3. Press the top part of the provided blood transfer pipette.
4. Touch the blood by the pipette tip while pressing the pipette.
5. Fill the pipette with blood sample up to the blue marked line by releasing slowly.

**Access Bio, Inc.**

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Website: www.accessbio.net

Technical Support
Tel: 888-898-1270 (Toll Free)
Email: TShelp@accessbio.net

In the USA, this product has not been FDA cleared or approved but has been authorized for emergency use by FDA under an EUA for use by authorized laboratories. This product has been authorized only for detecting the presence of IgM and IgG antibodies to SARS-CoV-2, not for any other viruses or pathogens. The emergency use of this product is only authorized for the duration of the declaration that circumstances exist justifying the authorization of emergency use of in vitro diagnostics for detection and/or diagnosis of COVID-19 under Section 564(b)(1) of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. § 360bbb-3(b)(1), unless the declaration is terminated, or authorization is revoked sooner.
Quick Reference Instructions for **CareStart™ COVID-19 IgM/IgG Rapid Test for the Detection of SARS-CoV-2 IgM/IgG Antibody**

**TEST PROCEDURES**

6. Add the blood sample to the sample well “S” by pressing the top part of the blood transfer pipette.

7. Open the cap and invert the assay buffer bottle and hold vertically above the sample well. Add one (1) drop of the assay buffer solution to the sample well “S” immediately after sample loading.

8. Start a timer. Read the result at 10 minutes. The test result should not be read after 15 minutes.

**RESULT INTERPRETATION**

- **Positive**
  - IgM Positive
  - IgG Positive
  - IgM, IgG Positive

- **Negative**

- **Invalid**

**NOTES:**
- The color intensity in the test region will vary.
- Any faint colored line(s) in the test region(s) should be considered as positive.

**EXTERNAL CONTROL TESTING PROCEDURES**

User should refer to the instructions for use for the external control testing.

1. Open a control vial containing the control reagents.

2. Transfer the control solution from the control vial:
   a) using a provided blood transfer pipette:
      - Press the top part of the provided blood transfer pipette and touch the control solution by the pipette tip while pressing the pipette. Releasing the press slowly to fill the pipette with sample up to the blue marked line (approximately 10 μL). Add the control sample to the sample well “S” of the test device by pressing the top part of the blood transfer pipette.
   b) using a micropipette:
      - Transfer 10 μL of the control solution to the sample well “S” of the test device using a micropipette.

3. Open the cap and invert the assay buffer bottle and hold vertically above the sample well. Squeeze the bottle gently to add one (1) drop of the assay buffer solution to the sample well “S” immediately after sample loading.

4. Start a timer. Read the result at 10 minutes. The test results should not be read earlier than 10 minutes. Test results should not be read after 15 minutes.

**Expected Results:**

- **Positive Control**
  - C
  - G
  - M

- **Negative Control**
  - C
  - G
  - M

**NOTES:**
- If the test result for either the Negative Control or the Positive Control is not as expected, the test should be repeated using a new Test Device. If the test result for any of the controls is not as expected upon retesting, contact Technical Support.
CareStart™
COVID-19
Antibody External Controls

Package Insert
(Instructions for Use)
# Table of Contents

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CareStart™ COVID-19 Antibody External Controls

For use under Emergency Use Authorization only
For in vitro diagnostic use only
For prescription use only

Intended Use

CareStart™ COVID-19 Antibody External Controls are intended to be used for in vitro diagnostic use in the quality control of the CareStart™ COVID-19 IgM/IgG, Rapid Response™ Liberty COVID-19 IgG/IgM, and CareStart™ EZ COVID-19 IgM/IgG tests.

For in vitro diagnostic use.

When to use

Run the Controls under the following circumstances:

- when a new lot of the test kit is opened
- whenever a new shipment of test kits arrives
- for each new operator before testing
- if the temperature of the test storage area falls outside of the recommended storage conditions
- at periodic intervals as dictated by the user facility.

It is the responsibility of each laboratory using the CareStart™ COVID-19 IgM/IgG, Rapid Response™ Liberty COVID-19 IgG/IgM, or CareStart™ EZ COVID-19 IgM/IgG tests to establish an adequate quality assurance program to ensure the performance of the device under its specific locations and conditions of use.

