# **MAGLUMI IAA (CLIA)**





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CE

FOR PROFESSIONAL USE ONLY Store at 2...8 °C



COMPLETELY READ THE INSTRUCTIONS BEFORE PROCEEDING

# SYMBOLS EXPLANATIONS



Manufacturer

Authorized Representative in Europe

Contents of kit



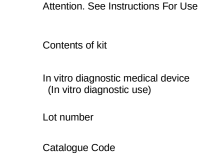
LOT

CONT

In vitro diagnostic medical device







Expiry date (Use by...)

Temperature limitation ( store at 2...8 °C)

Number of tests

Keep away from sunlight

Keep upright

# INTENDED USE

The kit has been designed for the quantitative determination of Human Insulin Autoantibodies (IAA) in human serum.

The method can be used for samples over the range of 0-175 II I/ml

The test has to be performed on the MAGLUMI chemiluminescence immunoassay (CLIA) fully auto analyzer (Including MAGLUMI 1000, MAGLUMI 2000, MAGLUMI 2000 Plus and new developed models).

# SUMMARY AND EXPLANATION OF THE TEST

Type 1 diabetes, commonly referred to as insulin-dependent diabetes (IDDM), is caused by pancreatic beta-cell destruction that leads to an absolute insulin deficiency. The clinical onset of diabetes does not occur until 80% to 90% of these cells have been destroyed. Prior to clinical onset, type 1 diabetes is often characterized by circulating autoantibodies against a variety of islet cell antigens, including glutamic acid decarboxylase (GAD), tyrosine phosphatase (IA2), and insulin. The autoimmune destruction of the insulin-producing pancreatic beta cells is thought to be the primary cause of type 1 diabetes. The presence of these autoantibodies provides early evidence of autoimmune disease activity, and their measurement can be useful in assisting the physician with the prediction, diagnosis, and management of patients with diabetes. Insulin is the only beta-cell specific autoantigen thus far identified. Antibodies to insulin are found predominantly, though not exclusively, in young children developing type 1 diabetes. In insulin-naive (untreated) patients, the prevalence of antibodies to insulin is almost 100% in very voung individuals and almost absent in adult onset of type 1 diabetes. Because the risk of diabetes is increased with the presence of each additional autoantibody marker, the positive predictive value of insulin antibody measurement is increased when measured in conjunction with antibodies to GAD and IA-2.

# **PRINCIPLE OF THE TEST**

Competition sandwich immunoluminometric assay:

Use an anti-Insulin monoclonal antibody to label FITC, and use another monoclonal antibody to label ABEI. The anti-Insulin in undiluted sample gets reaction with a certain amount of free Insulin antigen firstly, and then adds ABEI and FITC labels; compete to conjugate the binding sites with free INSULIN antigen. The amount of anti-Insulin in sample is inversely proportional to the amount of sandwich of ABEI, free Insulin and FITC. More anti-Insulin in sample, fewer sandwiches formed, and then less RLU value. Subsequently, the starter reagents are added and a flash chemiluminescent reaction is initiated. The light signal is measured by a photomultiplier as RLU within 3 seconds and is proportional to the concentration of anti- Insulin present in controls or samples.



**KIT COMPONENTS** Material Supplies

••		
Reagent Integral for 100 determinations		
Nano magnetic microbeads: TRIS buffer,		
1.2% (W/V), 0.2%NaN <sub>3</sub> , coat sheep anti- FITC	2.5ml	
polyclonal antibody.		
Calibrator Low: bovine serum, 0.2%NaN <sub>3</sub> .	2.5ml	
Calibrator High: bovine serum, 0.2%NaN <sub>3</sub> .	2.5ml	
FITC Label: Anti-Insulin monoclonal antibody		
labeled FITC contains BSA, 0.2%NaN <sub>3</sub> .	10.5ml	
ABEI Label: Anti-Insulin monoclonal antibody		
labeled ABEI contains BSA, 0.2%NaN <sub>3</sub> .	10.5ml	

