

# Mercodia High Range Rat Insulin ELISA

Directions for Use

10-1145-01 Reagents for 96 determinations

For Research Use Only

Please note. The assigned
Please note. The assigned
Concentrations for the
Calibrators are lot-specific

Manufactured by

Mercodia AB Sylveniusgatan 8A SE-754 50 Uppsala Sweden

# Explanation of symbols used on labels

Σ = 96	Reagents for 96 determinations
$\subseteq$	Expiry date
	Store between 2–8°C
LOT	Lot No.

#### Intended Use

Mercodia High Range Rat Insulin ELISA provides a method for the quantitative determination of insulin in rat serum or plasma.

## Summary and explanation of the test

Insulin is the principal hormone responsible for the control of glucose metabolism. It is synthesised in the B-cells of the islets of Langerhans as the precursor, proinsulin, which is processed to form C-peptide and insulin. Both are secreted in equimolar amounts into the portal circulation. The mature insulin molecule comprises two polypeptide chains, the A chain and the B chain. The two chains are linked together by two inter-chain disulphide bridges. There is also an intrachain disulphide bridge in the A chain.

Secretion of insulin is mainly controlled by plasma glucose concentration, and the hormone has a number of important metabolic actions. Its principal function is to control the uptake and utilisation of glucose in peripheral tissues via the glucose transporter. This and other hypo-glycaemic activities, such as the inhibition of hepatic gluconeogenesis and glycogenolysis are counteracted by the hyperglycaemic hormones including glucagon, epinephrine (adrenaline), growth hormone and cortisol.

## Principle of the procedure

Mercodia High Range Rat Insulin ELISA is a solid phase two-site enzyme immunoassay. It is based on the direct sandwich technique in which two monoclonal antibodies are directed against separate antigenic determinants on the insulin molecule. During incubation insulin in the sample reacts with peroxidase-conjugated anti-insulin antibodies and anti-insulin antibodies bound to the microtitration well. A simple washing step removes unbound enzyme labeled antibody. The bound conjugate is detected by reaction with 3,3',5,5'-tetramethylbenzidine. The reaction is stopped by adding acid to give a colorimetric endpoint that is read spectrophotometrically.

## Warnings and precautions

- · For research use only.
- · All samples should be handled as capable of transmitting disease.
- · Each well can only be used once.
- The Stop Solution contains <5% Sulphuric acid.</li>

The Stop Solution is labeled:



#### Danger

H318 - Causes serious eve damage.

H315 - Causes skin irritation.

P280 - Wear protective gloves. Wear eye or face protection.

P264 - Wash hands thoroughly after handling.

P302 + P352 + P362 + P364 - IF ON SKIN: Wash with plenty of soap and

water. Take off contaminated clothing and wash it before reuse.

P332 + P313 – If skin irritation occurs: Get medical attention.

P305 + P351 + P338 + P310 – IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing. Immediately call a POISON CENTER or physician.

 The Enzyme Conjugate Buffer, Cal 0, 1, 2, 3, 4, 5 and Wash Buffer contain <0.06% 5-chloro-2-methyl-4-isothiazolin-3-one and 2-methyl-2H-isothiazol-3-one (3:1).

The Enzyme Conjugate Buffer, the Calibrators and Wash Buffer are labeled:



#### Warning

H317 - May cause an allergic skin reaction.

P280 - Wear protective gloves.

P261 - Avoid breathing vapour.

P272 – Contaminated work clothing should not be allowed out of the workplace.

P302 + P352 - IF ON SKIN: Wash with plenty of soap and water.

P333 + P313 - If skin irritation or rash occurs: Get medical attention.

P501 - Dispose of contents and container in accordance with all local,

regional, national and international regulations.

## Material required but not provided

- Pipettes with appropriate volumes (repeating pipettes preferred for addition of enzyme conjugate 1X solution, Substrate TMB and Stop Solution)
- · Tubes, beakers and cylinders for reagent preparation
- · Redistilled water
- Magnetic stirrer
- Vortex mixer
- · Microplate reader with 450 nm filter
- Microplate shaker (700-900 cycles per minute, orbital movement)
- Microplate washing device with overflow function (recommended but not required)

## Reagents

Each Mercodia High Range Rat Insulin ELISA kit (10-1145-01) contains reagents for 96 wells, sufficient for 42 samples and one calibrator curve in duplicate. For larger series of assays, use pooled reagents from packages bearing identical lot numbers. The expiry date for the complete kit is stated on the outer label. The recommended storage temperature is 2–8°C.

