BioCheck SARS-CoV-2 IgG Antibody Test Kit

Instructions for Use

For Use under an Emergency Use Authorization (EUA) Only
For Prescription Use Only
For in vitro Diagnostic Use Only

REF  BCM-1001

IVD

Σ 10
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Symbols List
The following symbols may appear on the product labels or instructions for use.

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="Image" alt="IVD" /></td>
<td>In vitro diagnostic equipment</td>
<td><img src="Image" alt="REF" /></td>
<td>Catalog Number</td>
</tr>
<tr>
<td><img src="Image" alt="LOT" /></td>
<td>Batch Code</td>
<td><img src="Image" alt="SN" /></td>
<td>Serial Number</td>
</tr>
<tr>
<td><img src="Image" alt="Power on" /></td>
<td>Power on</td>
<td><img src="Image" alt="Power off" /></td>
<td>Power off</td>
</tr>
<tr>
<td><img src="Image" alt="Do Not Re-Use" /></td>
<td>Contains Sufficient for $&lt;n&gt;$ Tests</td>
<td><img src="Image" alt="CE-Marking of Conformity" /></td>
<td>Authorized Representative in the European Community</td>
</tr>
<tr>
<td><img src="Image" alt="Temperature Limit" /></td>
<td>Use-By Date</td>
<td><img src="Image" alt="Consult Electronic Instructions for Use" /></td>
<td>Caution: Federal law restricts this device to sale by or on the order of a licensed healthcare practitioner</td>
</tr>
<tr>
<td><img src="Image" alt="Rx only" /></td>
<td>Manufacturer</td>
<td><img src="Image" alt="Date of Manufacture" /></td>
<td>Distributor</td>
</tr>
<tr>
<td><img src="Image" alt="Importer" /></td>
<td>High temperature, do not touch these areas</td>
<td><img src="Image" alt="Biological hazard" /></td>
<td>Caution: General Warning</td>
</tr>
<tr>
<td><img src="Image" alt="Caution: General Warning" /></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Intended Use

The BioCheck SARS-CoV-2 IgG Antibody Test Kit is a chemiluminescent immunoassay intended for the qualitative detection of IgG antibodies to SARS-CoV-2 in human serum using the MS-Fast Automated Chemiluminescent Immunoassay Analyzing System.

The BioCheck SARS-CoV-2 IgG Antibody Test Kit 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 BioCheck SARS-CoV-2 IgG Test Kit should not be used to diagnose acute SARS CoV-2 infection.

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

Results are for the detection of SARS-CoV-2 IgG antibodies. The IgG antibodies to SARS-CoV-2 are generally detectable in 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 BioCheck SARS-CoV-2 IgG Antibody Test Kit 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 BioCheck SARS-CoV-2 IgG Antibody Test Kit 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 IgG assay.

The BioCheck SARS-CoV-2 IgG Antibody Test Kit is only for use under the Food and Drug Administration’s Emergency Use Authorization.

Background

Severe Acute Respiratory Syndrome Coronavirus 2 (SARS-CoV-2) is a new virus strain that has not been previously identified in humans and belongs to the coronavirus subfamily of the Coronaviridae family. Although it has better sequence conservation compared with the SARS virus in 2003 and MERS virus in 2012, it has low structural homology in the partial protein primary structure of the 19 proteins encoded. Combined with the recently published epidemiological and toxicological evidence, it is suggested that SARS-CoV-2 belongs to a relatively specific coronavirus, and current research shows that the new virus has more than 85% homology with bat SARS-like coronavirus (bat-SL-CoVZC45). This bat SARS-like coronavirus presents the characteristics of strong infection ability, fast transmission speed, and a large number of infections.
SARS-CoV-2 is highly contagious, and the human population is generally susceptible. It is mainly transmitted through respiratory droplets and contact routes, and the virus infects human cells via binding to angiotensin converting enzyme 2 (ACE2).\textsuperscript{1,2} People who are infected with SARS-CoV-2 may be asymptomatic or express signs and symptoms of lower respiratory tract infection such as dry cough, fever, and dyspnea (shortness of breath). In severe cases, the virus can cause acute respiratory distress syndrome (ARDS) and sepsis. Symptomatic, pre-symptomatic, and asymptomatic SARS-CoV-2 carriers all can be potential sources for viral transmission.\textsuperscript{3} Real-time reverse transcription polymerase chain reaction (rRT-PCR) detection of viral genes is the current gold standard for the diagnosis of COVID-19. Upper respiratory specimens, such as nasopharyngeal swabs and oropharyngeal swabs, are commonly used for diagnostic testing.\textsuperscript{2}

The potential public health threat posed by COVID-19 is high, both globally and to the United States. To respond effectively to the COVID-19 outbreak, rapid detection of cases and contacts, appropriate clinical management and infection control, and implementation of community mitigation efforts are critical.

The BioCheck SARS-CoV-2 IgG Antibody Test Kit is intended for the qualitative detection of antibodies which could be indicative recent or prior infection.

**Summary and Principles of the Procedure**

The BioCheck SARS-CoV-2 IgG Antibody Test Kit uses an indirect method to determine the presence of SARS-CoV-2 IgG antibodies in a human serum.

In the cassette, the SARS-CoV-2 IgG antibodies in a patient serum sample bind to biotinylated SARS-CoV-2 S1 antigens. Excess streptavidin coated magnetic beads bind with high affinity to biotinylated SARS-CoV-2 S1 antigen to form complexes. After incubation at 37°C for 10 minutes, the non-specific components in the serum are removed by washing. Next, alkaline phosphatase labeled anti-human IgG antibodies are added to the complexes for incubation at 37°C for 10 minutes, during which time the anti-human IgG antibodies bind to the complexes. This is followed by a second wash to remove excess alkaline phosphatase labeled anti-human IgG antibodies. A luminescent substrate is added, and the enzymes in the complexes catalyze the luminescent substrate to form unstable excited state intermediates. When the excited intermediates return to the ground state, photons are emitted. The number of photons produced is positively correlated with the level of SARS-CoV-2 IgG antibodies in the sample. The more SARS-CoV-2 IgG antibodies present in the sample, the more IgG antibodies that are captured and enzyme-labeled. Producing more enzymes to catalyze substrates, during which more photons (RLU) will be produced. The number of photons (RLU) produced is directly correlated with the level of SARS-CoV-2 IgG antibodies in the sample.
Figure 1: Schematic Diagram of the Reaction Principle of BioCheck SARS-CoV-2 IgG Antibody Test Kit

