Check your package **insert & kit box for lot** specific instructions

KOMA ELISA Kit User Manual





Mouse MIP-2 ELISA kit

Catalog No.	K0331217
Lot No.	34025
Quantity	96 tests
Storage	4°C
Standard Range	15.625-1000 pg/ml

[Important Notice]

- Please read this User Manual carefully prior to performing the assay.
- The reagent preparation method might be different from lot to lot, so please check the lot and follow the instructions given in this manual.
- Do not mix or interchange reagents between different lots.
- The kit is intended for Research Use Only.

Table of Contents

DESCRIPTION	3
KIT COMPONENTS	3
STORAGE AND STABILITY	3
STANDARD RANGE	3
SAMPLE PREPARATION	4
REAGENT PREPARATION	5
ELISA PROCEDURE	7
CALCULATION OF RESULTS	9
CROSS REACTIVITY	9
TROUBLESHOOTING	10
PLATE LAYOUT	11
CERTIFICATE OF ANALYSIS	12

DESCRIPTION

This kit contains all the necessary reagents required for performing quantitative measurement of the protein from samples including serum, plasma, culture medium or other biological fluids in a sandwich ELISA format.

KIT COMPONENTS

Component	Amount
Pre-Coated 96 well ELISA microplate	1 Plate
Biotinylated Affinity Purified Detection Antibody (Lyophilized)	2 EA
Recombinant Standard Protein (Lyophilized)	2 EA
Streptavidin-HRP Conjugate (0.6 ml)	1 EA
Assay Diluent (50 ml) : 1% BSA in PBS	1 EA
Assay Dileunt G (10 ml) : N/A	N/A
TMB or pink-ONE Solution (10 ml)	1 EA
Stop Solution (10 ml)	1 EA
Wash Buffer Concentrate (20X, 50 ml) to make 1 liter	1 EA
Plate Sealer	3 EA

STORAGE AND STABILITY

- Store kit at 4°C immediately upon receipt.
- The shelf life of the kit is one year from date of shipment.
- Expiry of the kit is stated on labels.

STANDARD RANGE

Standard Range 15.625-1000 pg/ml

SAMPLE PREPARATION

- Store all samples on ice after preparation and use immediately or aliquot and store at -80°C.
- Avoid repeated freeze-thaw cycles.

1) Cell culture supernatants

Centrifuge cell culture media at 1,500 rpm for 10 minutes at 4°C to remove particulates.

Immediately aliquot supernatants and store at -80°C.

2) Serum

Samples should be collected into a serum separator tube. After clot formation, centrifuge samples at 3,000 rpm for 10 minutes at 4°C to remove clots. Immediately aliquot supernatants and store at -80°C.

3) Plasma

Collect plasma using anti-coagulant (citrate, EDTA or heparin). Centrifuge samples at 3,000 rpm for 15 minutes at 4°C. Immediately aliquot supernatants and store at -80°C.

REAGENT PREPARATION

- Do not mix or substitute Assay Diluent from other kit lots.
- All reagents should be prepared right before use, and diluted solution should be used immediately.

1) Standard Protein

Reconstitute 1 vial of Standard protein in 0.1 ml sterile water to a concentration of 200,000 pg/ml. Then dilute in Assay Diluent at 1:2 serial dilutions as follows. The standard diluent buffer serves as zero standard.

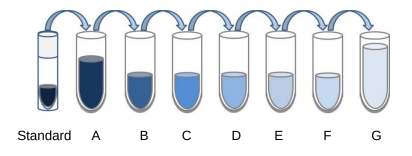


Table	1.
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Step	Dilution Method						tration
Step A	0.005	ml of Standard	+	0.995	ml of Assay Diluent	1000	pg/ml
Step B	0.5	ml of Step A	+	0.5	ml of Assay Diluent	500	pg/ml
Step C	0.5	ml of Step B	+	0.5	ml of Assay Diluent	250	pg/ml
Step D	0.5	ml of Step C	+	0.5	ml of Assay Diluent	125	pg/ml
Step E	0.5	ml of Step D	+	0.5	ml of Assay Diluent	62.5	pg/ml
Step F	0.5	ml of Step E	+	0.5	ml of Assay Diluent	31.25	pg/ml
Step G	0.5	ml of Step F	+	0.5	ml of Assay Diluent	15.625	pg/ml

2) Detection Antibody

Reconstitute 1 vial of Detection Antibody in 0.25 ml sterile water, and dilute 1:100 in Assay Diluent.

NOTE: Reconstituted solutions are stable at -20°C for up to 1 month. Do not repeat freezing and thawing.

3) Streptavidin-HRP

Dilute the Streptavidin-HRP conjugate 1:100 in Assay Diluent.

4) Wash Buffer

Dilute the 20X Wash Buffer Concentrate in 1 L distilled water.

ELISA PROCEDURE

 Washing: Add 200 ul of Washing Solution to each well. Aspirate the wells to remove liquid and wash the plate 3 times using 300 ul of Washing Solution per well. After the last wash, invert plate to remove residual solution and blot on paper towel.
 NOTE: Do not let the well dry completely and go immediately to the

NOTE: Do not let the well dry completely and go immediately to the next step.