Summary and Explanation of the External Controls

CareStart™ COVID-19 Antibody External Controls consist of a Positive Control and a Negative Control. External Controls are used as quality control. The purpose of quality control is to ensure the proper performance of the test. The positive control is a mixture of human chimeric SARS-CoV-2 spike S1 reactive IgM and recombinant human SARS-CoV-2 spike S1 reactive IgG antibodies in heat-inactivated SARS-CoV-2 antibody-negative confirmed serum. Negative Control is heat-inactivated SARS-CoV-2 antibody-negative confirmed serum.

Reagents and Materials Provided

<table>
<thead>
<tr>
<th>Contents Name</th>
<th>Quantity (in a kit)</th>
<th>Description</th>
</tr>
</thead>
</table>

Page 3 of 8
**Positive Control**  
1 vial each  
Mixture of human chimeric SARS-CoV-2 spike S1 reactive IgM and recombinant human SARS-CoV-2 spike S1 reactive IgG antibodies in heat-inactivated SARS-CoV-2 antibody-negative confirmed serum.  
< 0.1% sodium azide as a preservative.

**Negative Control**  
1 vial each  
Heat inactivated SARS-CoV-2 antibody-negative confirmed serum.  
< 0.1% sodium azide as a preservative.

### Package insert
1 each  
Instructions for use

* * Materials not supplied
  - 20 µl Micropipette
  - Timer
  - Pair of gloves

### Warnings and Precautions

- For *in vitro* diagnostic use.
- For laboratory/ professional use only.
- For use under Emergency Use Authorization only.
- This product has not been FDA cleared or approved, but has been authorized for emergency use by FDA under an EUA for use by authorized laboratories.
- This product is for use with a test authorized only for detecting the presence of IgM and IgG antibodies to SARS-CoV2, not for any other viruses or pathogens.
- The emergency use of this product is only authorized for the duration of the declaration that circumstances exist justifying the authorization of emergency use of in vitro diagnostics for detection and/or diagnosis of COVID-19 under Section 564(b)(1) of the Federal Food, Drug and Cosmetic Act, 21 U.S.C. § 360bbb-3(b)(1), unless the declaration is terminated or authorization is revoked sooner.
- *CareStart™ COVID-19 Antibody External Controls* are only available for use with the *CareStart™ COVID-19 IgM/IgG, Rapid Response™ Liberty COVID-19 IgG/IgM,* and *CareStart™ EZ COVID-19 IgM/IgG* tests.
- This control is provided for quality assurance only and must not be used for calibration.
- Read this package insert and the test kit package insert completely before using these controls. In order to obtain accurate results, the test must follow this package insert. Handle the External Controls as though they are capable of transmitting infectious agents.
- Do not eat, drink, or smoke in the area where the external controls are handled.
- Nitrile or latex gloves should be worn when performing this test.
- Dispose of used contents as biohazardous wastes in accordance with federal, state, and local requirements.
- Wipe all spills thoroughly with a freshly prepared solution of 10% bleach or other appropriate disinfectants.
- Use of the control reagents manufactured by any other sources may not produce the required results, and therefore, will not meet the requirements for an adequate quality.
assurance program for the CareStart™ COVID-19 IgM/IgG, Rapid Response™ Liberty COVID-19 IgG/IgM, and CareStart™ EZ COVID-19 IgM/IgG tests.

- Reagents contain sodium azide, which is harmful if inhaled, swallowed, or exposed to skin. If there is contact with skin, wash immediately with plenty of water. Sodium azide may react with lead and copper plumbing to form highly explosive metal azides. On disposal, flush with a large volume of water to prevent azide build-up.

Storage and Stability

- Store the External Controls as packaged between 2 - 8°C.
- The External Controls are stable until the expiration date printed on the outer packaging. Do not use beyond the expiration date.
- Open the External Control vials only when performing tests.
- Recap and store the External Control vials in their original package at 2 - 8°C after use.

Test Procedures

Procedural Notes

- Perform test procedures as indicated in the CareStart™ COVID-19 IgM/IgG, Rapid Response™ Liberty COVID-19 IgG/IgM, and CareStart™ EZ COVID-19 IgM/IgG package inserts.