Free antigen: contains purified INSULIN antigen, 0.2% NaN <sub>3</sub>	6.5ml	
All reagents are provided ready-to-use		

Reagent Vials in kit box		
Internal Quality Control: containing BSA,		
0.2%NaN <sub>3</sub> . (target value refer to Quality	2.0ml	
Control Information date sheet)		

#### Accessories Required But Not Provided

MAGLUMI Reaction Module	REF: 630003
MAGLUMI Starter 1+2	REF: 130299004M
MAGLUMI Wash Concentrate	REF: 130299005M
MAGLUMI Light Check	REF: 130299006M



# Preparation of the Reagent Integral

Before the sealing is removed, gentle and careful horizontal shaking of the Reagent Integral is essential (avoid foam formation!) Remove the sealing and turn the small wheel of the magnetic microbeads compartment to and fro, until the colour of the suspension has changed into brown. Place the Integral into the reagent area and let it stand there for 30 min. During this time, the magnetic microbeads are automatically agitated and completely resuspended.

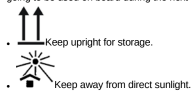
Do not interchange integral component from different reagents or lots!

### Storage and Stability

Sealed: Stored at 2-8

• Opened: Stable for 4 weeks. To ensure the best kit performance, it is recommended to place opened kits in the refrigerator if it's not going to be used on board during the next 12 hours.

°C until the expiry date.



# CALIBRATION AND TRACEABILITY

#### 1)Traceability

To perform an accurate calibration, we have provided the test calibrators standardized against the SNIBE internal reference substance

Calibrators in the Reagent Kit are from Sigma.

#### 2) 2-Point Recalibration

Via the measurement of calibrators, the predefined master curve is adjusted (recalibrated) to a new, instrument-specific measurement level with each calibration.

#### 3) Frequency of Recalibration

- After each exchange of lot (Reagent Integral or Starter Reagents).
- Every 2 weeks and/or each time a new Integral is used (recommendation).
- After each servicing of the MAGLUMI Fully Auto analyzer.
- If controls are beyond the expected range.

# SPECIMEN COLLECTION AND PREPARATION

#### Sample material: serum

Collect samples using standard procedures. Store at 2-8

below - 20 °C Avoid repeated freezing and thawing cycles, stored samples should be thoroughly mixed prior to use (Vortex mixer). 044120530-v1.0-EN Please ask local representative of SNIBE for more details if you have any doubt.

#### Vacuum Tubes

(a) Blank tubes are recommended type for collecting samples.(b) Please ask SNIBE for advice if special additive must be used in sample collecting.

## Specimen Conditions

- Do not use specimens with the following conditions:
- (a) heat-inactivated specimens;
- (b) Cadaver specimens or body fluids other than human serum;(c) Obvious microbial contamination.
- Use caution when handling patient specimens to prevent cross contamination. Use of disposable pipettes or pipette tips is recommended.
- Inspect all samples for bubbles. Remove bubbles with an applicator stick prior to analysis. Use a new applicator stick for each sample to prevent cross contamination.
- Serum specimens should be free of fibrin, red blood cells or other particulate matter.
- Ensure that complete clot formation in serum specimens has taken place prior to centrifugation. Some specimens, especially those from patients receiving anticoagulant or thrombolytic therapy, may exhibit increased clotting time. If the specimen is centrifuged before a complete clot forms, the presence of fibrin may cause erroneous results.

#### **Preparation for Analysis**

- Patient specimens with a cloudy or turbid appearance must be centrifuged prior to testing. Following centrifugation, avoid the lipid layer (if present) when pipetting the specimen into a sample cup or secondary tube.
- Specimens must be mixed thoroughly after thawing by low speed vortexing or by gently inverting, and centrifuged prior to use to remove red blood cells or particulate matter to ensure consistency in the results. Multiple freeze-thaw cycles of specimens should be avoided.
- All samples (patient specimens or controls) should be tested within 3 hours of being placed on board the MAGLUMI System. Refer to the SNIBE service for a more detailed discussion of onboard sample storage constraints.

