

Rat IgG ELISA Kit

Catalog NO.: RK00172

version: 2.0

This package insert must be read in its entirety before using this product

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Introduction

The kit is a sandwich enzyme immunoassay for in vitro quantitative measurement of lgG in rat serum, plasma, cell culture supernatants and other biological fluids.

Principle of the Assay

This assay employs the quantitative sandwich enzyme immunoassay technique. An antibody specific for rat lgG has been pre-coated onto a microplate.Standards and samples are pipetted into the wells and any lgG present is bound by the immobilized antibody. After washing away any unbound substances, and then a detection antibody specific for lgG is added to the wells and binds to the combination of capture antibody lgG in sample. Following a wash to remove any unbound combination, and enzyme conjugate is added to the wells. Following incubation and wash steps, a substrate solution is added to the wells and color develops in proportion to the amount of lgG bound in the initial step. The color development is stopped and the absorbance is measured.



Material Provided & Storage Conditions

Unopened kits can be stored at 2-8°C for 1 year, and opened products must be used within 1 month.

Part	Size	Cat. No.	Storage of opened/reconstituted material
Rat lgG Antibody Coated Plate	8×12	RM007 16	Put the unused slats back in the aluminum foil bag with the desiccant and reseal them. They can be stored at 2-8°C for 1 month.
Rat lgG Standard Lyophilized	2 vials	RM007 13	lt is not recommended to use again after redissolving.
Rat lgG Concentrated Biotin Conjugate Antibody (100×)	1 ×120ul	RM007 14	Store at 2-8°C for 1 month *

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Streptavidin-HRP Concentrated (100×)	1 ×120ul	RM007 15	Store at 2-8°C for 1 month *
Standard/Sampl e Diluent (R1)(4×)	1 ×20mL	RM180 00	
Biotin-Conjugate Antibody Diluent (R2)	1 ×12mL	RM000 24	
Streptavidin-HRP Diluent(R3)	1 ×12mL	RM000 25	Store at 2-8°C for 1 month
Wash Buffer(20x)	1 ×30mL	RM000 26	*
TMB Substrate	1 ×12mL	RM000 27	
Stop Solution	1 ×6mL	RM000 28	
Plate Sealers	4 Strips		
Specification	1		



Other Supplies Required

- Microplate reader capable of measuring absorbance at 450 nm, with the correction wavelength set at 630 nm or 570 nm.
- 2. Pipettes and pipette tips.
- 3. Deionized or distilled water.
- Squirt bottle, manifold dispenser, or automated microplate washer.
- 5. Incubator.
- 6. Test tubes for dilution of standards and samples.



Precautions

- Any variation in diluent, operator, pipetting technique, washing technique, incubation time or temperature, and kit age can cause variation in binding.
- Variations in sample collection, processing, and storage may cause sample value differences.
- Reagents may be harmful, if ingested, rinse it with an excess amount of tap water.
- 4. Stop Solution contains strong acid. Wear eye, hand, and face protection.
- Please perform simple centrifugation to collect the liquid before use.
- 6. Do not mix or substitute reagents with those from other lots or other sources.
- Adequate mixing is particularly important for good result.
 Use a mini-vortexer at the lowest frequency.
- Mix the sample and all components in the kits adequately, and use clean plastic container to prepare all diluents.



- Both the sample and standard should be assayed in duplicate, and reagents should be added in sequence in accordance with the requirement of the specification.
- 10. Reuse of dissolved standard is not recommended.
- 11. The kit should not be used beyond the expiration date on the kit label.
- 12. The kit should be away from light when it is stored or incubated.
- To reduce the likelihood of blood-borne transmission of infectious agents, handle all serum, plasma, and other biological fluids in accordance with NCCLS regulations.
- 14. To avoid cross contamination, please use disposable pipette tips.
- 15. Please prepare all the kit components according to the Specification. If the kits will be used several times, please seal the rest strips and preserve with desiccants. Do use up within 2 months.
- This assay is designed to eliminate interference by other factors present in biological samples.
- 17. Until all factors have been tested in this assay, the possibility of interference cannot be excluded.
- 18. The 48T kit is also suitable for the specification.



Sample Collection & Storage

The sample collection and storage conditions listed below are intended as general guidelines. Sample stability has not been evaluated.

Cell Culture Supernatant: Remove particulates by centrifugation. Assay immediately or aliquot and store samples at \leq -20 °C. Avoid repeated freeze-thaw cycles.

Serum: Use a serum separator tube (SST) and allow samples to clot for 30 minutes at room temperature before centrifugation for 15 minutes at 1000 x g. Remove serum and assay immediately or aliquot and store samples at \leq -20 °C. Avoid repeated freeze-thaw cycles.

Plasma: Collect plasma using EDTA or Heparin as an anticoagulant. Centrifuge for 15 minutes at $1000 \times g$ within 30 minutes of collection. Assay immediately or aliquot and store samples at \leq -20°C. Avoid repeated freeze-thaw cycles. (Note:



Citrate plasma has not been validated for use in this assay.)