Coated Plate 1 plate 96 wel	Is Ready for Use
Mouse monoclonal anti-insulin 8-well For unused microplate strips, reseal the bag using adhes use within 2 months.	strips
Calibrators 1, 2, 3, 4, 5 5 vials 500 µ Rat insulin Color coded yellow Concentration stated on vial label	L Ready for Use
Calibrator 0 1 vial 5 mL Color coded yellow	Ready for Use
Enzyme Conjugate 11X 1 vial 600 μl Peroxidase conjugated mouse monoclonal anti-insulin	Preparation, see below
Enzyme Conjugate Buffer 1 vial 6 mL Color coded blue	Ready for use
Wash Buffer 21X 1 bottle 50 mL Storage after dilution: 2–8°C for 2 months.	Dilute with 1000 mL redistilled water to make wash buffer 1X solution.
Substrate TMB 1 bottle 22 mL Colorless solution Note! Light sensitive!	Ready for Use
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	Ready for Use

## Preparation of enzyme conjugate 1X solution

Prepare the needed volume of enzyme conjugate 1X solution by dilution of Enzyme Conjugate 11X (1+10) in Enzyme Conjugate Buffer according to the table below. When preparing enzyme conjugate 1X solution for the whole plate, pour all of the Enzyme Conjugate Buffer into the Enzyme Conjugate 11X vial. Mix gently.

Number of strips	Enzyme Conjugate 11X	Enzyme Conjugate Buffer
12 strips	1 vial	1 vial
6 strips	300 μL	3.0 mL
4 strips	200 μL	2.0 mL

Storage after dilution: 2-8°C for 2 months.

## Specimen collection and handling

#### Serum

Collect blood by venipuncture, allow to clot, and separate the serum by centrifugation. Samples can be stored at 2–8°C up to 24 hours. For longer periods store samples at -20°C. Avoid repeated freezing and thawing.

#### Plasma

Collect blood by venipuncture into tubes containing heparin or EDTA as anticoagulant, and separate the plasma fraction by centrifugation. Samples can be stored at 2–8°C up to 24 hours. For longer periods store samples at -20°C. Avoid repeated freezing and thawing.

## Preparation of samples

No dilution is normally required for serum and plasma samples, however, samples with a concentration above Calibrator 5 should be diluted in Calibrator 0 (or Mercodia Diabetes Sample Buffer, 10-1195-01).

## Test procedure

All reagents and samples must be brought to room temperature before use. Prepare a calibrator curve for each assay run. The product has been optimized and validated without plate sealer.

- Prepare enzyme conjugate 1X solution (according to the tables on previous pages), wash buffer 1X solution and samples.
- Prepare sufficient microplate wells to accommodate Calibrators, controls and samples in duplicate.
- 3. Pipette 10  $\mu$ L each of Calibrators, controls and samples into appropriate wells.
- 4. Add 50 µL of enzyme conjugate 1X solution into each well.
- 5. Incubate on a plate shaker (700–900 rpm) for 2 hours at room temperature (18–25°C).
- Wash 6 times with 700 µL wash buffer 1X solution per well using an automatic plate washer with overflow-wash function. Do not include soak step in washing procedure.

Or manually.

Discard the reaction volume by inverting the microplate over a sink. Add 350  $\mu$ L wash buffer 1X solution to each well. Discard the wash solution, tap firmly several times against absorbent paper to remove excess liquid. Repeat 5 times. Avoid prolonged soaking during washing procedure.

- 7. Add 200 µL Substrate TMB into each well.
- 8. Incubate 15 minutes on the bench at room temperature (18-25°C).
- 9. Add 50  $\mu$ L Stop Solution to each well. Place the plate on the shaker for approximately 5 seconds to ensure mixing.
- Read optical density at 450 nm and calculate results. Read within 30 minutes.

Note! Be extra careful not to contaminate the Substrate TMB with enzyme conjugate solution.

## Internal quality control

Commercial controls and/or internal serum pools with low, intermediate and high insulin concentrations should routinely be assayed as samples, and results charted from day to day. It is good laboratory practice to record the following data for each assay: kit lot number, dilution and/or reconstitution dates of kit components, OD values for the blank, Calibrators and controls.

#### Calculation of results

The concentration of insulin is obtained by computerized data reduction of the absorbance for the Calibrators, except for Calibrator O, versus the concentration using cubic spline regression.

## Example of results

Wells	Identity	A <sub>450 nm</sub>	Mean conc. µg/L
1A-B	Calibrator 0	0.070/0.072	
1C-D	Calibrator 1*	0.115/0.116	
1E-F	Calibrator 2*	0.194/0.200	
1G-H	Calibrator 3*	0.567/0.580	
2A-B	Calibrator 4*	1.310/1.321	
2C-D	Calibrator 5*	2.480/2.452	
2E-F	Sample 1	0.601/0.618	32.2
2G-H	Sample 2	0.330/0.331	15.3

<sup>\*</sup>Concentration stated on vial label.

## Example of calibrator curve

A typical calibrator curve is shown here. Do not use this curve to determine actual assay results.



