

18. TROUBLE SHOOTING CHART

| PROBLEM                             | POSSIBLE CAUSE                                                                      | SOLUTION                                                                                                                  |
|-------------------------------------|-------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------|
| 1. Controls out of validation limit | a) Controls/ calibrator deterioration due to improper storage or used               | Use controls/ calibrator within 30 days once opened and Check storage after expiry. temp. It should be 2-8°C.             |
|                                     | b) Cross contamination                                                              | Pipette carefully and do not of Controls interchange caps.                                                                |
|                                     | c) Reagents deterioration due to improper storage or used after expiry.             | Use reagents within 30 days once opened and Check storage temp. It should be 2-8°C.                                       |
|                                     | d) Magnetic microsphere are not properly mixed before loading in the analyzer.      | Ensure proper mixing of bottle containing microspheres by gentle shaking/ inversion before use.                           |
| 2) High Myoglobin test results      | a) Use of turbid, lipaemic or hemolyzed sample.                                     | Use clear fresh sample. Refer specimen collection, handling and processing for more details.                              |
|                                     | b) Sample position is wrongly defined while loading the sample details in analyzer. | check the sample position and run the test meticulously.                                                                  |
|                                     | c) Magnetic microsphere are not properly mixed before loading in the analyzer.      | Ensure proper mixing of bottle containing microspheres by gentle shaking/ inversion before use.                           |
|                                     | d) Wrong Sample identification                                                      | Make sample I.D. at the time of sample Collection.                                                                        |
| 3) Low Myoglobin test results       | a)Sample deterioration due to improper Storage or microbially contaminated sample.  | Use clear fresh sample immediately after collection. Refer Specimen collection, and handling processing for more details. |
|                                     | b) Sample position is wrongly defined while loading the sample details in analyzer. | check the sample position and run the test meticulously.                                                                  |
|                                     | c) Magnetic microsphere are not properly mixed before loading in the analyzer.      | Ensure proper mixing of bottle containing microspheres by gentle shaking/ inversion before use.                           |
|                                     | d) Wrong Sample identification                                                      | Make sample I.D. at the time of sample Collection.                                                                        |

*in vitro* diagnostic Reagent, not for medicinal use

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# MYOGLOBIN iClia

*Chemiluminescence microparticle immunoassay for Quantitative Measurement of Myoglobin in Human Serum/Plasma*

1. INTRODUCTION

Myoglobin is a type of ferrous hemoglobin that exists in the heart and striated muscle of the human body. It is mainly responsible for the storage and transportation of oxygen in the muscle cells. When strenuous exercise, skeletal muscle injury or myocardial injury, the level of serum MYO increases, so it plays a certain role in the early diagnosis of patients with acute muscle injury and myocardial infarction. In the early stage of myocardial infarction, the rapid and sensitive detection of myoglobin is an important supplement to the electrocardiogram. Commonly used clinical detection methods include colloidal gold method, immunoturbidimetric method, chemiluminescence immunoassay (CLIA), etc.

2. INTENDED USE

Myoglobin iClia Kit is intended for the *in vitro* quantitative measurement of Myoglobin in human serum/plasma as an aid in the diagnosis of Myocardial Infarction in conjunction with other laboratory and clinical findings. This kit is only operational in conjunction with J.Mitra CLIA 181 Analyzer.

3. PRINCIPLE

Myoglobin iClia Kit is chemiluminescent immunoassay based on the “Sandwich” principle. The magnetic microspheres are coated with Anti-Myoglobin antibodies. The samples are added in the assay cup containing Assay buffer , coated microspheres followed by addition of AE conjugate (Anti- Myoglobin antibodies linked to acridinium ester). A sandwich complex is formed wherein Myoglobin (from serum sample) is “trapped” or “sandwiched” between the microspheres coated antibody and antibody labelled with AE conjugate. Unbound conjugate is then washed off with wash buffer. The amount of bound AE conjugate is proportional to the concentration of Myoglobin present in the sample. Finally pre-trigger and trigger solution containing hydrogen peroxide and sodium hydroxide solution is added to the reaction mixture. The resulting chemiluminescent reaction is measured as relative Light units (RLUs). There is a direct relationship between the amount of Myoglobin present in the sample and the RLUs detected by the optical system. Results are calculated automatically based on the established calibration curve and Myoglobin concentration in the sample expressed as ng/ mL.