Warnings and Precautions

General Precautions
- For in vitro diagnostic use under the FDA Emergency Use Authorization.
- For Prescription Use Only
- For in vitro Diagnostic Use Only
- Caution: Federal Law restricts this device to sale by or on the order of a licensed practitioner.
- This test has not been FDA cleared or approved; this test has been authorized by FDA under an EUA for use by laboratories certified under CLIA and meet requirements to perform moderate or high complexity tests.
- This test has been authorized only for the presence of IgG antibodies against SARS-CoV-2, not for any other viruses or pathogens.
- This test is only authorized for the duration of the declaration that circumstances exist justifying the authorization of emergency use of in vitro diagnostic tests for detection and/or diagnosis of COVID-19 under Section 564(b)(1) of the Act, 21 U.S.C. § 360bbb-3(b)(1), unless the authorization is terminated or revoked sooner.
- Laboratories within the United States and its territories are required to report all results to the appropriate public health authorities.
• Negative results do not rule out acute SARS-CoV-2 infection, particularly in those who have been in contact with the virus. Testing with a molecular diagnostic should be performed to evaluate for acute infection in symptomatic individuals.
• The BioCheck SARS-CoV-2 IgG Antibody Test Kit is intended for single use only. Do not reuse.
• 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.
• Consider any materials of human origin as infectious and handle using standard biosafety procedures. No test method can offer complete assurance that hepatitis B virus, hepatitis C virus (HCV), human immunodeficiency virus (HIV 1+2) or other infectious agents are absent.
• Handle, use, store, and dispose of solid and liquid waste from samples and test components in accordance with procedures defined by appropriate national biohazard safety guidelines or regulations.
• The entire test procedure should be performed according to general biosafety guidelines of clinical labs. Protect cuts, abrasions and other skin damages and mucosa from contamination and infection.
• The BioCheck SARS-CoV-2 IgG Antibody Test Kit contain ProClin 300 (CAS 55965-84-9), which may cause an allergic skin reaction. Wear protective gloves when handling the kits. If skin contact occurs, wash with plenty of soap and water. If skin irritation or rash occurs, get medical advice/attention. Wash contaminated clothing before reuse.
• Do not substitute or mix reagents from different kit lots or from other manufacturers. In each kit, sufficient quantities of each component are provided.
• Test result of the same sample using kits from different manufacturers may have variations due to differences in factors such as methodology or antibody specificity.
• Contaminated or suspected contaminated serum specimens or reagents can produce erroneous results.
• Testing must comply with standard laboratory guidelines to prevent cross-contamination.
• Do not use kit after the expiration date.
• Read the package insert and operation manual carefully prior to performing test.

Safety Precautions
• Trained, qualified laboratory personnel should wear appropriate protective gear (gloves, lab coat, safety goggles) and exercise standard precautions for handling and disposing of infectious or potentially infectious materials (e.g. patient samples or consumables).
• All human source materials used in the preparation of the negative control have tested negative for antibodies to HIV 1&2, Hepatitis C and Hepatitis B surface antigen. However, no test method can ensure 100% efficiency. Therefore, all human controls and antigen should be handled as potentially infectious material. The Centers for Disease Control and Prevention and the National Institutes of Health recommend that potentially infectious agents be handled at the Biosafety Level 2.
• Materials used in the tests, including reagent cartridges and the sample collection device, should be disposed of according to federal, state and local regulations.
• Safety Data Sheets (SDS) are available upon request. If the product is a kit or is supplied with more than one component, please refer to the SDS for each component for hazard information.
CAUTION: You must have read and understood the contents of the instructions for use before attempting to run this assay. The contents of this document may be modified by BioCheck at any time. Check www.biocheckinc.com for the latest information.

Materials Provided

The BioCheck SARS-CoV-2 IgG Antibody Test Kit contains the following materials sufficient to perform ten (10) tests.

Kit Components

Table 1. Key Components of the Biocheck SARS-CoV-2 IgG Antibody Test Kit

<table>
<thead>
<tr>
<th>Kit Components</th>
<th>Quantity</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cartridge</td>
<td>10 IgG cartridges</td>
<td>See Table 2.</td>
</tr>
<tr>
<td>Positive control IgG</td>
<td>200μL×1</td>
<td>BSA-containing buffer with human SARS-CoV-2 S1 IgG antibody. Batch specific</td>
</tr>
<tr>
<td>Negative control IgG</td>
<td>200μL×1</td>
<td>BSA-containing buffer. Batch specific</td>
</tr>
<tr>
<td>QR code label</td>
<td>1 labels</td>
<td>Contain positive controls and negative control reference values for IgG on the side of the kit box</td>
</tr>
</tbody>
</table>

Components of the Test Cartridge

The BioCheck SARS-CoV-2 IgG Antibody Test Kit cartridges (Figure 2) contain all the reagent components and accessories needed to run a test as shown in Table 2.

Table 2. Components of the BioCheck SARS-CoV-2 IgG Antibody Test Kit Cartridges

<table>
<thead>
<tr>
<th>#</th>
<th>Test Cartridge Components</th>
<th>Quantity</th>
<th>Main Composition</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Sample Well</td>
<td>1</td>
<td>N/A</td>
</tr>
<tr>
<td>2</td>
<td>Pipette tip</td>
<td>1</td>
<td>N/A</td>
</tr>
<tr>
<td>3</td>
<td>Eluting sleeve</td>
<td>1</td>
<td>N/A</td>
</tr>
<tr>
<td>4</td>
<td>Wash buffer</td>
<td>4mL</td>
<td>0.95% Tween-20, 0.17%, Proclin 300 in Tris buffer</td>
</tr>
<tr>
<td>5</td>
<td>Luminescent substrate</td>
<td>180μL</td>
<td>APS-5 substrate</td>
</tr>
<tr>
<td>6</td>
<td>Magnetic separation reagent</td>
<td>60μL</td>
<td>Streptavidin labeled magnetic beads in PBS buffer</td>
</tr>
<tr>
<td>#</td>
<td>Test Cartridge Components</td>
<td>Quantity</td>
<td>Main Composition</td>
</tr>
<tr>
<td>----</td>
<td>---------------------------</td>
<td>----------</td>
<td>------------------------------------------------------------</td>
</tr>
<tr>
<td>7</td>
<td>Reagent A</td>
<td>80μL</td>
<td>Biotinylated SARS-CoV-2 S1 antigen, 0.1% Proclin 300 in Tris buffer</td>
</tr>
<tr>
<td>8</td>
<td>Reagent B IgG</td>
<td>130μL</td>
<td>ALP labeled Anti-human IgG antibody, 0.1% Proclin 300 in Tris buffer</td>
</tr>
<tr>
<td>9</td>
<td>Sample diluent</td>
<td>540μL</td>
<td>0.9% NaCl in buffer</td>
</tr>
<tr>
<td>10</td>
<td>Sample diluent</td>
<td>405μL</td>
<td>0.9% NaCl in buffer</td>
</tr>
<tr>
<td>11</td>
<td>Empty</td>
<td>N/A</td>
<td>Reaction Well</td>
</tr>
<tr>
<td>12</td>
<td>Empty</td>
<td>N/A</td>
<td>Wash Well</td>
</tr>
<tr>
<td>13</td>
<td>Empty</td>
<td>N/A</td>
<td>Wash Well</td>
</tr>
<tr>
<td>14</td>
<td>Empty</td>
<td>N/A</td>
<td>Reaction Well</td>
</tr>
<tr>
<td>15</td>
<td>Reading aperture</td>
<td>1</td>
<td>N/A</td>
</tr>
</tbody>
</table>

**Materials Required But Not Provided**

MS Fast Chemiluminescent Immunoassay Analyzing System (MS Fast Analyzer) manufactured by Sophonix Co., Ltd. in Beijing, China

- Micropipette and tips for sample addition
- Sample collection materials and centrifuge

**Reagent Storage, Handling and Stability**

Proper storage and handling of reagents and samples are essential for the performance of the BioCheck SARS-CoV-2 IgG Antibody Test Kit.