- 2) **Reaction:** Add 100 ul of standard, blank and sample to each well in duplicate. Cover the plate with the Plate Sealer. Incubate at room temperature for at least 2 hours.
- Washing: Aspirate the wells to remove liquid and wash the plate 4 times as in step 1.
 NOTE: Vigorous washing of the plate after incubation steps is essential to obtaining low background values.
- 4) **Detection:** Add 100 ul of the diluted detection antibody per well. Then cover the plate with the Plate Sealer. Incubate at room temperature for 2 hours.
- 5) Washing: Aspirate and wash plate 4 times as in step 1.
- 6) **Conjugates:** Add 100 ul of the diluted Streptavidin-HRP per well. Cover the plate with the Plate Sealer. Incubate 30 minutes at room temperature (or at 37°C for 30 minutes).
- 7) Washing: Aspirate and wash plate 4 times as in step 1.
- 8) **Color Development:** Add 100 ul of TMB or pink-ONE TMB solution to each well. Incubate at room temperature for a proper color development. Add 100 ul of the stop solution to each well.

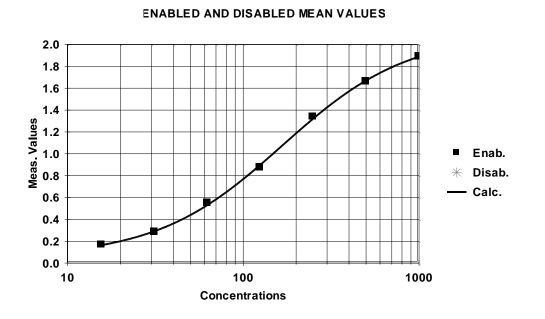
NOTE: Please be aware that the color may develop more quickly or more slowly than the recommended incubation time depending on the localized room temperature. Please monitor the color development to optimize the incubation time.

NOTE: Stop Solution (H_2SO_4) is a caustic material. Eye, hand, face and clothing protection should be worn when handling this reagent.

9) Reading: Using a microplate reader, measure observance at 450 nm.

CALCULATION OF RESULTS

Create a standard curve by reducing the data using ELISA reader's computer software capable of generating Standard curve-fit. A standard curve should be generated for each set of samples.



Mosue MIP-2 (pg/ml) (11 minutes color development)

CROSS REACTIVITY

To define the specificity of this ELISA, several proteins were tested for cross reactivity at 50 ng/ml.

Human :	GRO-α/MGSA, GRO-β, GRO-r, NAP-2
Mouse :	CXCL-16, IP-10, JE/MCP-1, KC, LIX, MIG, MIP-1α, MIP-1β, MIP-1r, MIP-3α, MIP-3β, SDF-1α, SDF-1β
Rat : Others :	GRO/KC, GRO-β N/A

TROUBLESHOOTING

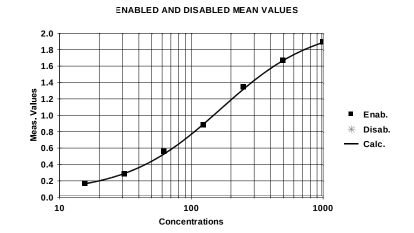
Problem	Probable Cause	Solution		
	Reagents not fresh or contaminated	Ensure proper preparation of		
Low OD No Signal	Incubation time not long enough	reagents. Ensure sufficient incubation times.		
		Reagent solutions should be at RT		
	Incubation temperature too low	before use.		
	Stop solution not added	Addition of stop solution		
	Inadequate standard dilution	Ensure proper dilution of Standard.		
High OD	Inadequate incubation time of detection antibody, Streptavidin- HRP or Substrate	Decrease incubation time.		
	Inadequate washing	Increase the stringency of washes.		
	Inaccurate pipetting	Ensure accurate pipetting of volume and avoid air bubbles.		
Poor consistency	Inadequate mixing of samples	Mix samples thoroughly before pipetting		
of replicates	High particulate matter of samples	Mix samples thoroughly and remove particulates by centrifugation.		
	Cross-well contamination	Use fresh plate sealers or pipette tips		
	Contamination of reagents or	Use a clean container before		
	samples	addition into wells.		
	Insufficient plates washing	Ensure proper washing of each well		
High	Too much concentrated detection antibody and Streptavidin-HRP	Ensure proper dilution of detection antibody or conjugate and incubation time.		
background	Substrate solution or stop solution is not fresh	Use fresh substrate and stop solution.		
	Plate left too long before reading	Read on the plate reader right after		
	on the plate reader	the experiment.		
	Incubation temperature is too high	Decrease incubation temperature of substrate.		
	Samples contain no or below	If samples are below detectable		
Poor	detectable levels of analyte or	levels, higher sample volume. If samples are higher than detectable		
standard	samples contain analyte			
curve	concentrations greater than the	levels, it may require dilution and		
	highest standard point.	reanalysis.		

PLATE LAYOUT

	1	2	3	4	5	6	7	8	9	10	11	12
Α												
В												
С												
D												
Е												
F												
G												
Н												

CERTIFICATE OF ANALYSIS

Product	Mouse MIP-2 ELISA kit
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Mosue MIP-2 (pg/ml) (11 minutes color development)

Layout map for calibrators sheet

		OD (450nm)		
А	Cal_1	1000	pg/ml	1.889
В	Cal_2	500	pg/ml	1.663
С	Cal_3	250	pg/ml	1.338
D	Cal_4	125	pg/ml	0.876
E	Cal_5	62.5	pg/ml	0.551
F	Cal_6	31.25	pg/ml	0.283
G	Cal_7	15.625	pg/ml	0.167
Н	Black	0	pg/ml	0.047