Other biological fluids: Centrifuge samples for 20 minutes at $1,000 \times g$. Collect the supernatants and assay immediately or store samples in aliquot at -20°C or -80°C for later use. Avoid repeated freeze-thaw cycles.

Note: It is suggested that all samples in one experiment be collected at the same time of the day. Avoid hemolytic and hyperlipidemia sample for serum and plasma.

Reagent Preparation

Bring all reagents to room temperature before use. If crystals have formed in the concentrate, Bring the reagent to room temperature and mix gently until the crystals have completely dissolved.



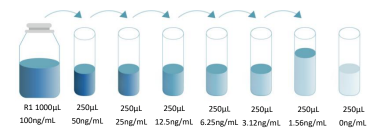
Standard/Sample Diluent (R1)(4x) -Dilute 1:4 with double distilled or deionized water before use, for example : Add 5 mL of Standard/Sample Diluent Concentrate to 15mL of deionized or distilled water to prepare 20 mL of Standard/Sample Diluent .

Standard - Reconstitute the Standard Lyophilized with 1.0 mL Standard/Sample Diluent(R1)(1x). This reconstitution produces a stock solution of 100 ng/mL. Mix the standard to ensure complete reconstitution and allow the standard to sit for a minimum of 15 minutes with gentle agitation prior to making dilutions.

Use the 100 ng/mL standard stock to produce a dilution series (below) with Standard/Sample Diluent(R1)(1x). Mix each tube thoroughly and change pipette tips between each transfer (recommended concentration for standard curve: 100, 50, 25, 12.5, 6.25, 3.12,1.56, 0 ng/mL). Use diluted standards within 60 minutes of preparation.

Std 250 μL 250 μL 250 μL 250 μL 250 μL 250 μL





Working Biotin Conjugate Antibody - Dilute 1:100 of Concentrated Biotin Conjugate Antibody (100x) with Biotin-Conjugate Antibody Diluent (R2) before use, for example: Add 20 μ L of Concentrated Biotin Conjugate Antibody (100x) to 1980 μ L Biotin-Conjugate Antibody Diluent (R2) to prepare 2000 μ L Working Biotin Conjugate Antibody Buffer.

Working Streptavidin-HRP - Dilute 1:100 of Concentrated Streptavidin-HRP (100x) with Streptavidin-HRP Diluent (R3) before use, for example: Add 20 μ L of Concentrated Streptavidin-HRP (100x) to 1980 μ L Streptavidin-HRP Diluent (R3) to prepare 2000 μ L Working Streptavidin-HRP Buffer.

Wash Buffer - If crystals have formed in the concentrate, warm

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to room temperature and mix gently until the crystals have completely dissolved. Dilute 1:20 with double distilled or deionized water before use, for example : Add 20 mL of Wash Buffer Concentrate to 380 mL of deionized or distilled water to prepare 400 mL of Wash Buffer.



Assay Procedure

Bring all reagents and samples to room temperature before use. It is recommended that all standards, controls, and samples be assayed in duplicate.

- Prepare all reagents, working standards, and samples as directed in the previous sections.
- Remove excess microplate strips from the plate frame, return them to the foil pouch containing the desiccant pack, and reseal properly.
- Add wash buffer 350 µL/well, aspirate each well after holding 40 seconds, repeating the process two times for a total of three washes.
- Add 100 μL Standard/sample Diluent (R1)(1x) in a blank well.
- Add 100 µL different concentration of standard or sample in other wells, Cover with the adhesive strip provided. Incubate for 2 hours at 37°C. record the plate layout of standards and sample assay.
- Prepare the Concentrated Biotin Conjugate Antibody (100x) Working Solution 15 minutes early before use.
- 7. Repeat the aspiration/wash as in step 3.
- 8. Add 100 μ L Working Biotin Conjugate Antibody in each

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well, cover with new adhesive Sealer provided.Incubate for 1 hour at 37°C.

- Prepare the Streptavidin-HRP Concentrated (100x)
 Working Solution 15minutes early before use.
- 10. Repeat the aspiration/wash as in step 3.
- Add 100 µL Working Streptavidin-HRP in each well, cover with new adhesive Sealer provided.Incubate for 0.5 hour at 37℃.
- 12. Repeat the aspiration/wash as in step 3.
- During the incubation, turn on the microplate reader to warm up for 30 minutes before measuring.
- Add 100 μ L TMB Substrate to each well. Incubate for 15-20 minutes at 37°C .Protect from light.
- 15. Add 50 µL Stop Solution, determine the optical density of each well within 5 minutes, using a Microplate reader set to 450 nm. If wavelength correction is available, set to 570 nm or 630 nm. If wavelength correction is not available, subtract readings at 570 nm or 630 nm from the readings at 450 nm. This subtraction will correct for optical imperfections in the plate. Readings made directly at 450 nm without correction may cause higher value and less accurate result.