4. DESCRIPTION OF SYMBOLS USED

The following are graphical symbols used in or found on J. Mitra diagnostic products and packing. These symbols are the most common ones appearing on medical devices and their packing. They are explained in more detail in the European Standard EN ISO 15223-1:2021.

|  |                                              |  |                                               |
|--|----------------------------------------------|--|-----------------------------------------------|
|  | Manufactured By                              |  | <i>In vitro</i> diagnostic medical device     |
|  | No. of tests<br>Lot Number                   |  | Instruction for use                           |
|  | Batch Number                                 |  | Temperature Limitation                        |
|  | Manufacturing Date                           |  | Caution - See instruction for use             |
|  | Expiry Date                                  |  | Catalogue Number                              |
|  | Do not use if package is damaged             |  | Keep away from sunlight                       |
|  | Contains biological Material of Human Origin |  | Contains biological Material of Animal Origin |
|  | Country of Manufacture                       |  | Keep Dry                                      |

5. KIT PRESENTATION

- 25 Test Pack
- 50 Test Pack
- 100 Test Pack

6. KIT & ITS COMPONENTS

| COMPONENT                   | DESCRIPTION                                                                     |
|-----------------------------|---------------------------------------------------------------------------------|
| <b>Microparticle Buffer</b> | Magnetic Microspheres coated with anti-Myoglobin antibodies with preservatives. |
| <b>Assay Buffer</b>         | Tris buffer and BSA with preservative.                                          |
| <b>AE Conjugate</b>         | anti-Myoglobin antibodies linked to acridinium ester with Protein stabilizers.  |
| <b>Control-1</b>            | Freeze-dried powder contain Myoglobin antigen.                                  |
| <b>Control-2</b>            | Freeze-dried powder contain Myoglobin antigen.                                  |
| <b>Calibrator-1</b>         | Freeze-dried powder contain Myoglobin antigen.                                  |
| <b>Calibrator-2</b>         | Freeze-dried powder contain Myoglobin antigen.                                  |
| <b>Reagent Plugs</b>        | Silicon caps to cover the opened reagents.                                      |

7. STORAGE AND STABILITY

The shelf-life of the kit is 12 months from the date of manufacturing, when stored at 2-8°C. **Once the kit is opened, onboard stability of reagents, calibrator and control is 30 days at 2-8°C.**

8. ADDITIONAL MATERIAL AND INSTRUMENTS REQUIRED

- **Pre-Trigger Solution:** Hydrogen peroxide solution.
- **Trigger Solution:** Sodium hydroxide solution.
- **Wash Buffer:** Phosphate buffered saline solution with surfactant.
- **Assay Cup**
- **Sample Diluent**
- **J. Mitra CLIA Analyzer**

*All materials and analyzer to be used for running the Myoglobin iClia shall be from J. Mitra & Co. Pvt. Ltd.*

9. SPECIMEN COLLECTION & HANDLING

1. Only human serum or plasma samples should be used for the test.
2. For serum collection use serum vacutainer. While preparing serum samples, remove the serum from the clot as soon as possible to avoid hemolysis. Fresh serum/plasma samples are preferred.
3. For plasma collection: use Dipotassium EDTA, Tripotassium EDTA, Sodium heparin and lithium heparin gel vacutainer.
4. Specimens should be free of microbial contamination and may be stored at 2-8°C for one week, or frozen at -20°C or lower. Avoid repeated freezing and thawing.
5. Do not use heat inactivated samples as their use may give false results. Hemolyzed and Icteric hyperlipemic samples may give erroneous results.
6. Serum specimens from patients receiving anticoagulant or thrombolytic therapy may contain fibrin due to incomplete clot formation.
7. Always use clear specimens. Centrifuge viscous/ thick or turbid specimen at 10,000 RPM for 15 minutes prior to use to avoid inconsistent result.
8. Use of disposable pipettes or pipette tips is recommended to prevent cross contamination.

10. SPECIMEN PROCESSING

(A) FROZEN SAMPLE

Myoglobin iClia test is best used with fresh samples that have not been frozen and thawed. However most frozen samples will perform well if the procedure suggested below is followed.