- Inspect package upon arrival. Kit components should be intact and complete with no leaking, damage, or contamination. All labels should be legible.
- If the foil seals are pierced, do not use the test cartridge.
- Remove kit from the shipping box and immediately store in a dark environment at temperatures between 2-8°C.
- Positive and negative controls are stable for 7 days at 2-8°C after opening.
- Test must be used immediately after opening.
- Do not use the kit after the expiration date. See the packaging label for the product expiration date.

**Sample Requirements**

**Sample Collection and Preparation**

- The BioCheck SARS-CoV-2 IgG Antibody Test Kit should only be performed on human serum samples.
- Collect specimens using universal precautions.¹
- Follow recommended procedures for collection of diagnostic blood specimens by venipuncture.²
- Follow instructions provided with your collection device for use and processing of the sample.³
- Ensure that complete clot formation in specimens has taken place prior to centrifugation.⁴
- Carefully transfer serum to a tightly sealed sterile container for storage.⁵
Avoid use of icteric, hyperlipemic, hemolyzed, or contaminated sera which may cause erroneous results.

Sample Stability and Handling

- Serum samples should be tested within 4 hours at room temperature (approximate range 15-30°C (59-86°F)) after blood collection. If testing cannot be performed immediately, store the serum samples at 2-8°C for up to 24 hours.
- For long-term storage, store small aliquots of samples below -20°C. Avoid repeated freeze/thaw cycles of aliquots. Freeze-thaw damage can result if specimens are frozen in self-defrosting freezers. Thaw and mix samples well prior to use.

Quality Control

External Controls (Positive and Negative Controls)

QC should be performed upon receipt of each new lot or shipment. The detected positive control Relative Light Units (RLU) will be calculated with the standard RLU of the positive control in the QR code to obtain a correction coefficient (M, M = b / a). If the RLU of the negative control after correction by the correction coefficient falls within the specified range of the negative control, the correction proved to be accurate, thus reducing the error caused by different instruments.

The RLU of the sample corrected by the correction coefficient is compared with the cut-off value of the product, and the result is judged to be negative or positive. The above calculation process will be completed automatically by MS-Fast Analyzer, and the result will be reported. The negative and positive controls should result in negative and positive test results, respectively.

If the quality control test results of the positive or negative controls are not within the expected range, repeat the failed control test under the same conditions. A new QC test should be run to obtain the expected positive or negative result. If QC test failure persists, stop using the kit lot and contact the U.S distributor, Accelerate Diagnostics.

Positive and Negative Quality Control (QC) Testing

1. A rack of QC cartridges will take 30 minutes from test start until result acquisition.
2. Unpack the BioCheck SARS-CoV-2 Antibody Test Kit QC contents and equilibrate all materials to room temperature (15-30° C (59-86°F)) before use.
3. Insert four cartridges (two cartridges for positive controls, two cartridges for negative controls) into the reagent rack.

![Figure 3: Inserting Controls Cartridges into Reagent Rack](image)
4. Using a micropipette with a fresh tip for each control, add 80μL of the appropriate controls to the corresponding sample wells of the cartridges.

5. Open the door of the MS Fast Analyzer by sliding it to the left as shown in Figure 6.
6. Insert the reagent rack, then close the sliding door as shown in Figure 7.

![Figure 7: Inserting Rack and Closing Instrument Door](image)

NOTE: The test result of the Negative Control must be negative (-) and the test result of the Positive Control must be positive (+). Otherwise, the kit is invalid, and troubleshooting is necessary to ensure proper function of the instrument reagents, control materials, or parameter settings.

NOTE: If this is the first time the lot or shipment is run:
- From the Main Window, select “Setting”.
- In the Setting Window, select “Reagent Management”.
- Select “SARS-CoV-2 IgG”.
- Click “New batch”.
- Scan each of the QR code labels on the side of the box with the handheld scanner. If the label cannot be scanned, the information can be manually input using the “Manual Input” button.
- Select “Save Data”.
- Repeat for the other antibody kit. Click “Back” when both QR code labels’ information has been saved.

7. From the “Run” screen, double click column corresponding to the cartridge position.
8. Confirm the detected position is correct, then select the project name, lot number, and sample type (i.e., QC).
9. Select positive or negative control and then click “Confirm”. The system will populate the required fields for IgG. Repeat until all controls have been entered.
10. Click “Run” on the Home screen. The MS-Fast Analyzer will automatically identify the test project. When the run is complete, the QC results will automatically print. The QC report can be exported and reprinted.
11. The MS-Fast report will indicate whether the control is valid or not. If the control is valid, the serum sample can be tested next. If the control is not valid, cause for the invalid control should be determined. See the MS-Fast User Manual for more information.
The expected test results for valid negative and positive controls are given below in Table 3.

### Table 3. Expected RLU of IgG Positive and Negative Controls for Valid Runs

<table>
<thead>
<tr>
<th>IgG Control</th>
<th>RLU</th>
<th>Interpretation and Next Steps</th>
</tr>
</thead>
<tbody>
<tr>
<td>If both QC Negative Controls</td>
<td>&lt;23,000</td>
<td>Valid QC Result</td>
</tr>
<tr>
<td>If one QC Negative Control</td>
<td>&lt;23,000</td>
<td>Invalid QC Result. Repeat in duplicate</td>
</tr>
<tr>
<td>If both QC Negative Controls</td>
<td>≥23,000</td>
<td>Invalid QC Result. Repeat in duplicate</td>
</tr>
<tr>
<td>If both QC Positive Control</td>
<td>30,000 to 120,000</td>
<td>Valid QC Result</td>
</tr>
<tr>
<td>If one QC Positive Control</td>
<td>&lt;30,000 or &gt;120,000</td>
<td>Invalid QC Result. Repeat in duplicate</td>
</tr>
<tr>
<td>If both QC Positive Control</td>
<td>&lt;30,000 or &gt;120,000</td>
<td>Invalid QC Result. Repeat in duplicate</td>
</tr>
</tbody>
</table>

**NOTE:** When the negative QC and/or positive QC is not within range of the expected Relative Light Unit (RLU), the instrument should not be used to test samples. The QC should be repeated and if the QC fails again, please contact Accelerate Diagnostics Technical Support. The negative and positive controls should show correct results in order to have valid runs.