#### Storage

- If testing will be delayed for more than 8 hours, remove serum or plasma from the serum or plasma separator, red blood cells or clot. Specimens removed from the separator gel, cells or clot may be stored up to 24 hours at 2-8°C.
- Specimens can be stored up to 30 days frozen at -20°C or colder.

#### Shipping

 Before shipping specimens, it is recommended that specimens be removed from the serum or plasma separator, red blood cells or clot. When shipped, specimens must be packaged and labeled in compliance with applicable state, federal and international regulations covering the transport of clinical specimens and infectious substances. Specimens must be shipped frozen (dry ice). Do not exceed the storage time limitations identified in this section of the package insert.

## WARNING AND PRECAUTIONS FOR USERS



• For use in *IN-VITRO* diagnostic procedures only.

°C: 24 hour a state of the instruction of the instructions in this package insert.

#### Safety Precautions

**CAUTION:** This product requires the handling of human specimens.

- The calibrators in this kit are prepared from bovine serum products. However, because no test method can offer complete assurance that HIV, Hepatitis B Virus or other infectious agents are absent; these reagents should be considered a potential biohazard and handled with the same precautions as applied to any serum or plasma specimen.
- All samples, biological reagents and materials used in the assay must be considered potentially able to transmit infectious agents. They should therefore be disposed of in accordance with the prevailing regulations and guidelines of the agencies holding jurisdiction over the laboratory, and the regulations of each country. Disposable materials must be incinerated; liquid waste must be decontaminated with sodium hypochlorite at a final concentration of 5% for at least half an hour. Any materials to be reused must be autoclaved using an overkill approach (USP 24, 2000, p.2143). A minimum of one hour at 121 considered adequate, though the users must check the effectiveness of their decontamination cycle by initially validating it and routinely using biological indicators.
- It is recommended that all human sourced materials be considered potentially infectious and handled in accordance with the OSHA Standard on Bloodborne Pathogens13. Biosafety Level 214 or other appropriate biosafety practices should be used for materials that contain or are suspected of containing infectious agents.
- This product contains Sodium Azide; this material and its container must be disposed of in a safe way.
- Safety data sheets are available on request.

#### Handling Precautions

• Do not use reagent kits beyond the expiration date.

- Do not mix reagents from different reagent kits.
- Prior to loading the Reagent Kit on the system for the first time, the microbeads requires mixing to re-suspend microbeads that have settled during shipment.
- For microbeads mixing instructions, refer to the KIT COMPONENTS, Preparation of the Reagent Integral section of this package insert.
- To avoid contamination, wear clean gloves when operating with a reagent kit and sample.
- Over time, residual liquids may dry on the kit surface, please pay attention the silicon film still exists on the surface of the kit.
- For a detailed discussion of handling precautions during system operation, refer to the SNIBE service information.

## TEST PROCEDURE

To ensure proper test performance, strictly adhere to the operating instructions of the MAGLUMI Fully Auto analyzer. Each test parameter is identified via a RFID tag on the Reagent Integral. For further information please refer to the MAGLUMI Chemiluminescence Analyzer Operating Instructions.

40µl	Sample, calibrator or controls
+40µl	Free antigen
15 min	Incubation
+80µl	ABEI
+80µl	FITC
+20µl	Nano magnetic microbeads
15 min	Incubation
400µl	Cycle washing
3 s	Measurement

## DILUTION

Sample dilution by analyzer is not available in this reagent kit Samples with concentrations above the measuring range can be diluted manually. After manual dilution, multiply the result by the dilution factor.

Please choose applicable diluents or ask SNIBE for advice before manual dilution must be processed.

## QUALITY CONTROL

Observe quality control guidelines for medical laboratories

Use suitable controls for in-house quality control. Controls should be run at least once every 24 hours when the test is in use, once per reagent kit and after every calibration. The control intervals should be adapted to each laboratory's individual requirements. Values obtained should fall within the defined ranges. Each laboratory should establish guidelines for corrective measures to be taken if values fall outside the range.