#### Conversion factor

1 µg corresponds to 0.174 nmol:

μg/L	3	7.5	30	75	150
nmol/L	0.52	1.30	5.22	13.05	26.1

## Limitations of the procedure

#### Performance limitations

Grossly lipemic, icteric or hemolyzed samples do not interfere in the assay. However, hemolysis in serum and plasma samples may result in a degradation of insulin which could give falsely low values and contributes to higher inter assay variation. The degradation is dependent on time, temperature and the hemoglobin concentration. Keep hemolyzed samples cold or on ice to prevent the insulin degradation.

## **Expected values**

Good practice dictates that each laboratory establishes its own expected range of values.

#### Performance characteristics

#### **Detection limit**

Detection limit is defined as the Capability of Detection according to ISO11843-Part 1. Capability of Detection should be seen as part of a method validation, rather than the lowest concentration that can be measured. The detection limit is  $\leq$ 1.5 µg/L as determined by the methodology described in ISO11843- Part 4. Concentration of samples with absorbance below Calibrator 1 should not be calculated, instead expressed as less or equal to ( $\leq$ ) the concentration indicated on the vial for Calibrator 1.

## Recovery

Recovery upon addition is 106%. Recovery upon dilution is 100%.

#### Hook effect

Samples with a concentration of up to at least 576  $\mu$ g/L can be measured without giving falsely low results.

#### Precision

Each sample was analyzed in 4 replicates on 5 different occasions. The analysis were done in one laboratory with 1 kit lot and by 1 technician.

		Coefficient of variation		
Sample	Mean value μg/L	Repeatability %*	Within laboratory %**	
1	16.2	4.5	6.1	
2	32.7	4.8	5.4	

<sup>\*</sup>Within assay variation

## Reproducibility

Each sample was analyzed in 4 replicates on 107 different occasions. The analysis were done in two laboratories with 18 kit lots and by 6 technicians.

		Co	pefficient of variat	ion
Sample	Mean value µg/L	within assay %	between assay %	total assay %
1	14.3	4.6	3.4	3.9
2	30.8	3.9	2.9	3.7

<sup>\*\*</sup>Total assay variation

## Specificity

The following cross reactions have been found:

	Crossreaction
IGF-I	n.d.
IGF-II	n.d.
Mouse C-peptide I	n.d.
Mouse C-peptide II	n.d.
Rat C-peptide I	n.d.
Rat C-peptide II	n.d.
Mouse insulin	75%
Rat insulin	100%
Mouse proinsulin I	33%
Mouse proinsulin II	51%
Rat proinsulin I	8%
Rat proinsulin II	51%
Bovine insulin	78%
Porcine insulin	476%
Ovine insulin	179%
Human C-peptide	n.d.
Human insulin	167%
Human proinsulin	75%
Insulin lispro (Humalog® Eli Lilly)	167%

n.d. = not detected

#### Calibration

Mercodia High Range Rat Insulin ELISA is calibrated against an in-house reference preparation of rat insulin.

## Warranty

The performance data presented here was obtained using the procedure indicated. Any change or modification in the procedure not recommended by Mercodia AB may affect the results, in which event Mercodia AB disclaims all warranties expressed, implied or statutory, including the implied warranty of merchantability and fitness for use.

Mercodia AB and its authorized distributors, in such event, shall not be liable for damages indirect of consequential.

## References

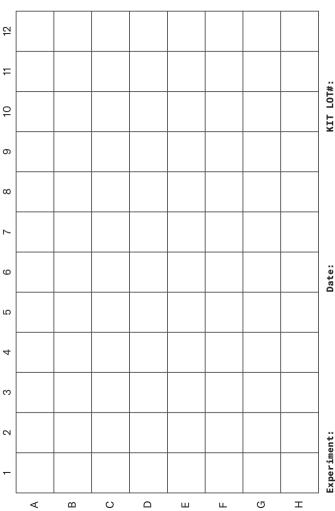
Kullin M *et al.* (2003) Protection of rat pancreatic islets by potassium channel openers against alloxan, sodium nitroprusside and interleukin-1beta mediated suppression-possible involvement of the mitochondrial membrane potential. *Diabetologia* 46:80-88.

von Mach MA et al. (2003) Cryopreservation of islets of Langerhans: Optimization of protocols using rat pancreatic tissue. EXCLI Journal 2:6-21.

Further references can be found on our website: www.mercodia.com

Notes		

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Date: Experiment:

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## Summary of protocol sheet Mercodia High Range Rat Insulin ELISA

Add Calibrators, controls* and samples	10 μL
Add enzyme conjugate 1X solution	50 μL
Incubate	2 hours at 18–25°C on a plate shaker, 700–900 rpm
Wash	700 μL, 6 times
Add Substrate TMB	200 μL
Incubate	15 minutes at 18–25°C
Add Stop Solution	50 μL Shake for 5 seconds to ensure mixing
Measure A <sub>450 nm</sub>	Evaluate results

<sup>\*</sup>not provided

For full details see page 7

For technical support please contact: support@mercodia.com