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### Procedure

#### Sample Testing

1. A rack of samples will take 30 minutes from test start until result acquisition.
2. The instrument can run up to eight samples at a time.
3. Unpack the IgG test cartridges and equilibrate all materials to room temperature (15-30°C (59-86°F)) before use. Return all unused reagents to the refrigerator.
4. Insert one cartridge for each sample into the reagent rack. Ensure that test strips are locked into the rack (user should hear a click when inserted properly).
5. Using a micropipette, add 80μL serum to the sample well of each cartridge as shown in Figure 8.

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![Figure 8: Test Cartridge Showing Sample Well](image)

6. Slide door of the instrument to the left to open.
7. Insert the reagent rack.
8. From the Sample screen select the SARS-CoV-2 IgG project icon.
9. If using patient or sample barcodes, highlight the “Scan” field and scan in the unique patient (or sample) identifier barcodes using the external barcode scanner. Scan each barcode in the order of the samples in the rack. Click “Exit”.
10. If using manual sample entry, enter the number of samples to test in the “Sample #” field.
11. Push “add data” and double click the “Sample ID” field to manually edit the “Sample ID” field.
12. Click “Done” when complete and then click “Exit”.
13. Close the sliding door and click on “Run”. The MS-Fast Analyzer will automatically identify the test project and provide the final result when testing is complete.

**Interpretation of Results**

**Cut-Off**
The BioCheck SARS-CoV-2 IgG Antibody Test Kit assay uses the receiver operating characteristic (ROC) curve method to determine the positive cut-off value based on a selection of 131 clinically diagnosed serum specimens, including 45 positive serum samples and 86 negative serum samples. Statistical processing was performed to obtain the ROC curve, and the point with the largest Youden index was selected as the critical point, which is the positive cut-off value of the kit. The RLU corresponding to the SARS-CoV-2 IgG cut-off is 26000. When the RLU of the sample test result is ≥26000, the result is positive. When the RLU of the sample test result is <26000, the result is negative.

**Valid Results**
The MS-Fast Analyzer produces the test result when the assay run is complete. Patient sample test results should only be interpreted if the QC test result was valid. The expected test results for valid negative and positive QC results are given in Table 4.

<table>
<thead>
<tr>
<th>IgG Test</th>
<th>RLU Expected</th>
<th>Test Result Interpretation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Positive</td>
<td>≥26000</td>
<td>Positive for SARS-CoV-2 IgG antibodies</td>
</tr>
<tr>
<td>Negative</td>
<td>&lt;26000</td>
<td>Negative for SARS-CoV-2 IgG antibodies</td>
</tr>
</tbody>
</table>

**Invalid Runs**
When the negative control or positive control is not within range of the expected RLU, the test result is not valid and, therefore, the assay should not be used to report results. The positive and negative controls should show passing results prior to running patient specimens.

**Limitations of the Procedure**

- The product can only be used with MS-Fast Automated Chemiluminescent Immunoassay Analyzing System.
- The BioCheck SARS-CoV-2 IgG Antibody Test Kit has not been evaluated for specimens other than human serum.
- Samples that are contaminated or suspected of contamination cannot be used for testing.
- The assay should not be used to diagnose or exclude acute infection. Results are not intended to be used as the sole basis for patient management decisions.
Negative results do not preclude SARS-CoV-2 infection and should not be used as the sole basis for patient management decisions.

A negative or non-reactive 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 the virus has undergone minor amino acid mutation(s) in the epitope recognized by the antibody detected by the test.

The clinical significance of the test results of the kit needs to be analyzed in combination with other test indicators and clinical manifestations.

A positive result may not indicate previous SARS-CoV-2 infection. Consider other information, including clinical history and local disease prevalence, in assessing the need for a second, alternative serology test to confirm an immune response.

It is not known at this time if the presence of antibodies to SARS-CoV-2 confers immunity to reinfection.

Assay performance characteristics have not been established for testing cord blood, for testing neonates, for prenatal screening, or for general population screening.

Results from immunosuppressed patients must be interpreted with caution.

Failure to observe proper procedures for sample collection, preparation, storage, handling, and/or transportation may cause incorrect results.

The kit has not been tested with all types of blood collection tubes that are commercially available. For different blood collection tubes from different manufacturers, different results may be obtained due to different raw materials and additives.

Not for the screening of donated blood.

Conditions of Authorization for the Laboratory


Authorized laboratories using the BioCheck SARS-CoV-2 IgG Antibody Test Kit must adhere to the Conditions of Authorization indicated in the Letter of Authorization as listed below:

- Authorized laboratories using the BioCheck SARS-CoV-2 IgG Antibody Test Kit will 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.

- Authorized laboratories will use the BioCheck SARS-CoV-2 IgG Antibody Test Kit 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 product are not permitted.

- Authorized laboratories that receive the BioCheck SARS-CoV-2 IgG Antibody Test Kit will notify the relevant public health authorities of their intent to run the assay prior to initiating testing.
• Authorized laboratories using the BioCheck SARS-CoV-2 IgG Antibody Test Kit will have a process in place for reporting test results to healthcare providers and relevant public health authorities, as appropriate.

• Authorized laboratories will collect information on the performance of the BioCheck SARS-CoV-2 IgG Antibody Test Kit and report to DMD/OHT7-OIR/OPEQ/CDRH (via email: CDRH EUA Reporting@fda.hhs.gov) and BioCheck Technical Support (https://BioCheckInc.com) any suspected occurrence of false reactive or false non-reactive results and significant deviations from the established performance characteristics of the assay of which they become aware.

• All laboratory personnel using the BioCheck SARS-CoV-2 IgG Antibody Test Kit must be appropriately trained in immunoassay techniques and use appropriate laboratory and personal protective equipment when handling this kit, and use the BioCheck SARS-CoV-2 IgG Antibody Test Kit 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 the BioCheck SARS-CoV-2 IgG Antibody Test Kit.

• BioCheck Inc., authorized distributors, and authorized laboratories using the BioCheck SARS-CoV-2 IgG Antibody Test Kit will 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.

a The letter of authorization refers to, “Laboratories certified under the Clinical Laboratory Improvement Amendments of 1988 (CLIA), 42 U.S.C. §263a, that meet requirements to perform moderate and high complexity tests” as “authorized laboratories”.

Performance Characteristics

Clinical Performance
Clinical studies were conducted using a total of two hundred fifty-three (253) patient samples obtained from China and the United States. One hundred fifty-three (153) of the samples were collected from individuals at two clinical trial institutions in China. Of the 153 samples, 110 had tested positive for SARS-CoV-2 and the other 43 samples tested negative for SARS-CoV-2, both using a comparator PCR testing method. An additional one hundred (100) known negative samples collected in the United States prior to December 2019 from a biobank tested negative for SARS-CoV-2 using a comparator PCR testing method.

