Alere™ Triage® MeterPro Troponin I METHOD AND SAMPLE COLLECTION

1. PURPOSE AND SCOPE

The purpose of this document is to describe the procedure for performing a Troponin I using the Alere Triage MeterPro analyser. The Alere Triage MeterPro analyser is to be used by trained health care professionals in accordance with a facility’s policies and procedures. This document will be focusing on Troponin I from the multi-marker panels, and further information on the other parameters can be found in the appropriate product inserts.

2. HAZARDS

Patient Samples
All patient samples should be treated as potentially infectious and handled appropriately. Some quality control solutions contain human source material. Personal Protective Equipment should be worn when processing samples, quality control testing and maintenance procedures.

3. CLINICAL SIGNIFICANCE

Acute Coronary Syndrome (ACS) is a term used to describe a group of conditions resulting from insufficient blood flow to the heart muscle. These conditions range from atypical chest discomfort and non specific electrocardiographic changes to a large ST-segment elevation, myocardial infarction and cardiogenic shock. Symptoms can include chest pain including tightness and heaviness in the chest, discomfort in the arms and upper body, shortness of breath and other constitutional symptoms including sweating, nausea and light-headedness.

Diagnosis of ACS is based on a complete medical history, physical examination, electrocardiogram to evaluate the electrical activity of the heart and blood tests to evaluate the presence of biological markers resulting from cardiac cell injury.

Troponin T & I are members of a group of cardiac regulatory proteins which function to regulate the calcium mediated interaction of muscle filaments actin and myosin resulting in contraction and relaxation of striated muscle. Troponin T is almost exclusive to the myocardium, with small amounts expressed in skeletal muscle not detectable in current Troponin T assays. Insufficient blood flow and oxygen supply to the heart muscle causes necrosis of the myocardium and subsequent release of Troponin T & I into the bloodstream.

Troponin T in the bloodstream rises to detectable levels after 4-6 hours, peaks at 10-12 hours and can be detected for up to 14 days post infarction. Troponin I is released from necrotic cardiac myocytes into the bloodstream within hours (~4-8 hours) after the onset of chest pain. The peak TnI concentration is generally reached in 12-48 hours. Troponin I serum levels can remain elevated for up to 4–7 days.

The diagnostic utility of Troponin T & I to detect myocardial necrosis and to enable
risk stratification in patients with ACS is well established.\textsuperscript{5,8} Furthermore, the use of Troponin T as a prognostic indicator for recurrence of ischaemic events and death in ACS patients is increasing.\textsuperscript{5,9}

Results from PoCT devices measuring Troponin T & I should always be used in conjunction with clinical presentation, history and other diagnostic information.

4. TEST PRINCIPLE

The Alere Triage Troponin I Test is a single use fluorescence immunoassay device designed to determine the concentration of Troponin I in EDTA anticoagulated whole blood or plasma specimens.

The test procedure involves the addition of several drops of an EDTA anticoagulated whole blood or plasma specimen to the sample port on the Test Device. After addition of the specimen, the whole blood cells are separated from the plasma using a filter contained in the Test Device. The specimen reacts with fluorescent antibody conjugates and flows through the Test Device by capillary action. Complexes of each fluorescent antibody conjugate are captured on a discrete zone resulting in a binding assay.

The concentration of Troponin I in the specimen is directly proportional to the fluorescence detected; therefore, a greater amount of fluorescence indicates a higher Troponin I concentration. Light from a laser hits a test device that has been inserted in the meter, which causes the fluorescent dye in the test device to give off energy. The more energy the fluorescent dye gives off, the stronger the signal. The results are displayed on the Meter screen in approximately 20 minutes.

4.1 Interference

- Severely haemolysed specimens should be avoided. When a sample appears to be severely haemolysed, another specimen should be obtained and tested
- As with any assay employing mouse antibodies, the possibility exists for interference by human anti-mouse antibodies (HAMA) in the sample. The test has been formulated to minimize this interference; however, specimens from patients who have been routinely exposed to animals or to animal serum products may contain heterophile antibodies which may cause erroneous results
- The hematocrit between 30% and 55% had no significant effect on the recovery of Troponin I.
- No high dose hook effect was observed with the Alere Triage panel assays for Troponin I up to 2,100 ng/mL.
- Further information of substances that were tested can be found in the corresponding Test Device package insert

Important! It is possible that other substances and/or factors not listed above may interfere with the test and cause false results.