Allow the frozen sample to thaw in a vertical position in the rack. Do not shake the sample. This allows particles to settle to the bottom. Centrifuge the sample at 10,000 rpm for 15 minutes.

**(B) TRANSPORTATION**

If the specimen is to be transported, it should be packed in compliance with the current Government regulations regarding transport of aetiologic agents.

**11. WARNING & PRECAUTION**

 **CAUTION:** THIS KIT CONTAINS MATERIALS OF HUMAN ORIGIN. NO TEST METHOD CAN OFFER COMPLETE ASSURANCE THAT HUMAN BLOOD PRODUCTS WILL NOT TRANSMIT INFECTION. NEGATIVE CONTROL, POSITIVE CONTROL & ALL THE SAMPLES TO BE TESTED SHOULD BE HANDLED AS THOUGH CAPABLE OF TRANSMITTING INFECTION.

1. The use of disposable gloves and proper biohazardous clothing is STRONGLY RECOMMENDED while running the test.
2. In case there is a cut or wound in hand, DO NOT PERFORM THE TEST.
3. Do not smoke, drink or eat in areas where specimens or kit reagents are being handled.
4. Tests are for *in vitro* diagnostic use only and should be run by competent person only.
5. Do not pipette by mouth.
6. All materials used in the assay and samples should be decontaminated in 5% sodium hypochlorite solution for 30-60 min. before disposal or by autoclaving at 121°C at 15psi for 60 minutes. Do not autoclave materials or solution containing sodium hypochlorite. They should be disposed off in accordance with established safety procedures.
7. Wash hands thoroughly with soap or any suitable detergent, after the use of the kit. Consult a physician immediately in case of accident or contact with eyes, in the event that contaminated material are ingested or come in contact with skin puncture or wounds.
8. Spills should be decontaminated promptly with Sodium Hypochlorite or any other suitable disinfectant.

**12. PRECAUTIONS FOR USE & REAGENT HANDLING**

1. Do not use kit components beyond the expiration date which is printed on the kit.
2. Store the reagents & samples at 2-8°C.
3. Do not pool reagents from within a batch or between different batches, as they are optimised for individual batch to give best results.
4. Before loading the reagent kit in the clia analyzer for the first time, ensure proper mixing of microparticle bottle to resuspend microparticles that may have settled during transport or storage.
5. Once reagents are opened, reagent plug must be used to prevent reagent evaporation and contamination and to ensure reagent integrity. Reliability of assay results cannot be guaranteed if reagent plugss are not used according to the instructions given.
6. Mark the test specimen with patient's name or identification number. Improper identification may lead to wrong result reporting.
7. To avoid contamination, wear clean gloves when placing a reagent plug on an uncapped reagent bottle.
8. Once a reagent plug has been placed on an open reagent bottle, do not invert the bottle as this will result in reagent leakage and may compromise assay results.
9. Reagents may be stored on or off the Chemiluminescence immunoassay analyzer. If reagents are removed from the analyzer, store them at 2-8°C (with Reagent plugs) in an upright position. For reagents stored off the system, it is recommended that they should be stored in their original trays and boxes to ensure they remain upright. If the microparticle bottle does not remain upright (with a Reagent plugs placed) while in refrigerated storage off the system, the reagent kit must be discarded.
10. Run Myoglobin Control-1 & Myoglobin Control-2 in each assay to evaluate validity of the kit.
11. Distilled or deionised water must be used for wash buffer preparation.

12. Avoid strong light exposure during the assay.
13. In case of any doubt the run should be repeated.

**13. TEST PROCEDURE**

**Reconstitution of Controls and calibrators:**

- i) Use 3 ml of antigen diluent to dissolve lyophilized antigen well and desolve completely. Put the cap and let it stand for 10 minutes. Mix solution thoroughly before use. The working antigen is stable for 7 days at 2-8°C and 2 months at -20°C (only 2 freeze thaw of liquid antigen are allowed at -20°C).
- ii) Transfer reconstituted antigen into the RC bottle by opening soft purple cap. After transferring reconstituted antigen and place the purple plug before use.