Test Procedures: CareStart™ COVID-19 IgM/IgG

1. Open a control vial containing the control reagents.

2. Transfer the control solution from the control vial:

   a) using a transfer pipette provided in the test kit:
      Press the top part of the transfer pipette and touch the control solution by the pipette tip while pressing the pipette. Releasing the press slowly to fill the pipette with sample up to the blue marked line (approximately 10 µl). Add the control sample to the sample well “S” of the test device by pressing the top part of the transfer pipette.

   b) using a micropipette:
      Transfer 10 µl of the control solution to the sample well “S” of the test device using a micropipette.

3. Open the cap and invert the assay buffer bottle and hold vertically above the sample well. Squeeze the bottle gently to add one (1) drop of the assay buffer solution to the sample well “S” immediately after sample loading.

4. Start a timer. Read the result at 10 minutes. The test results should not be read earlier than
10 minutes. Test results should not be read after 15 minutes.

5. Reseal the control vials and store them in their original container at 2 ~ 8°C.

**Test Procedures:** *CareStart™ EZ COVID-19 IgM/IgG*

1. Gently turn and pull the green sterility tab out, then discard it.

2. Push the grey button firmly against the hard flat surface to retract the lancet.

3. Place the test device on the QRI.

4. Open a control vial containing the control solutions.

5. Transfer the control solution from the control vial:
   
   a) *using a transfer pipette provided in the test kit:*
   
   Press the top part of the transfer pipette and touch the control solution by the pipette tip while pressing the pipette. Releasing the press slowly to fill the pipette with sample up to the blue marked line (approximately 10 µl). Add the control sample to the sample well of the test device by pressing the top part of the transfer pipette.

   b) *using a micropipette:*
   
   Transfer 10 µl of the control solution to the sample well of the test device using a micropipette.

6. Twist off the tip of assay buffer vial to open.

7. Add 4 drops in the sample well.

8. Start a timer. Read the result at 15 minutes. The test result should not be read after 20 minutes.

**Expected Results**

*Positive Control: three distinct lines appear.*

One red-colored line next to “C”, one purple-colored line next to “M”, and one purple-colored line next to “G” indicates SARS-CoV-2 IgM and IgG positive result.
Negative Control: one distinct line appears.

Only one red-colored line next to “C” indicates a negative result.

NOTE: If the test result for either the Negative Control or the Positive Control is not as expected, the test should be repeated using a new Test Device. If the test result for any of the controls is not as expected upon retesting, contact Technical Support.

Technical Support

For questions, or to report a problem, please call Technical Support at +1-888-898-1270 (Available Hours: Mon. to Fri.: 8 a.m. – 5 p.m.) or TShelp@accessbio.net (24/7 available).
Ordering Information

- SCLM-02571: CareStart™ COVID-19 Antibody External Controls, 25 Test
- SCLM-10071: CareStart™ COVID-19 Antibody External Controls, 100 Test

Description of Symbols

<table>
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<tr>
<th>Symbol</th>
<th>Descriptions</th>
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| ![IVD](image1.png) | *In vitro* diagnostic medical device  
Indicates a medical device that is intended to be used as an *in vitro* diagnostic medical device. | ![REF](image2.png) | Catalog number  
Indicates the manufacturer's catalog number so that the medical device can be identified. |
| ![i](image3.png) | Consult instructions for use  
Indicates the need for the user to consult the instructions for use. | ![!] | Caution  
Indicates the need for the user to consult the instructions for use for important cautionary information such as warnings and precautions that cannot, for a variety of reasons, be presented on the medical device itself. |
| ![Manufacturer](image4.png) | Manufacturer  
Indicates the medical device manufacturer. | ![Use by date](image5.png) | Use by date  
Indicates the date after which the medical device is not to be used. |
| ![LOT](image6.png) | Batch code  
Indicates the manufacturer's batch code so that the batch or lot can be identified. | ![Temperature limit](image7.png) | Temperature limit  
Indicates the temperature limits to which the medical device can be safely exposed. |
| ![Σ](image8.png) | Contains sufficient for <n> tests  
Indicates the total number of IVD tests that can be performed with the IVD. | |

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