4.2 Accuracy & Precision

This product fulfills the requirements of the EU Directives.
Specific details for the accuracy and precision for the different Test Panels can be found in the corresponding Test Device package insert.

5. INSTRUMENT

Product specifications
5.1 Operating Conditions and Technical Data

<table>
<thead>
<tr>
<th>Specification</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature range</td>
<td>15° – 30°C</td>
</tr>
<tr>
<td>Relative humidity</td>
<td>10 - 85% (non-condensing)</td>
</tr>
<tr>
<td>Maximum altitude</td>
<td>Not stated</td>
</tr>
<tr>
<td>Position</td>
<td>Place meter on a level, vibration-free surface, away from direct sunlight</td>
</tr>
<tr>
<td>Measuring range</td>
<td>0.05 - 30 ng/mL</td>
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<tr>
<td>Sample size</td>
<td>Not stated</td>
</tr>
<tr>
<td>Test time</td>
<td>20 minutes</td>
</tr>
<tr>
<td>Memory</td>
<td>750 patient records, 200 QC sample results, 70 QC device results, 250 misc. test results, 600 User IDs</td>
</tr>
<tr>
<td>Barcode scanner</td>
<td>Optional external scanner</td>
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<tr>
<td>Interface</td>
<td>RS-232 computer interface port</td>
</tr>
<tr>
<td>Battery operation</td>
<td>Yes – 4x AA batteries</td>
</tr>
<tr>
<td>Mains connection</td>
<td>Yes – 6v at 1 amp</td>
</tr>
<tr>
<td>Number of tests with fully charged batteries</td>
<td>Approx. 100 tests</td>
</tr>
<tr>
<td>Safety class</td>
<td>Not stated</td>
</tr>
<tr>
<td>Automatic power-off</td>
<td>Yes – if left unused for 2 hours. Programmable for 0.5, 1, 2, or 4 hours</td>
</tr>
<tr>
<td>Dimensions</td>
<td>225 x 190 x 70 mm</td>
</tr>
<tr>
<td>Weight</td>
<td>700g excl. batteries</td>
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5.2 Storage and transport conditions

<table>
<thead>
<tr>
<th>Temperature range</th>
<th>Not stated</th>
</tr>
</thead>
<tbody>
<tr>
<td>Meter (In original container)</td>
<td>Not stated</td>
</tr>
</tbody>
</table>

6. SPECIMEN REQUIREMENTS

Venous whole blood or plasma collected in EDTA tubes is acceptable. Other blood collection tubes containing other additives and other blood specimen types have not been evaluated.

Venepuncture (see suitable anticoagulants above)

- Skin surface must be cleaned with an alcohol swab and dried well prior to collection to ensure there are no substances on the skin surface.
- Ensure sample is properly mixed and at room temperature before testing.
- Test the sample within 1 hour of collection.
- Transport specimens at room temperature or chilled and avoid extreme temperatures.
- Avoid using severely haemolysed specimens whenever possible. If a specimen appears to be severely haemolysed, another specimen should be obtained and tested.

7. CARTRIDGES / REAGENTS

7.1 Storage and handling

- Perform Troponin I tests using the following Test Devices: Alere Triage Cardiac Panel (CK-MB, Myoglobin, Troponin I), Alere Triage CardioProfiler Panel (CK-MB, Myoglobin, Troponin I & BNP), or the Alere Triage Profiler SOB Panel (CK-MB, Myoglobin, Troponin I, BNP and D-Dimer).
- Unopened Test Devices should be refrigerated at 2° - 8°C and are stable up to the expiration date.
- Once removed from refrigeration, the pouched Alere Triage Cardiac Panel, CardioProfiler and SOB Test Devices are stable for up to 14 days at room temperature, but not beyond the expiration date printed on the pouch.
- Once equilibrated to room temperature, do not return the Test Device to refrigeration.
- Optimal results will be achieved by performing testing at temperatures between 20-24°C.
- Test Devices must be at room temperature before use (i.e. allow the unopened Test Device to sit at room temperature for at least 15 minutes).
- If a kit containing multiple Test Devices is removed from refrigeration, allow the kit to reach room temperature before use. This will take a minimum of 60 minutes.
- Keep the Test Device in the sealed pouch until ready for use. Discard after single use.
- The transfer pipette should be used for one patient specimen only. Discard after single use.
• Discard the Test Devices if they are past the used by date. Expired Test Devices can produce incorrect results.
• The QC Device is light sensitive and should be stored in its black opaque case when not in use.

