

ElisaRSR™ 2 Screen ICA™

2 Screen Islet Cell Autoantibody ELISA Kit - Instructions for use



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INTENDED USE

The RSR 2 Screen Islet Cell autoantibody (2 Screen) ELISA kit is intended for use by professional persons only, for quantitative determination of both GAD and IA-2 autoantibodies in human serum.

Autoantibodies to pancreatic beta cell antigens are important serological markers of type 1 diabetes mellitus. The antigens recognised by these antibodies include insulin, glutamic acid decarboxylase (GAD₆₅ kDa isoform), the islet cell antigen named IA-2 or ICA-512 and zinc transporter 8 (ZnT8). RSR's 2 Screen ELISA allows simultaneous measurement of GAD and IA-2 autoantibodies in the same sample.

REFERENCES

S. Chen et al

Sensitive non-isotopic assays for autoantibodies to IA2 and to a combination of both IA2 and GAD $_{65}$. Clinica Chimica Acta 2005 357: 74-83

C. Törn et al

Diabetes Antibody Standardization Program: evaluation of assays for autoantibodies to glutamic acid decarboxylase and islet antigen-2. Diabetologia 2008 51:846-852.

PATENTS

The following patents apply:

European patent EP 1 448 993 B1, Chinese patent ZL 02822274.1, Indian patent 226484, Japanese patent 5711449 and US patents US 8,129,132 B2, US 9,435,797 B2 and US 10,481,156 B2.

ASSAY PRINCIPLE

In RSR's 2 Screen ELISA, GAD and IA-2 autoantibodies (Ab) in patient sera, calibrators and controls are allowed to interact with GAD65 and coated onto ELISA plate wells (1st incubation). The samples are then discarded, leaving any GAD or IA-2 autoantibodies in the patient sera, calibrators or controls bound to the GAD₆₅ and IA-2 coated wells. A mixture of GAD₆₅-Biotin and IA-2-Biotin is then added and during a second incubation step (through the ability of GAD and IA-2 autoantibodies to act divalently), a bridge is formed between the GAD65 or IA-2 bound to the wells and GAD65-Biotin or IA-

2-Biotin respectively. The amount of $GAD_{65}/IA-2$ -Biotin bound is determined in a third incubation step by the addition of Streptavidin Peroxidase (SA-POD), which binds specifically to Biotin.

Excess unbound SA-POD is then washed away and addition of 3,3',5,5' tetramethylbenzidine (TMB) results in formation of a blue colour. This reaction is stopped by addition of stop solution causing the well contents to turn from blue to yellow. The absorbance of the yellow reaction mixture at 450nm is then read using an ELISA plate reader. A higher absorbance indicates the presence of GAD or IA-2 Ab in the test sample. Reading at 405nm allows quantitation of high absorbances.

STORAGE AND PREPARATION OF TEST SERUM SAMPLES

Sera to be analysed should be assayed soon after separation or stored, preferably in aliquots, at or below -20°C. 100μL is sufficient for one assay (duplicate 50µL determinations). Repeated freeze thawing or increases in storage temperature must be avoided. Do not use lipaemic or haemolysed serum samples. Do not use plasma in the assay. When required, thaw test sera at temperature and mix gently to ensure homogeneity. Centrifuge serum prior to assay (preferably for 5 min at 10-15,000 rpm in a microfuge) to remove particulate matter. Please do not omit this centrifugation step if sera are cloudy or contain particulates.

SYMBOLS

STIVIBULS	
Symbol	Meaning
C€	EC Declaration of Conformity
IVD	In Vitro Diagnostic Device
REF	Catalogue Number
LOT	Lot Number
[]i	Consult Instructions
***	Manufactured By
23	Expiry Date
2°C	Store
CONTROL +	Positive Control
CONTROL .	Negative Control

MATERIALS REQUIRED AND NOT SUPPLIED

Pipettes capable of dispensing $25\mu L$, $50 \mu L$ and 100μL.