For the 110 positive samples obtained from clinical trial institutions in China,109 IgG test results were positive. For the 143 negative samples, comprising 43 pre-pandemic samples from China and 100 pre-pandemic samples from the United States, 143 IgG were negative.

By comparing with the diagnostic results of a nucleic acid detection method, the positive percent agreement (PPA) and negative percent agreement (NPA) of the BioCheck SARS–CoV-2 IgG Antibody Test Kit were determined and shown in Table 5 and Table 6.
### Table 5. IgG PPA

<table>
<thead>
<tr>
<th>Days Post Symptom Onset</th>
<th># PCR Positive At Any Time</th>
<th>BioCheck SARS-CoV-2 IgG Test Kit</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td># Of Positive Results</td>
</tr>
<tr>
<td>≤7</td>
<td>7</td>
<td>7</td>
</tr>
<tr>
<td>8-14</td>
<td>16</td>
<td>16</td>
</tr>
<tr>
<td>≥15</td>
<td>9</td>
<td>9</td>
</tr>
<tr>
<td>unknown</td>
<td>78</td>
<td>77</td>
</tr>
</tbody>
</table>

**Overall IgG PPA:** (99.09% 109/110); [95% CI (95.03%-99.84%)]

### Table 6. IgG NPA

<table>
<thead>
<tr>
<th># PCR Negative</th>
<th>BioCheck SARS-CoV-2 IgG Test Kit</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td># Of Negative Results</td>
</tr>
<tr>
<td>143</td>
<td>143</td>
</tr>
</tbody>
</table>

### Analytical Performance

#### Cross-Reactivity

Cross-reactivity of the BioCheck SARS-CoV-2 IgG Antibody Test Kit was evaluated by judging positive and negative results of specimens serologically positive for other known coronaviruses, respiratory pathogens (both bacterial and viral), and other blood-borne pathogens against the SARS-CoV-2 IgG test.

No cross-reactivity was detected when anti-sera for the following infectious agents was tested on the SARS-CoV-2 IgG test: Human coronavirus panel 229E/HKU1, Human coronavirus panel OC43/NL63, Parainfluenza 1, Parainfluenza 3, Parainfluenza 4, Adenovirus, Rhinovirus, *Mycoplasma pneumonia*, HBV, HCV, Influenza A, Influenza B, and Respiratory Syncytial Virus.

### Table 7. Cross-Reactivity

<table>
<thead>
<tr>
<th>Conditions</th>
<th>Number of Samples</th>
</tr>
</thead>
<tbody>
<tr>
<td>anti-influenza A IgG</td>
<td>5</td>
</tr>
<tr>
<td>anti-influenza B IgG</td>
<td>5</td>
</tr>
<tr>
<td>anti-HCV IgG</td>
<td>5</td>
</tr>
<tr>
<td>anti-HBV IgG</td>
<td>5</td>
</tr>
<tr>
<td>anti-Haemophilus influenzae IgG</td>
<td>5</td>
</tr>
<tr>
<td>Human coronavirus panel 229E/HKU1</td>
<td>5</td>
</tr>
<tr>
<td>Human coronavirus panel OC43/NL63</td>
<td>5</td>
</tr>
<tr>
<td>ANA</td>
<td>5</td>
</tr>
<tr>
<td>anti-respiratory syncytial virus IgG</td>
<td>5</td>
</tr>
</tbody>
</table>

In addition, one hundred (100) known negative samples were collected in the US prior to December 2019 from a population with a high prevalence of vaccination and/or infection and were tested to evaluate potential cross-reactivity. The specificity of IgG test was 100%.
Interference

Low titer SARS-CoV-2 IgG antibody positive serum samples and SARS-CoV-2 IgG antibody negative serum samples were spiked with one of the following substances to specified concentrations. No false-positivity or false-negativity was found.

Table 8. Interference

<table>
<thead>
<tr>
<th>Interfering Substance</th>
<th>Concentration IgG</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hemoglobin</td>
<td>10 mg/mL</td>
</tr>
<tr>
<td>Bilirubin, conjugated</td>
<td>0.4 mg/mL</td>
</tr>
<tr>
<td>Bilirubin, unconjugated</td>
<td>0.4 mg/mL</td>
</tr>
<tr>
<td>Triglycerides</td>
<td>15 mg/mL</td>
</tr>
<tr>
<td>Cholesterol</td>
<td>4 mg/mL</td>
</tr>
<tr>
<td>Human anti-mouse antibody</td>
<td>800 ng/mL</td>
</tr>
<tr>
<td>Rheumatoid factor</td>
<td>2000 IU/mL</td>
</tr>
<tr>
<td>Human serum albumin</td>
<td>60 mg/mL</td>
</tr>
<tr>
<td>Histamine hydrochloride</td>
<td>4 mg/mL</td>
</tr>
<tr>
<td>α-IFN</td>
<td>200 mg/L</td>
</tr>
<tr>
<td>Zanamivir</td>
<td>1 mg/L</td>
</tr>
<tr>
<td>Oseltamivir carboxylate</td>
<td>1 mg/L</td>
</tr>
<tr>
<td>Abidol</td>
<td>40 mg/L</td>
</tr>
<tr>
<td>Levofoxacin</td>
<td>200 mg/L</td>
</tr>
<tr>
<td>Ceftriaxone</td>
<td>400 mg/L</td>
</tr>
<tr>
<td>Meropenem</td>
<td>200 mg/L</td>
</tr>
<tr>
<td>Tobramycin</td>
<td>10 mg/L</td>
</tr>
<tr>
<td>Ribavirin</td>
<td>40 mg/L</td>
</tr>
<tr>
<td>Biotin</td>
<td>3500 ng/mL</td>
</tr>
</tbody>
</table>

Reproducibility

For the BioCheck SARS-CoV-2 IgG Antibody Test Kit, five (5) positive samples and two (2) negative samples, respectively, were selected and tested in duplicate by two (2) different operators at two (2) different labs with three (3) different lots/batches of kits for five (5) consecutive days, resulting in a total of 30 test results per sample. Inter/ intra assay, inter-lab, and inter-lot reproducibility was determined and shown in Table 9.
Table 9. IgG Reproducibility