7.2 Storing information about test devices
Meter data is updated via a disposable Code Chip module. The Code Chip module contains microchip circuitry embedded into a plastic housing. When inserted into the meter Code Chip module port and activated, the information is transferred into the meter’s memory. A Code Chip module typically needs to be installed only once per box of strips and remains in the meter memory. A Code Chip module does not need to remain in the meter while performing tests.

8. CALIBRATION
In the self-test mode the meter scans an Internal Calibration Chip. Each calibration chip scan is used to validate and adjust, if necessary, the meter calibration. Operator calibration is not necessary.

9. Quality CONTROL

Quality control material (perform as per your organisation protocol)
Accurately testing known levels of Troponin I ensures that the system and your technique used in testing give accurate results on patient tests. The control solutions have defined (known) values. The results for these solutions must first fall within a certain acceptable range in order to allow valid patient testing.

A quality control test should be performed every time a new shipment of test devices are received, when a new lot number of test devices are used, if the clinical picture does not correlate with the patient test results, after major maintenance, and at a minimum of once a month.

Enrolling in an External Quality Assurance Program is encouraged to objectively compare results with other users using the same method of testing. If an External Quality Assurance Program is not available, monthly lab comparison is encouraged.

The Alere Triage MeterPro uses the following methods for quality:
• Built-in QC Check
• Code Chip modules
• Electronic QC Device test
• Control solutions

9.1 Built-in internal QC check
The Test Devices contains a built-in internal QC Check, which is performed automatically by the Meter during every assay. These assay controls in each test device monitors the assay procedure and the reagent integrity. If the results are found to be in specification, the result is allowed to be used, but if the test device’s internal quality control (QC) results are unacceptable, the results for the affected analyte(s) will not appear on the screen (instead will be flagged with an exclamation mark symbol).
9.2 Installing the Reagent, QC Sample or QC Device Code Chip modules

- From the ‘Main Menu’ select ‘Install Code Chip’ using the up/down arrow keys and press ‘Enter’ to confirm selection.
- Use the new Code Chip module that comes with every new box of test devices.
- Compare the lot number on the code chip with the corresponding lot number on the test strip pouch.
- Insert the Code Chip module into the Code Chip module Port in the lower left front corner of the meter.
- The meter will display a confirmation message that the information was installed into the meter and press the ‘Enter’.
- Remove the Code Chip module from the Code Chip module Port.
- Place the Code Chip module back into its original container for storage.

**NOTE:** The meter will direct the user to install a Code Chip module if it is attempting to run a test and does not have the data in its memory.

9.3 Electronic QC Device test

The QC Device should be run on each day of patient testing. The QC Device Code Chip Module should be performed before running a QC Device test, as stated in section 9.2.

To perform a QC Device test:

- Press the ‘Power’ button to turn on the meter.
- Select ‘Run Test’ using the up/down arrow keys and press the ‘Enter’ key to confirm selection.
- Type in your identification number (User ID) if prompted.
- Select ‘QC Device’ using the up/down arrow keys and press the ‘Enter’ key to confirm selection.
- Gently insert the QC Device into the meter until you feel the QC Device catch on the pin and hear a ‘click’.
- Press the ‘Enter’ key to start the test.

**NOTE:** The meter will prompt the user to install the QC Device Code Chip module if the QC Device has not been run before.

- The meter pulls in the QC Device and scans it. The test device may partially move in and out of the meter several times.

**NOTE:** After the test device has been inserted, do not push the device in further or attempt to pull it out. The device may be ejected by returning to the Main Menu and pressing the ‘Exit’ key.

- When the test is complete, the meter will beep, eject the device and display a Pass or Fail result on the meter’s screen.
- Press the ‘Print’ key to make a printed copy of the results.
• Remove the QC Device from the meter and place in the QC Device Box. Do not discard the QC Device.
  **NOTE:** The QC Device is light sensitive and should be stored in its black opaque case when not in use. Keep the QC Device free of contaminants, as dust, lint, fibers and other small particles may interfere with the QC Device.
• If the QC Device tests fail, wipe the QC Device clean with a lint free cloth to remove any oils, dust, fibers, or fingerprints. Do not apply any liquid to the QC Device. After cleaning the device, repeat the QC Device test.
  **NOTE:** The QC Device does not expire.