Assay Procedure Summary

Prepare the standard and reagents

Wash 3 times

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Add 100ul of standards or test samples to each well

Incubate for 2 hours at 37°C, then wash 3 times

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Add 100ul Working Biotin Conjugate Antibody

Incubate for 1 hour at 37°C, then wash 3 times

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Add 100ul Working Streptavidin-HRP Incubate for 0.5 hour at 37°C, then wash 3 times

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Add 100ul Substrate Solution

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Add 50ul Stop Solution

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Detect the optical density within 5 minutes under 450nm. Correction Wavelength set at 570nm or 630nm

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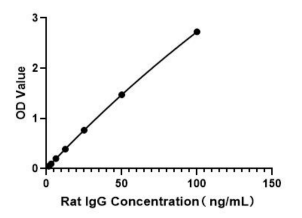


Calculation of Results

- Average the duplicate readings for each standard, control and sample, and subtract the average zero standard optical density (O.D.).
- 2. Create a standard curve by reducing the data using computer software capable of generating a four-parameter logistic (4-PL) curve-fit. As an alternative, construct a standard curve by plotting the mean absorbance for each standard on the Y-axis against the concentration on the X-axis and draw a best fit curve through the points on a log/log graph. The data may be linearized by plotting the log of the lgG concentrations versus the log of the O.D. on a linear scale, and the best fit line can be determined by regression analysis.
- If samples have been diluted, the concentration read from the standard curve must be multiplied by the dilution factor.



Typical Data



The standard curves are provided for demonstration only. A standard curve should be generated for each set of lgG assayed.

Detection Range

1.56-100 ng/mL



Sensitivity

The minimum detectable dose (MDD) of lgG typically less than 0.78 ng/mL. The MDD was determined by adding two standard deviations to the mean optical density value of twenty zero standard replicates and calculating the corresponding concentration.

Specificity

This method has high sensitivity and specificity for IgG detection, and there is no obvious cross-reaction or interference between IgG and analogue.

Note:

Due to the limitations of existing technology and knowledge, it is impossible to complete the detection of cross-reactions between IgG and all analogues, so cross-reactions may still exist.



Precision

Intra-plate Precision

3 samples with low, middle and high level lgG were tested 20 times on one plate, respectively.

Intra-Assay: CV<10%

Inter-plate Precision

3 samples with low, middle and high level lgG were tested on 3 different plates, 20 replicates in each plate.

Inter-Assay: CV<15%

	Intra-Assay Precision			Inter-	Assay Pre	cision
Sample	1	2	3	1	2	3
n	20	20	20	20	20	20
Mean(ng/mL)	26.5	52.1	55	6.5	18.9	49.3
Standard deviation	1.04	2.1	1.5	0.51	1.8	3.02
CV(%)	3.9	4.0	2.7	7.8	9.5	6.1



Recovery

Matrices listed below were spiked with certain level of lgG and the recovery rates were calculated by comparing the measured value to the expected amount of lgG in samples.

Sample	Average Recovery (%)	Range (%)
Cell Culture Media(n=5)	100	87-113
Serum(n=5)	94	82-106



Linearity

The linearity of the kit was assayed by testing samples spiked with appropriate concentration of lgG and their serial dilutions. The results were demonstrated by the percentage of calculated concentration to the expected.

/	/	Cell Culture Media(n=5)	Serum(n=5)
1:2	Average of Expected (%)	91	86
	Range (%)	80-102	82-90
1:4	Average of Expected(%)	94	90
	Range (%)	90-98	82-98
1:8	Average of Expected(%)	93	88
	Range (%)	85-100	81-95
1:16	Average of Expected(%)	98	91
	Range (%)	90-106	83-99



Trouble Shooting

Problem	Possible Cause	Solution
		Sufficiently wash plates as
		required. Ensure appropriate
	Insufficient washing	duration and number of washes.
		Ensure appropriate volume of
		wash buffer in each well.
High	Incorrect incubation	Check whether the duration and
Background	procedure	temperature of incubation are set
	procedure	up as required.
	Cross-contamination	Be careful of the operations that
	of samples and	could cause cross-contamination.
		Use fresh reagents and repeat the
	reagents	tests.
		Check the concentration and
	Incorrect use of	dilution ratio of reagents. Make
	reagents	sure to use reagents in proper
		order.
No singular		Warm the reader up before use.
No signal or weak signal	Incorrect use of	Make sure to set up appropriate
	microplate reader	main wavelength and correction
		wavelength.
	Insufficient colour	Optimum duration of colour
	reaction time	reaction should be limited to 15-25
	reaction time	minutes.



	Read too late after stopping the colour reaction Matrix effect of	Read the plate in 5 minutes after stopping the reaction. Use positive control.
Too much signal	samples Contamination of TMB substrate	Check if TMB substrate solution turns blue. Use new TMB substrate solution.
	Plate sealers reused	Use a fresh new sealer in each step of experiments.
	Protein concentration in sample is too high	Do pre-test and dilute samples in optimum dilution ratio.
Poor Duplicates	Uneven addition of samples Impurities and precipitates in samples	Check the pipette. Periodically calibrate the pipette. Centrifuge samples before use.
	Inadequate mixing of reagents	Mix all samples and reagents well before loading.

*For research purposes only. Not for therapeutic or diagnostic purposes.