**Assay Procedure**

1. Refer to the Clia-181 user manual for detailed information on preparing the analyzer.
2. Before loading the Myoglobin iClia reagent tray on the analyzer for the first time, mix contents of the microparticle buffer bottle to resuspend microparticles that may have settled during transporation/ storage. Once the microparticles have been loaded, no further mixing is required.

**Important Note: Swirl the microparticle (RA) bottle 30 times. Visually inspect the bottle to ensure microspheres are resuspended. If microspheres are still adhered to the bottle, continue to Swirl the bottle until the microspheres have been completely resuspended. If the microspheres do not resuspend, DO NOT USE. Once the microspheres have been resuspended, remove the cap and place the reagent plug on the bottle to make it ready to use. Remove the cap of (RA), (RB), (RC) and (RD) bottles and place the reagent plugs before use as follow:**

- |                        |          |                           |
|------------------------|----------|---------------------------|
| <b>(RA) &amp; (RB)</b> | <b>:</b> | <b>Natural color plug</b> |
| <b>(RC)</b>            | <b>:</b> | <b>Purple color plug</b>  |
| <b>(RD)</b>            | <b>:</b> | <b>Brown color plug</b>   |
3. Load the Myoglobin iClia reagent tray on the Chemiluminescence immunoassay analyzer.
  4. Verify that all necessary reagents are available in the reagent tray.
  5. Ensure that adequate sample volume (not less than 250 µL) is present in sample tube prior to running the test.
  6. Sample volume required for each additional test from same sample tube is 15 µL.
  7. The Myoglobin test-specific parameters are stored in reagent barcode placed on the reagent tray and read through barcode reader. In cases, the barcode cannot be read, contact customer support at: 011-47130300, 500 or write us at: jmitra@jmitra.co.in.
  8. Run calibration, if required.
  9. Mix Myoglobin iClia calibrators and controls by gentle inversion before use. Open the the cap and place the Myoglobin Calibrator-1 & Myoglobin Calibrator-2 and Myoglobin Control-1 & Myoglobin Control-2 vials into each respective sample positions. Read the barcode for calibrator and controls provided with the kit.
  10. Press START. The test result for first sample will be obtained at 15 minutes.
  11. The Chemiluminescence immunoassay analyzer performs all the functions automatically and calculate the results.

**Calibration**

1. Traceability: This assay has been standardized against the Roche Myoglobin reagent kit.
2. Every Myoglobin iClia kit has a two-dimension code label containing the predefined master curve of the particular reagent lot.
3. Test all 2 Calibrators in triplicate. Both Myoglobin Control-1 and Myoglobin Control-2 must be tested in each run to evaluate the assay calibration. Ensure that controls values are within the validity range specified in the Myoglobin iClia QC data sheet given in the Clia Analyzer.
4. Once calibration is accepted (within range) and stored, all subsequent samples may be tested without further calibration unless:

5. Recalibrate the analyzer in following conditions:
  - a) After each exchange/use of new lot (Test reagent and pritrigger/ Trigger solution/wash buffer).
  - b) Every 15 days at the time of any component to be changed.
  - c) Controls are out of validation range.
  - d) Required by pertinent regulations.
  - e) After specified service procedures have been performed or maintenance to critical part or subsystems that might influence the performance of the Myoglobin iClia.

**Result Calculation**

The analyzer automatically calculates the concentration of each sample. The results are given in ng/ml.

**Result Interpretation**

If sample concentration is lower than the lower limit of the linear range, report the result <5.00 ng/mL, while > 1000.00 ng/mL when it is higher than the upper limit of linear range.

**Determination of Reference Interval**

Reference Interval of this assay is considered as <115 ng/mL for healthy people, which is established referring to literatures, based on the rest results of more than 60 clinical samples.

Due to the differences in geography, race, gender or age, it is suggested each laboratory establish its own reference interval or conduct verification of the existing reference interval.

**14. PERFORMANCE CHARACTERISTICS**

- Assay results obtained in individual laboratories may vary from data presented in this product insert.

**Limit of Blank (LoB)**

- The Limit of Blank was determined in accordance with the CLSI (Clinical and Laboratory Standards Institute) EP17-A requirements.
- The Limit of Blank is the 95th percentile value from n >20 measurements of analyte free samples over several independent series. The Limit of Blank corresponds to the concentration below which analyte-free samples are found with a probability of 95%.
- The observed LoB value was ≤2.000 ng/ml.