9.4 Running control solutions
The QC Device Code Chip Module should be performed before running a QC Device test, as stated in section 9.2
The control solutions has two levels:
• Alere Triage Total 5 Control 1
• Alere Triage Total 5 Control 2

Store the controls frozen at -20°C or colder in a non-defrosting freezer. Do not store near the freezer door. The reagents are stable until the date on the box. The reagents should not be refrozen and it is recommended that each tube be used once and discarded. Frozen control material must be brought to room temperature (at least 30 minutes) prior to use.

Preparing the control solution
• Remove the control solution from the freezer and allow to thaw and warm to room temperature (19°-25°C) for at least 30 minutes.
  **NOTE:** Use within 1 hour of removal from the freezer.
• Mix thoroughly by inverting the control solution tube

Preparing the meter
• Power the meter on by pressing the ‘Power’ key. Press the ‘Enter’ key to run self-test.
• Select ‘Run Test’ using the up/down arrow keys and press
• Type in your identification number (User ID) if prompted.
• Select ‘QC Sample’ using the up/down arrow keys and press the ‘Enter’ key to confirm selection
• Enter the QC lot number from the label on the side of the vial containing the QC Sample and press the ‘Enter’ key to confirm the number.
  **NOTE:** Only enter the four digit numeric value of the QC lot number Do not enter the preceding alpha character
  **NOTE:** If you enter a QC lot number for which there is no data in the meter’s memory, you will see a message on the meter’s screen: ‘No QC Sample Data in Memory’. Press ‘Enter’ to continue and install Code Chip module.
Applying the control sample

**NOTE:** The unopened Test Devices should sit at room temperature for at least 15 minutes before performing a test.

- Open the pouch and label the Test Device.
- Place the Test Device on a level, horizontal surface.
- Hold the tube with the tip facing upwards, ensuring that all material is at the bottom of the tube.
- Snap off the tab
- Turn the tube over and dispense the entire contents into the sample port of the Test Device.
- Ensure the specimen has been absorbed completely before moving the Test Device.

Inserting the Test Device

- Hold the Test Device by the edges and gently insert the Test Device into the meter until you feel the Test Device catch on the pin and hear a ‘click’.
  **NOTE:** After the test device has been inserted, do not push the device in further or attempt to pull it out. The device may be ejected by returning to the Main Menu and pressing ‘Exit’.
- Press ‘Enter’ to start the test
- The meter pulls in the test device and scans it
  **NOTE:** The Test Device must be inserted into the Meter within 30 minutes from the time the sample was added

Results

**NOTE:** Do not move the meter while a test is in progress.

- The results are displayed on the screen, and automatically stored in the meter’s memory.
- Results should fall within the expected ranges provided on the Expected Values card provided with the D-Dimer Controls Kit
If any of the QC Sample results are out of range, the results for that particular analyte will be displayed as light text on a dark background.

9.5 Laboratory Comparison
If your policy states you must perform laboratory comparison then perform a venepuncture sample for the laboratory and store in an EDTA blood tube. Perform venepuncture and run sample on the Alere Triage MeterPro and send a specimen to the lab for Troponin I testing. Collection of both samples should occur at the same time. Record and compare the results ensuring they are in acceptable range for your organization.

10. TEST PROCEDURE

To change the meter settings, insert the Supervisor CODE CHIP module and refer to the Alere Triage MeterPro User Manual for instructions.

NOTE: Prior to commencing patient testing, remove the Supervisor CODE CHIP module from the meter, return it to the storage box and place in a safe place for future use.

10.1 Performing the Test

The Reagent Code Chip Module should be performed before running a patient test, as stated in section 9.2

Preparing the meter

- Power the meter on by pressing the ‘Power’ key. Press the ‘Enter’ key to run self-test.
- Select ‘Run Test’ using the up/down arrow keys and press the ‘Enter’ key to confirm.
- Type in your identification number (User ID) if prompted.
- Select ‘Patient Sample’ using the up/down arrow keys and press the ‘Enter’ key to confirm selection.
- Type or (scan) the patient’s identification (Patient ID) then press the ‘Enter’ key to confirm.

NOTE: To correct the patient ID, press the ‘Delete’ key to clear the entire ID or select the incorrect character using the keys. Then type in the correct ID.

Applying the blood sample

- Open the pouch and label the Test Device with the patient identification number.
- Place the Test Device on a level, horizontal surface.
- Using the transfer pipette, squeeze the larger (top) bulb completely and insert the tip into the specimen.