Means of measuring out various volumes to reconstitute or dilute reagents supplied.

Pure water

ELISA Plate reader suitable for 96 well formats and capable of measuring at 450nm and 405nm ELISA Plate shaker, capable of 500 shakes/min (not an orbital shaker).

ELISA Plate cover

PREPARATION OF REAGENTS SUPPLIED

Store	unopened kit and components at 2 - 8°C
_	GADes and IA-2 Coated Wells 12 breakapart strips of 8 wells (96 in total) in a frame and sealed in foil bag. Allow to stand at room temperature (20 – 25°C) for at least 30 minutes before opening. Ensure stripwells are firmly fitted into
A	frame provided. After opening return any unused wells to the original foil packet with desiccant provided and seal with adhesive tape. Place foil bag in the self-seal plastic bag and store at 2-8°C for up to 8 months.
В	Reaction Enhancer 4 mL coloured red Ready for use
C1- 6	Calibrators 4, 10, 20, 70, 145 and 450 u/mL (units are NIBSC 97/550) 6 x 0.7 mL Ready for use
D1	GAD Ab Positive Control 0.7 mL Ready for use
D2	IA-2 Ab Positive Control 0.7 mL Ready for use
D3	Negative Control 0.7 mL Ready for use
E	GAD ₆₅ /IA-2-Biotin (GAD ₆₅ Biotin plus IA-2 Biotin) 3 vials lyophilised Reconstitute each vial with the amount of reconstitution buffer for GAD ₆₅ /IA-2-Biotin (F) shown on the vial label. When more than one vial is used, pool the reconstituted vials and mix gently before use. Use on day of reconstitution.
F	Reconstitution Buffer for GADes/IA-2-Biotin 2 x 15 mL coloured blue Ready for use
G	Streptavidin Peroxidase (SA-POD) 1 x 0.7 mL Concentrated Dilute 1 in 20 with diluent for SA-POD (H). For example, 0.5mL (G) + 9.5mL (H). Store at 2 - 8°C for up to 18 weeks after dilution.
н	Diluent for SA-POD 15 mL Ready for use

	Peroxidase Substrate (TMB)
'	15 mL Ready for use
	Concentrated Wash Solution
	125 mL
J	Concentrated
	Dilute 10 X with pure water before use.
	Store at 2 – 8°C up to kit expiry.
	Stop Solution
K	12 mL
	Ready for use

ASSAY PROCEDURE

Allow all reagents to stand at room temperature (20 - 25°C) for at least 30 minutes before use. A repeating Eppendorf type pipette is recommended

for s	teps 2	, 6, 9, 11 and 12.		
	1.	Pipette 50 μL of patient sera,		
		calibrators (C1-6) and controls (D1,		
		D2 and D3) into respective wells in		
		duplicate, leaving one well empty for		
		blank (see step 13).		
_	2.	Pipette 25 μL of reaction enhancer (B)		
Day 1		into each well (except blank).		
	3.	Cover the frame and shake the wells		
		for 5 seconds on an ELISA plate		
		shaker (500 shakes per min).		
	4.	Incubate the plate at 2 – 8°C (without		
		shaking) overnight (16-20 hours)		
	5.	After this overnight incubation,		
		aspirate the samples and wash the		
		plate 3 times with wash solution (J)		
		using a plate washer. (If a plate		
		washer is not available, discard the		
		samples by briskly inverting the frame		
		of stripwells over a suitable		
		receptacle, wash the wells 3 times		
		manually and after the final wash		
		invert the frame of wells and tap		
		gently on a clean dry absorbent		
		surface to remove excess wash		
		solution).		
Day 2	6.	Pipette 100μL of reconstituted		
۵		GAD ₆₅ /IA-2-Biotin (E) into each well		
		(except blank). Avoid splashing the		
		material out of the wells during		
		addition.		
	7.	Cover the plate, and incubate at 18 -		
		22 °C for 1 hour on an ELISA plate		
		shaker (500 shakes per min).		
	8.	Repeat wash step 5.		
	9.	Pipette 100μL of diluted SA-POD (G)		
		into each well (except blank) and		
		incubate at room temperature for 20		
		minutes, on an ELISA plate shaker		
		(500 shakes per min).		