# LIMITATIONS OF THE PROCEDURE

## 1) Limitations

Patients with malignancies may exhibit IAA values within the °C is ngmaligange. IAA concentrations may be elevated in case of liver cirrhosis, hepatitis or tyrosinaemia. Thus, IAA determination is more suitable for therapeutic monitoring and follow-up as well as for a comparison with histological results. IAA serum levels may only be interpreted in context with the clinical picture and other diagnostic procedures. The IAA assay should not be used as the only criterion for cancer screening.

#### 2) Interfering Substances

No interference with test results is seen by concentrations of bilirubin <0.06mg/ml, haemoglobin <16mg/dl or triglycerides <12.5mg/ml.

## 3) HAMA

Patient samples containing human anti-mouse antibodies (HAMA) may give falsely elevated or decreased values. Although HAMA-neutralising agents are added, extremely high HAMA serum concentrations may occasionally influence results.

# RESULTS

#### 1) Calculation of Results

The analyzer automatically calculates the IAA concentration in each sample by means of a calibration curve which is generated by a 2-point calibration master curve procedure. The results are expressed in IU/ml. For further information please refer to the MAGLUMI Chemiluminescence Analyzer Operating Instructions.

#### 2) Interpretation of Results

- Results of study in clinical centers with group of individuals, 95% of the results were: < 20IU/ml.
- Results may differ between laboratories due to variations in population and test method. If necessary, each laboratory should establish its own reference range.

# PERFORMANCE CHARACTERISTICS

#### 1) Precision

Intra-assay coefficient of variation was evaluated on 3 different levels of control serum repeatedly measured 20 times in the same run, calculating the coefficient of variation.

Intra-assay precision			
Control	Mean(IU/ml)	SD(IU/ml)	CV%
Level 1	10.56	2.26	6.89
Level 2	20.44	3.21	5.92
Level 3	105.15	6.05	5.76
Inter access coefficient of variation was evaluated on three batches			

Inter-assay coefficient of variation was evaluated on three batches of kits. Repeatedly measured 3 different levels of control serum 21 times, calculating the coefficient of variation.

Inter-assay precision			
Control	Mean(IU/ml)	SD(IU/ml)	CV%
Level 1	11.02	2.17	9.74
Level 2	23.41	4.11	9.01
Level 3	113.37	9.54	8.42

## 2) Analytical Sensitivity

The sensitivity is defined as the concentration of IAA equivalent to the mean RLU of 20 replicates of the zero standard plus two standard deviations corresponding to the concentration from the standard curve. The sensitivity is typically less than 2 IU/ml.

## 3) Specificity

The specificity of the IAA assay system was assessed by measuring the apparent response of the assay to various potentially cross reactive analytes.

Compound	Concentration	Cross reactivity
T3	100 ng/ml	0.6%
rT3	100 ng/ml	0.6%

#### 4) Recovery

Consider calibrator high of known concentration as a sample, dilute it by 1:2 ratio with diluents, and measure its diluted concentration for 10 times. Then calculate the recovery of measured concentration and expected concentration. The recovery should be within 90% -110%.

Expected	Mean Measuring	Recovery
54.562IU/ml	56.349IU/ml	103%

#### 5) Linearity

Use IAA calibrator to prepare the six-point standard curve, measuring all points' RLU except point A, and then do four-parameter linear fitting in double logarithm coordinate, the absolute linear correlation coefficient (r) should be bigger than 0.9800.

Calibrator	Concentration	Absolute linear
Point	IU/ml	correlation coefficient (r)
A	0	
В	8	r=0.9811
С	20	
D	50	
E	100	
F	175	

#### 6) Method comparison

A comparison of MAGLUMI IAA (y) with a commercially available Insulin test (x) using clinical samples gave the following correlations (IU/mI):

Linear regression

y = 0.98x+13.4 r = 0.972 Sy.x =22.4

Number of samples measured: 260 The sample concentrations were between 9 and 157 IU/ml.

#### REFERENCES

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