**Accuracy:** The accuracy of Myoglobin iClia was detected with 60 clinical specimens and compared with Roche CLIA. The co-relation co-efficient is >0.95.

**Precision**

**Intra Assay Variation**

Within run variation was determined by 10 replicate measurements of two different Myoglobin control sera( Low) and (High) in one assay in 3 different lots. The within assay variability is <10%.

**Inter Assay Variation**

Between run variation was determined by 10 replicate measurements in 10 sequential days of two different control sera( Low ) and (High) in 3 different lots. The between assay variability is <10%.

| Intra-Assay, n=10 |              |       | Inter-Assay, n=10×2 |              |       |
|-------------------|--------------|-------|---------------------|--------------|-------|
| Control           | Mean (ng/ml) | CV    | Sample              | Mean (ng/ml) | CV    |
| 1                 | 0.49         | 4.37% | 1                   | 0.52         | 6.19% |
| 2                 | 22.70        | 3.80% | 2                   | 22.84        | 7.56% |

**Inter machine (CLIA-181 Analyzer) Variation**

Between machine variation was determined by 3 replicate measurements of two different Myoglobin control sera( Low ) and (High) in 3 different lots in 3 different CLIA-181 Analyzer. The between machine variability is <10.0 %.

**Linearity**

The linearity was determined in accordance with the CLSI (Clinical and Laboratory Standards Institute) EP6-A requirements.

The linearity range was verified by more than 6 concentration levels which encompass or be equal to the minimum and the maximum values of linearity range and duplicate assays in triplicate in single run for each lot at all 6 levels.

The Myoglobin iClia kit has been demonstrated to be linear from is 5 ng/ml to 100 ng/ml, regression (R<sup>2</sup>) of more than ≥0.990.

**Interference**

A study was performed based on guidance from CLSI EP7-A2.

Potentially interfering substances were evaluated to determine whether Myoglobin concentrations were affected when using the Myoglobin iClia kit. Samples containing the potential interferents were prepared at two Myoglobin concentrations. The samples were assayed, and the Myoglobin concentrations of the spiked samples were compared to the reference samples.

| Potential Interferent | Interferent Concentration | % Interferent Bias |
|-----------------------|---------------------------|--------------------|
| Bilirubin             | 20 mg/dL                  | <10%               |
| Hb                    | 500 mg/dL                 | <10%               |
| Triglyceride          | 1000 mg/dL                | <10%               |
| Total protein         | 10 g/dL                   | <10%               |
| RF                    | 1000IU/mL                 | < 10%              |
| ANA                   | 400AU/mL                  | < 10%              |
| HAMA                  | 600ng/mL                  | < 10%              |

**15. LIMITATION OF THE TEST**

- The Myoglobin iCia should be used for detection of Myoglobin in serum or plasma only and not in other body fluids.
- Results should be used in conjunction with other data; e.g., symptoms, results of other tests, and clinical impressions. If the Myoglobin results are inconsistent with clinical evidence, additional testing is recommended.
- Clinical diagnosis should not be made on the findings of a single test result, but should be integrated with all clinical and laboratory findings.
- Samples of lipid, hemolysis or jaundice may result in incorrect results. Hemoglobin (150 mg/dL), triglyceride (1000 mg/dL), or bilirubin (40 mg/dL) will have no significant interference for the results.

**16. LIMITED EXPRESSED WARRANTY DISCLAIMER**

The manufacturer limits the warranty to the test kit, as much as that the test kit will function as an *in vitro* diagnostic assay within the limitations and specifications as described in the product instruction-manual, when used strictly in accordance with the instructions contained therein. The manufacturer disclaims any warranty expressed or implied including such expressed or implied warranty with respect to merchantability, fitness for use or implied utility for any purpose. The manufacture's liability is limited to either replacement of the product or refund of the purchase price of the product and in no case liable to for claim of any kind for an amount greater than the purchase price of the goods in respect of which damages are likely to be claimed.

The manufacturer shall not be liable to the purchaser or third parties for any injury, damage or economic loss, howsoever caused by the product in the use or in the application there of.

**17. REFERENCES**

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