	10.	After the incubation, wash the wells three times with diluted wash solution (J) as in step 5 (in the case of washing manually, use an additional final wash step with pure water to remove any foam).		
well (including blank) and incuba		Pipette 100µL of TMB (I) into each well (including blank) and incubate in the dark at room temperature for 20 minutes without shaking.		
Day 2 continued	12.	Pipette 100µL stop solution (K) into each well (including blank) and shake the plate for approximately 5 seconds on a plate shaker (500 shakes per min). Ensure substrate incubations are the same for each well.		
	13.	Within 10 minutes read the absorbance of each well at 405nm and then 450nm using an ELISA plate reader, blanked against a well containing 100µL of TMB substrate (I) and 100µL Stop solution (K) only.		

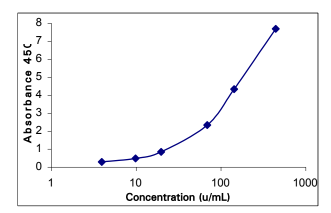
RESULT ANALYSIS

A calibration curve can be established by plotting calibrator concentration on the x-axis (log scale) against the absorbance of the calibrators on the yaxis (linear scale). The GAD and/or IA-2 Ab concentrations in patient sera can then be read off the calibration curve [Plotted at RSR as a spline log/lin curve (smoothing factor = 0)]. Other data reduction methods can be used. The negative control (D3) has a concentration of 0 u/mL, but can be assigned a value of 0.4 u/mL to facilitate computer processing of data. Absorbance readings at 405nm can be converted to 450nm absorbance values by multiplying by the appropriate factor (approximately 3.5, dependant on equipment being used). Values less than 25 u/mL should be read off a 450 nm curve.

Samples with high GADAb and IA-2Ab concentrations can be diluted in kit negative control (D3). For example, 15 μ L of sample plus 135 μ L of negative control to give a 10x dilution. Other dilutions (e.g. 100x) can be prepared from a 10x dilution or otherwise as appropriate. Some sera will not dilute in a linear way.

TYPICAL RESULTS (Example only; not to be used for calculation of actual results)

Calibrator	Absorbance	
u/mL	450nm	405nm
Negative Control	0.120	0.039
4	0.261	0.083
10	0.453	0.133
20	0.818	0.228
70	2.307	0.659
145	4.305	1.230
450	7.662	2.189



Index Calculation

If results are to be expressed as an index, only the $4\ \text{u/mL}$ calibrator need be included in the assay (all controls should still be included). The index values are calculated as follows:

$$Index = \frac{test \ sample \ absorbance \ at \ 450nm}{4 \ u/mL \ calibrator \ absorbance \ at \ 450nm}$$

Healthy blood donor sera give index values of less than 1 suggesting that index values of 1 or more can be considered positive for GADAb and/or IA-2 Ab.

ASSAY CUT OFF

	u/mL
Negative	< 4 u/mL
Positive	≥ 4.0 u/mL

This cut off has been validated at RSR. However each laboratory should establish its own normal and pathological reference ranges for GAD and/or IA-2 Ab levels. Also it is recommended that each laboratory include its own panel of control samples in the assay.

CLINICAL EVALUATION

Clinical Specificity and sensitivity

Sera from 70 healthy blood donors were all negative in the 2 Screen ELISA, although occasional healthy blood donors may have detectable GAD autoantibodies. Autoantibodies to GAD and/or IA2 were detected in 84% (n=216) of samples from patients with type 1 diabetes of various disease durations. In the DASP 2005 study, the RSR 2 Screen ELISA showed 98% (n=100) specificity and 96% (n=50) sensitivity.