<table>
<thead>
<tr>
<th>Sample ID</th>
<th>Mean Value</th>
<th>N</th>
<th>Inter-assay</th>
<th>Intra-assay</th>
<th>Between-Days</th>
<th>Between-Lot</th>
<th>Total Precision</th>
</tr>
</thead>
<tbody>
<tr>
<td>Low Positive</td>
<td>40231</td>
<td>30</td>
<td>3197</td>
<td>2720</td>
<td>824</td>
<td>3245</td>
<td>3450</td>
</tr>
<tr>
<td>Moderate Positive</td>
<td>66647</td>
<td>30</td>
<td>5663</td>
<td>5418</td>
<td>1420</td>
<td>5321</td>
<td>6180</td>
</tr>
<tr>
<td>High Positive</td>
<td>181598</td>
<td>30</td>
<td>12738</td>
<td>10533</td>
<td>4443</td>
<td>12793</td>
<td>13349</td>
</tr>
<tr>
<td>Positive</td>
<td>83621</td>
<td>30</td>
<td>5820</td>
<td>4945</td>
<td>1692</td>
<td>5639</td>
<td>6184</td>
</tr>
<tr>
<td>Negative</td>
<td>46216</td>
<td>30</td>
<td>3682</td>
<td>2662</td>
<td>2172</td>
<td>3728</td>
<td>4466</td>
</tr>
<tr>
<td>Negative</td>
<td>7498</td>
<td>30</td>
<td>470</td>
<td>383</td>
<td>239</td>
<td>476</td>
<td>524</td>
</tr>
<tr>
<td>Negative</td>
<td>22974</td>
<td>30</td>
<td>1266</td>
<td>1090</td>
<td>395</td>
<td>1250</td>
<td>1335</td>
</tr>
</tbody>
</table>

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<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>IVD</td>
<td>In vitro diagnostic equipment</td>
<td>REF</td>
<td>Catalog Number</td>
</tr>
<tr>
<td>LOT</td>
<td>Batch Code</td>
<td>SN</td>
<td>Serial Number</td>
</tr>
<tr>
<td></td>
<td>Power on</td>
<td>O</td>
<td>Power off</td>
</tr>
<tr>
<td></td>
<td>Do Not Re-Use</td>
<td>Σ n</td>
<td>Contains Sufficient for &lt;n&gt; Tests</td>
</tr>
<tr>
<td></td>
<td>CE-Marking of Conformity</td>
<td>EC REP</td>
<td>Authorized Representative in the European Community</td>
</tr>
<tr>
<td></td>
<td>Temperature Limit</td>
<td>Use-By Date</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Consult Electronic Instructions for Use</td>
<td>Rx only</td>
<td>Caution: Federal law restricts this device to sale by or on the order of a licensed healthcare practitioner</td>
</tr>
<tr>
<td></td>
<td>Manufacturer</td>
<td>Date of Manufacture</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Distributor</td>
<td>Importer</td>
<td></td>
</tr>
</tbody>
</table>

Symbols List
<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><img src="image1" alt="Symbol" /></td>
<td>High temperature, do not touch these areas</td>
<td><img src="image2" alt="Symbol" /></td>
<td>Biological hazard</td>
</tr>
<tr>
<td><img src="image3" alt="Symbol" /></td>
<td>Note, refer to the attached file</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
1 About this manual

This manual contains the MS-Fast Automated Chemiluminescent Immunoassay Analyzing System (hereinafter referred to as "MS-Fast") produced by Sophonix Co., Ltd. and distributed by Accelerate Diagnostics, Inc. Please read this manual carefully before attempting to operate MS-Fast.

2 About MS-Fast

This section includes the intended use, operating characteristics, and instrument performance of the MS-Fast Analyzer.

2.1 Intended Use

The MS-Fast Analyzer is an automated discrete loading chemiluminescence enzyme-linked immunosorbent assay (CLEIA) and magnetic particle separation analyzing system that uses indirect chemiluminescence based, on a chemiluminescent substrate (APS-5) and alkaline phosphatase. The BioCheck SARS-CoV-2 IgM and IgG Antibody Combo Test Kit is for qualitative detection and differentiation of IgG and IgM antibodies to SARS-CoV-2 in human serum using the MS-Fast Automated Chemiluminescent Immunoassay Analyzing System. It is intended for use as an aid in identifying individuals with an adaptive immune response to SARS-CoV-2, indicating recent or prior infection.

- The BioCheck SARS-CoV-2 IgM and IgG Antibody Combo Test Kit has not been FDA cleared or approved; this test has been authorized by FDA under an EUA for use by laboratories certified under CLIA and meets requirements to perform moderate or high complexity tests.
- The BioCheck SARS-CoV-2 IgM and IgG Antibody Combo Test Kit has been authorized only for the presence of IgG or IgM antibodies against SARS-CoV-2, not for any other viruses or pathogens.
- The BioCheck SARS-CoV-2 IgM and IgG Antibody Combo Test Kit is only authorized for the duration of the declaration that circumstances exist justifying the authorization of
emergency use of in vitro diagnostic tests for detection and/or diagnosis of COVID-19 under Section 564(b)(1) of the Act, 21 U.S.C. § 360bbb-3(b)(1), unless the authorization is terminated or revoked sooner.

### 2.2 Specifications and performance

<table>
<thead>
<tr>
<th>Instrument type</th>
<th>Automated chemiluminescent immunoassay analyzing system</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model</td>
<td>MS-Fast Analyzer</td>
</tr>
<tr>
<td>Detection batch</td>
<td>8-channel parallel detection</td>
</tr>
<tr>
<td>Sample material</td>
<td>Serum</td>
</tr>
<tr>
<td>Sample volume</td>
<td>80ul</td>
</tr>
<tr>
<td>Wavelength</td>
<td>300-650nm</td>
</tr>
<tr>
<td>Data storage</td>
<td>Patient data: more than 100,000 samples (tests)</td>
</tr>
<tr>
<td>QC data</td>
<td>More than 100,000 results</td>
</tr>
<tr>
<td>Size</td>
<td>508 (W) x 586 (D) x 343 (H) mm</td>
</tr>
<tr>
<td>Weight</td>
<td>40 kg</td>
</tr>
<tr>
<td>Electrical requirements</td>
<td>110/220 VAC</td>
</tr>
<tr>
<td>Rated power supply</td>
<td>110/220V 60/50 Hz 200W</td>
</tr>
<tr>
<td>Printer</td>
<td>Built in</td>
</tr>
<tr>
<td>Computer</td>
<td>Built in</td>
</tr>
<tr>
<td>Screen</td>
<td>7 in LCD touch screen (800 x 600) px</td>
</tr>
<tr>
<td>Operating System</td>
<td>Microsoft Windows10 Professional</td>
</tr>
</tbody>
</table>

### 3 Details of MS-Fast

This section describes the measurement principle, equipment structure, and accessories used in MS-Fast.

#### 3.1 Principle of detection

The detection principle used by MS-Fast is Chemiluminescent Enzyme Immunoassay (CLEIA), and the reagents used in each test are stored in a special, individually packaged cassette sealed with aluminum foil.

The MS-Fast detection principle is described by applying the sandwich immunoassay reaction mode. The sample is combined with an alkaline phosphatase-labeled antibody and a biotinylated antibody to form a sandwich structure. The excess streptavidin-coated magnetic particles form a complex with the biotinylated antibody, and after washing, the luminescent substrate is
catalytically cleaved by an enzyme in the complex to form an unstable excited intermediate. When the particle returns to the ground state, it emits photons. The light intensity of the reaction is detected by a photomultiplier tube, and the luminescence intensity of the analyte is converted into relative light units (RLU) by the instrument operating software.