- Release the bulb slowly. The transfer pipette barrel should fill completely with some fluid flowing into the smaller (lower) bulb.
- Place the tip of the transfer pipette into the sample port of the Test Device and squeeze the larger bulb completely. The entire volume of fluid in the transfer pipette barrel must flow into the sample port. The specimen in the smaller (lower) bulb will not be expelled.
- Remove the transfer pipette tip from the sample port and then release the larger (top) bulb.
- Discard the transfer pipette.
- Ensure the specimen has been absorbed completely before moving the Test Device.

**NOTE:** The Test Device must be inserted into the Meter within 30 minutes from the time the patient specimen was added

**Inserting the Test Device**
- Hold the Test Device by the edges and gently insert the Test Device into the meter until you feel the Test Device catch on the pin and hear a ‘click’ and press ‘Enter’.

**NOTE:** After the test device has been inserted, do not push the device in further or attempt to pull it out. The device may be ejected by returning to the Main Menu and pressing ‘Exit’.
- After the test device has been pulled into the meter, the meter will display a tick mark indicates tests which are selected and the absence of a tick mark indicates which tests have not been selected.
- The user will be prompted to select desired analyte(s) or deselect undesired analyte(s) by pressing the number to the left of the test to toggle between selected or deselected and press the ‘Enter’ key to confirm selection
- Press the ‘Enter’ key to start the test.

**NOTE:** If no key is pressed on the Select Tests screen, the meter waits 30 seconds then proceeds using only the selected default tests. Then, if no key is pressed on the Confirmation Screen, the test proceeds after waiting 30 seconds.

**Results**

**NOTE:** Do not move the meter while a test is in progress.
• The results are displayed on the screen, and save to the meter’s memory automatically.
• Results displayed below the cut off levels will have no box around the result. Results in the indeterminate range have a black border around the result. Abnormal results are in a filled black box.
• Results can be printed by pressing the ‘Print’ button.

**NOTE:** A panel with multiple analytes will still be able to report patient results on those analytes which passed QC. An analyte that failed QC will not be reported on patient tests (instead will be flagged with a # symbol).

11. RESULTS

11.1 Interpretation of results
Collectively, the diagnosis of myocardial infarction should include measurement of cardiac related proteins and other clinical information including patient history and electrocardiographic data. Other conditions that may result in elevated cardiac proteins are: cardiac contusions, myocarditis, invasive examination of the heart, coronary artery bypass surgery, congestive heart failure and unstable angina. Therefore, these data must be considered when interpreting the results. TnI is platform dependent so each site should establish a reference range that is representative of the patient population to be evaluated.

11.2 Transferring Data to a printer or computer
• To review previous results, turn on the meter by pressing the ‘Power’ button
• Select ‘Recall Results’ using the up/down arrow keys and press the ‘Enter’ button to confirm.
• Enter your User ID if prompted and press ‘Enter’
• Select ‘Panel Type’ using the up/down arrow keys
• Use the left/right arrow keys to select the desired panel.
• Select the results based on Patient ID, Test Device Lot Number (L/N), User ID, from date, till date.
• Use the left/right arrow keys view the results.
• Press the ‘Print’ key to make a printed copy or press the ‘Enter’ key to upload the selected results to LIS.

12. MAINTENANCE
• Clean the exterior using a damp sponge or cloth with mild soap and water solution
• Wipe the meter dry using a soft cloth or absorbent tissue.
• Do not allow water to seep into the printer.
- Do not immerse the meter in water or other liquids.
- If blood or other fluids are not allowed enough time to fully absorb into test devices, the device track door may occasionally require cleaning. Using a cotton swab dampened with isopropyl alcohol and a pair of tweezers, carefully lift the door and clean both front and back of the door.

Changing Paper
- Tear off any excess paper sticking out of the Meter
- Remove the paper compartment cover by pulling up on the cover as indicate by the arrow on the back of the cover.
- Remove unused paper or the empty paper spindle from paper compartment.
- Cut a clean, straight edge to feed into the printer. Do not cut paper at an angle, as the printer must sense the edge of the paper along the feed path.
- Insert the new roll of paper into the paper compartment.
- Position the paper such that the paper will feed from under the roll (as opposed to over the top of the roll)
- Insert the paper edge under the paper roller (platen) until it firmly seats or resistance is felt.
- Press the ‘Print’ key.
- Replace the cover of the printer and continue operation.

13. REFERENCES

This method has been adapted from the Alere Triage MeterPro System User Manual, Test Device and control solution package inserts.