Lower Detection Limit

The kit negative control was assayed 20 times and the mean and standard deviation calculated. The lower detection limit at +2 standard deviations was 0.43 u/mL.

Intra Assay Precision

Sample	u/mL (n = 25)	CV (%)
1	6.6	6.3
2	25.7	4.7

Inter Assay Precision

Sample	u/mL (n = 28)	CV (%)
3	115.2	3.4
4	21.2	4.4

Clinical Accuracy

Analysis of sera from patients with autoimmune diseases other than type 1 DM indicated no interference from autoantibodies to the TSH receptor, thyroglobulin, thyroid peroxidase, ds-DNA the acetylcholine receptor or from rheumatoid factor.

Interference

No interference was observed when samples were spiked with the following materials; haemoglobin up to 5mg/mL, bilirubin up to 20 mg/dL or intralipid up to 3000 mg/dL.

SAFETY CONSIDERATIONS

Streptavidin Peroxidase (SA-POD)

Signal word: Warning Hazard statement(s)

H317: May cause an allergic skin reaction

Precautionary statement(s)

P280: Wear protective gloves/protective clothing/

eye protection/face protection

P302 + P352: IF ON SKIN: Wash with plenty of

soap and water

P333 + P313: If skin irritation or rash occurs: Get

medical advice/attention

P362 + P364: Take off contaminated clothing

and wash it before reuse

Peroxidase Substrate (TMB)

Signal word: Danger Hazard statement(s)



H360: May damage fertility or the unborn child

Precautionary statement(s)

P280: Wear protective gloves/protective clothing/ eve protection/face protection

P308 + P313: IF exposed or concerned: Get

medical advice/attention

This kit is intended for *in vitro* use by professional persons only. Follow the instructions carefully. Observe expiry dates stated on the labels and the specified shelf life for coated wells, reconstituted reagents and diluted reagents. Refer to Safety Data Sheet for more detailed safety information. Material of human origin used in the preparation of the kit has been tested and found non reactive for HIV1 and 2 and HCV antibodies and HBsAg but should, none the less, be handled as potentially infectious. Wash hands thoroughly contamination has occurred and before leaving the laboratory. Sterilise all potentially contaminated waste, including test specimens before disposal. Material of animal origin used in the preparation of the kit has been obtained from animals certified as healthy but these materials should be handled as potentially infectious. Some components contain small quantities of sodium azide as preservative. With all kit components, avoid ingestion, inhalation, injection and contact with skin, eyes and clothing. Avoid formation of heavy metal azides in the drainage system by flushing any kit component away with copious amounts of water.

ASSAY PLAN

ASSAY PLAN		
Allow all reagents and samples to reach room temperature (20 – 25°C) before use		
Pipette:	50μL Calibrators, Controls, Patient Sera (except blanks)	
Pipette:	25μL Reaction Enhancer (except blanks)	
Mix:	Shake for 5 seconds at 500 shakes/min	
Incubate	Overnight (16-20) hours at 2 – 8°C (without shaking)	
Aspirate/Decant:	Plate	
Wash:	Plate three times (dry on absorbent material for manual wash)	
Pipette:	100μL GAD/IA-2 Biotin (reconstituted) into each well (except blanks)	
Incubate:	1 hour at 18 - 22 °C with shaking at 500 shakes/min	
Aspirate/Decant:	Plate	
Wash:	Plate three times (dry on absorbent material for manual wash)	
Pipette:	100μL SAPOD (diluted 1:20) into each well (except blanks)	
Incubate:	20 minutes at room temperature with shaking at 500 shakes/min	
Aspirate/Decant:	Plate	
Wash:	Plate three times, (additional rinse with pure water and dry on absorbent material for	
	manual wash)	
Pipette:	100μL TMB into each well (including blanks)	
Incubate:	20 minutes at room temperature in the dark (without shaking)	
Pipette:	100μL stop solution into each well (including blanks) and shake for 5 seconds	
Read absorbance at 405nm and 450nm within 10 minutes of stop solution addition.		