3.2 MS-Fast components

This section describes the components of the MS-Fast and their functions.

3.2.1 Front of the Device

The necessary components for operating the MS-Fast are located at the front and side of the unit; see the figure below. Component names and functions are described as follows:

- **Touchscreen computer** - Displays various information about the MS-Fast Analyzer and accepts input from the user operating the device. Select the appropriate program.
- **Sliding door** - Slide to the left to open; insert the rack after opening. The instrument automatically scans the barcode, and the user can read the cassette information and select the appropriate program.
- **Printer** - Prints results and other information.
3.2.2 Back of the Device

Supply voltage = 110VAC                  Supply voltage = 220VAC

The input and output ports of the device are mainly located on the back of the device, as shown in the figure above.

- **Switch** – This is the main power switch of the device. After the instrument is turned on, the computer powers up in about 1 minute.
- **USB** – Provides connection to the internal instrument computer.
- **110/220 VAC** – Provides connection to the power cable.
- **Fuse: F2AL250V/F3AL250V** – Protects the internal circuitry of the instrument.
- **Selected Switch** – For use by manufacturer or distributor

3.2.3 Accessories

Information on the main MS-Fast accessories is shown in the table below.

<table>
<thead>
<tr>
<th>Main Accessories Description</th>
<th>Quantity</th>
<th>Remarks</th>
</tr>
</thead>
<tbody>
<tr>
<td>Power cord</td>
<td>1</td>
<td>Standard</td>
</tr>
<tr>
<td>Fuse: F2AL250V/F3AL250V</td>
<td>2</td>
<td>Alternate replacement</td>
</tr>
<tr>
<td>Cartridge rack</td>
<td>1</td>
<td>Standard</td>
</tr>
</tbody>
</table>
4 Storage and Operation of the Equipment

4.1 Environmental conditions and placement instructions

4.1.1 Normal working conditions

1) Altitude: no more than 2000m
2) Power supply voltage: 110-220V 50/60 Hz 200W
3) Ambient temperature: 10 °C ~ 30 °C;
4) Relative humidity: ≤70%;
5) Atmospheric pressure: 85.0 kPa to 106.0 kPa;
6) Keep away from magnetic field interference sources, e.g., MRIs;

4.1.2 Placement instructions

1) The instrument should be placed on a flat, stable surface.
2) The rear panel of the instrument requires a distance of >200 mm for access the power switch and ventilation.
3) Avoid placing the MS-Fast in a place where it is exposed to direct sunlight, air conditioning, or other equipment.
4) Do not block the device vents, and ensure that there is no paper or other objects that may affect the air circulation in this area.
5) MS-Fast is placed at an altitude of no more than 2000 meters.
5 Software Description

5.1 Software Overview

This section describes the software used with MS-Fast.

5.1.1 Software Environment

1) Uses a custom 7-inch touch screen, with a resolution of 800X600 px

2) Microsoft Windows10 Professional operating system.

5.2 System startup

Power on the system using the power switch. If the Sophonix software does not automatically start, double-click the desktop shortcut “POCT_Software”.

![POCT Software Icon](image-url)
5.3 Log in

1) To log in, enter the username and password, you can select whether to save the password. When the "Confirm" button is clicked, the system checks the username and password, and then login to the system is completed after meeting the system requirements.

2) Usernames are used to distinguish system permissions: user permissions are divided into four levels: A, B, C, and D.
   - Permission A and B: for manufacturing or distributor use
   - Permission C: administrative permissions (user permissions, user settings, re-collection of calibration curves, etc.)
   - Permission D: user permission (sample detection, calibration, quality control, result query, report printing, adding new batch of reagents)
5.4 Initialization check

1) After login the system will initialize. A dialogue box will show “Initialization complete”, and click on the “Confirm” box to access the main window.

2) If an error, like the ones below, is encountered contact customer support.
5.5 Main window

1) Click "Run" to enter the device running window
2) Click on "Sample" to go to the Edit Samples window
3) Click "Result" to enter the results query window
4) Click "Setting" to enter the settings window.

5.6 Run window

The “Run” window is the core part of the normal operation and shows the status of the instrument. The whole operation process is carried out in this window.
5.6.1 Routine detection settings

All pertinent information, such as the project, sample, and batch number can be manually added on a channel by channel basis from the “Run” Window.

1) Double-click the "Project" or "Sample" positions to pop up a window for selection.
2) Confirm detection position is correct, and then select the project name, batch number, and sample type (Sample or QC).

### 5.7 Sample window

The “Sample” window allows you to add the sample and project information for the cassettes, as well as display sample and project information that has been set, edit patient information for samples, and more.

![Sample window interface](image)

#### 5.7.1 Scan Accession number

1) Select the project(s) for the cassette and then scan the accession number into the “Scan QR” toolbar.

2) If you do not want to wish to add an accession number, the cassette QR code can be scanned instead or use the “Add Data” button shown below.

3) Add data
Select the project and enter the number of samples. Click "Add Data" to automatically fill in "Testing Date", "Channel", "Sample ID".

**NOTE**: Date of testing and results follow the Year-Month-Day format.

5.7.2 Edit record

1) Double-click the record to go to the edit information interface.

2) Fill in the corresponding data, click the "Continue" button to save the current data, and load the data for the next cassette slot.

3) Click the "Cancel" button to cancel the current operation.

4) Click the "Done" button to close the current interface and return to the previous interface.
5.7.2 Set up project portfolio

1) Double-click the project portfolio button at the top to enter the portfolio Settings window.

2) Go to the portfolio settings.

3) Modify the combination name and select or deselect the project name, blue is selected, white is deselect. Click the "Confirm" button to save the current setting item information.

5.7.3 Delete data

Select a row record and click the "Delete" button to delete the selected record data.

5.7.4 Historical data

1) Click to enter the history interface for querying sample information. Update the testing date range and click search.
2) According to the date of detection, query all sample information for the period of time.

Double-click the record to go to the edit Information screen.

5.7.5 Exit
Exit the current interface

5.8 Result window
In the "Result" window, you can query the experimental results, including the test results, the calibration results and quality control results.
5.8.1 Find Sample

1) After entering the "Result" window, the software automatically displays to the "Find Sample" interface.
2) Under this interface, you can select the date of Detection, and “Project”, query by “Sample ID” or “Name”. The “Testing Date” is mandatory to see results, while all other search criteria is optional. If you do not enter information, you do not select the condition.

3) Click the “Search” button to retrieve the experimental results of all samples under the current selection condition and display them in the data area below.

4) Click “Upload to Host” to upload the results of the current query to the LIS system.

5) Click “Print List” to print the record of the current query by list.

6) Click “Export” to export the data and save the format as text.

7) Click “Export-Excel” to export the data and save the format as Excel.

5.8.2 Find QC

1) Click on “Find QC” to enter the “Find QC” window.
2) “Project Name”, “Start Time”, and “End Time” are required items, the rest are optional.

3) Click the “Search” button to retrieve all the quality control results from the selected conditions.
4) Click “Save QC chart” to save the currently displayed quality control charts.

1) Print
   Print list format.
2) PDF
   Save as PDF
3) Export
   Click the "Export" button to export the data and save the format as text.

5.9 Setting window

The “Setting” window includes Reagent Management, Hospital Information Setup, User Settings and Switch user.

1) Click the corresponding button to enter the corresponding window.

5.9.1 Reagent Management

In the “Reagent Management” interface, you can enable and disable projects; query project parameters; and add new batch reagents.
1) Add new batches

- Select the “Project List” on the left side of the screen, and the detailed information of the currently selected project will automatically be displayed.
- Click the “New Batch” button to select a new batch of the currently selected project.
• **Scan QR**: When entering the interface, scan the QR code on the kit with the handheld scan code gun. After the scan is successful, the current scan information is automatically added.

• **Manual Input**: If unable to be scanned, the information can be manually added via the “Manual Input” button.

**NOTE**: After scanning or manually inputting the batch information, the “Reagent Batch” and “Expiry to” values are unable to be changed.

- After inputting, click the “Save Data” button to save the current batch data.
- You can also add multiple batches of information at once. Repeat the above operation.
- Click the "Back" button to exit the current interface and return to the reagent management interface.
2) Add a user

- In setting, select the “User Setting” button to add a user and password.

- “Authority”: A is the highest level and D is the lowest level.

- The current user’s authority cannot be higher than the new User they are setting up.

- Modify the “Authority” and “Password” of the original users.

- Add a user, enter "User Name", select “Authority”, set "Password", the password can be empty.

- Click the "Confirm" button to save the current information.

- Click "Back" to close the current window and return to the settings window.
3) To update, modify, or delete

- Click on the corresponding tab to update a department, patient category, submitting doctor, examiner, or auditor.

- Go back to the login window and enter your username and password. Re-register.

5.10 Start initialization

- Click the "Initialization " button in the main window. System will be re-initialized.

5.11 Shut down

- Click the "Shut Down" button in the main window to access the shutdown option.
6 Instrument warnings and precautions

6.1 Precautions

1) Only trained, qualified laboratory personnel are authorized to use the instrument. Authorized personnel must wear appropriate protective gear (gloves, lab coat, safety goggles) and exercise standard precautions for handling and disposing of infectious or potentially infectious materials (e.g. patient samples or consumables).

2) The test cassette provided by the factory will not release toxic and harmful gases during the use of the instrument. The test sample is potentially infectious. The used test cassettes are treated according to the Medical Waste Management Regulations to avoid biohazard. If hazardous materials leak into the surface or inside the equipment, proper cleaning should be carried out (see section 8)

3) During cleaning of the instrument ensure that the instrument is in turned off.

4) The operator must follow the instructions in this manual in order to ensure safe operation.

5) MS-Fast is a medical analytical instrument for testing serum. Do not use the device for other purposes.
6) Do not use solvents, abrasive cleaning fluids, or rough cloth to clean any parts. Do not use cleaning agents or disinfectants that are not approved for cleaning or disinfections as specified in section 8.

7) The door should not be opened during the operation of the instrument to avoid injury caused by touching moving parts.

8) The user must not open service door or remove any covers from the instrument.

7 Troubleshooting

<table>
<thead>
<tr>
<th>Error Alarm</th>
<th>Cause of Issue</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Z motion abnormality</td>
<td>Magnetic rod motor or magnetic sleeve</td>
<td>1. The operator turns the machine on and off once, so that the instrument is reset again to see if the fault persists.</td>
</tr>
<tr>
<td></td>
<td>magnetic sleeve</td>
<td></td>
</tr>
<tr>
<td></td>
<td>motor movement abnormal</td>
<td></td>
</tr>
<tr>
<td>TIP movement abnormality</td>
<td>TIP motor motion abnormality</td>
<td>3. The operator turns the machine on and off once, so that the instrument is reset again to see if the fault persists.</td>
</tr>
<tr>
<td>Abnormal pump motion</td>
<td>Sample pump motor movement is abnormal</td>
<td>4. If the fault persists, please contact the factory after-sales engineer or an authorized technical service representative.</td>
</tr>
<tr>
<td>PMT movement abnormality</td>
<td>PMT motor motion abnormality</td>
<td>1. The operator turns the machine on and off once, so that the instrument is reset again to see if the fault persists.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. If the fault persists, please contact the factory after-sales engineer or an authorized technical service representative.</td>
</tr>
</tbody>
</table>
## Error Alarm

<table>
<thead>
<tr>
<th>Error Alarm</th>
<th>Cause of Issue</th>
<th>Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Y-axis motion anomaly</td>
<td>Y-axis motor movement is abnormal</td>
<td>1. The operator turns the machine on and off once, so that the instrument is reset again to see if the fault persists.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. If the fault persists, please contact the factory after-sales engineer or an authorized technical service representative.</td>
</tr>
<tr>
<td>X-axis motion abnormality</td>
<td>X-axis motor movement is abnormal</td>
<td>1. The operator turns the machine on and off once, so that the instrument is reset again to see if the fault persists.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. If the fault persists, please contact the factory after-sales engineer or an authorized technical service representative.</td>
</tr>
<tr>
<td>Fitting exception, data processing is unsuccessful</td>
<td>Reagent related data processing exception</td>
<td>1. The operator turns the machine on and off once, so that the instrument is reset again to see if the fault persists.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>2. If the fault persists, please contact the factory after-sales engineer or an authorized technical service representative.</td>
</tr>
<tr>
<td>Open the serial port exception</td>
<td>Thermal printer communication is abnormal</td>
<td>Contact the manufacturer's after-sales engineer or an authorized technical service representative.</td>
</tr>
</tbody>
</table>

### 8 Equipment maintenance

When servicing the equipment, turn off the power and wear appropriate safety clothing (gloves, test jackets, goggles, etc.) to keep the equipment accessories and reagent racks clean.

#### 8.1 Weekly maintenance

1) Clean the Reagent Rack

- Gently clean the surface of the reagent rack with a lint free wipe soaked with disinfectant 70% alcohol.
- Ensure all channels are disinfected with 70% alcohol when cleaning the channels of the reagent rack.
8.2 Monthly maintenance

1) Clean the surface of the device with 70% alcohol.

8.3 Recommended regular maintenance

To ensure proper operation and reliability of the MS-Fast instrument, follow recommendation for periodic maintenance by the manufacturer's after-sales engineer or an authorized MS-Fast technical service representative.

8.4 Replacing the internal printer paper

1) Power off the instrument before replacing printer paper.
2) Pull the printer cover located at the lower left of the front of the unit.
3) Remove the used paper roll and install new paper roll.
4) Push the retaining plate back into place and use the paper cutter to cut off excess paper.