

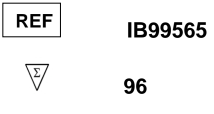


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Mouse TNF-a ELISA

Sandwich Enzyme-Linked Immunosorbent Assay for the detection of Mouse $TNF\mbox{-}\alpha$



Storage: 2-8°C



For research use only, not for use in diagnostic procedures.

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1. Introduction

Tumor necrosis factor (TNF)- α is an important cytokine in the innate immune response produced by macrophages, neutrophils, fibroblasts, keratinocytes, NK cells, T and B cells, and tumour cells. TNF- α mediates host responses in acute and chronic inflammatory conditions, and is a mediator of protection from infection and malignancy. TNF- α causes apoptotic cell death, cellular proliferation, differentiation, inflammation, tumorigenesis, and viral replication.

TNF- α is initially produced as a biologically active 26 kDa transmembrane protein, which is subsequently cleaved, principally by TNF- α -converting enzyme (TACE), to release the 17kDa free protein. These proteins are arranged in biologically active homotrimers that act on the ubiquitously expressed TNF- α receptors 1 and 2 (TNFR1 and TNFR2). These 17 kDa TNF protomers are composed of two antiparallel β -pleated sheets with antiparallel β -strands, forming a 'jelly roll' β -structure, typical for the TNF family.

TNF-R1 is constitutively expressed in most tissues, and can be fully activated by both the membrane-bound and soluble trimeric forms of TNF, while TNF-R2 is only found in cells of the immune system and respond to the membrane-bound form of the TNF homotrimer. Although their extracellular domains share structural and functional homology, their intracellular domains are distinct.

Tumor necrosis factor promotes the inflammatory response, which in turn causes many of the clinical problems associated with autoimmune disorders such as rheumatoid arthritis, ankylosing spondylitis, Crohn's disease, psoriasis and refractory asthma. These disorders are sometimes treated by using TNF inhibitors such as infliximab (Remicade) or adalimumab (Humira).

2. Principles of Method

The design of this assay is based on a sandwich Enzyme-Linked Immunosorbent Assay (ELISA). The microtiter plate provided in this kit has been pre-coated with a monoclonal antibody specific to mouse TNF- α . Samples are pipetted into these wells. Nonbound mouse TNF- α and other components of the sample should be removed by washing, then polyclonal antibody specific to mouse TNF- α added. In order to quantitatively determine the amount of mouse TNF- α present in the sample, streptavidin Horseradish Peroxidase (HRP) should be added to each microplate well. The final step, a TMB-substrate solution added to each well. Finally, a sulfuric acid solution is added and the resulting yellow colored product is measured at 450nm. Since the increases in absorbency is directly proportional to the amount of captured mouse TNF- α .

3. Intended Use

The IBL-AMERICA mouse TNF- α ELISA kit is to be used for the determination of mouse TNF- α in mouse serum, mouse plasma, cell lysate, culture supernatants and buffered solution. The assay will recognize native and recombinant mouse TNF- α .

This kit has been configured for research use only and is not to be used in diagnostic procedures.

4. Storage and Stability

All kit components of this kit are stable at 2 to 8°C. Any unused reconstituted standard should be discarded or frozen at -70°C. Standard can be frozen and thawed one time only without loss of immunoreactivity.

5. Chemical Hazard

- Stop solution: This reagent is an irritant to eyes, skin and mucous membranes. Avoid contact with eyes, skin and clothing. Wear suitable protective clothing, gloves and eye protection. In the event of contact with eyes or skin, wash immediately with plenty of water.
- Standard protein and 2nd antibody containing Sodium Azide as a preservative.

6. Kit Contents

Contents	Number	Volume
96 Well Plate	1 (in aluminum foil bag with desiccant)	
Washing Buffer	2	(20X) 25 ml
Standard Protein	1 Glass vial	(lyophilized)
Standard/Sample Dilution Buffer	1	25 ml
Secondary Antibody	1 Glass vial	(lyophilized)
Streptavidin HRP(X100)	1	150 μl
Secondary antibody/ streptavidin HRP Dilution Buffer	1	25 ml
Substrate (TMB)	1	15 ml
Stop Solution	1	15 ml
Protocol booklet		1
Plate sealers		2

- 1 96 Well Plate
 - : Mouse TNF- α microtiter plate, one plate of 96 wells (8 well strips x 12).

A plate using break-apart strips coated with a mouse monoclonal antibody specific to mouse TNF- α .

2 Standard Protein

: Recombinant mouse TNF- α .

- 3 Secondary Antibody
 - : Biotinylated anti mouse TNF- α antibody.
- (4) Streptavidin HRP(X100)
 - : Streptavidin Horseradish Peroxidase (HRP, enzyme)
- (5) Substrate (Stabilized chromogen)
 - : Tetramethylbenzidine (TMB) solution
- 6 Stop Solution
 - : 1N solution of sulfuric acid (H₂SO₄).
- ⑦ Plate sealer
 - : Adhesive sheet.
- Do not mix or interchange different reagents from various kit lots.

7. Materials Required But Not Provided

- ① Microtiter plate reader capable of measurement at or near 450nm.
- ② Calibrated, adjustable precision pipettes, preferably with disposable plastic tips (A manifold multi-channel pipette is desirable for large assays.)
- ③ Distilled or deionized water
- (4) Data analysis and graphing software
- (5) Vortex mixer
- (6) Polypropylene tubes for diluting and aliquoting standard
- Absorbent paper towels
- (8) Calibrated beakers and graduated cylinders of various sizes

8. Reagent Preparation

1) Mouse TNF-α standard

- 1. Reconstitute the lyophilized mouse TNF- α standard by adding 1 ml of *Standard/Sample Dilution Buffer* to make the 2 ng/ml standard stock solution. Allow solution to sit at RT for 5 minutes, then gently vortex to mix completely. Use within one hour of reconstituting.
- Prepare 1 ml of 500 pg/ml top standard by adding 250 μl of the above stock solution in 750 μl of *Standard/Sample Dilution Buffer*. Perform 2-fold serial dilutions of the top standards to make the standard curve within the range of this assay (7.81 pg/ml ~ 500 pg/ml) as below. *Standard/Sample Dilution Buffer* serves as the zero standard (0 pg/ml).

Standard	Add	Into	
500 pg/ml	250.0 µl of the std.(2 ng/ml)	750.0 µl of the Standard/Sample Dilution Buffer	
250 pg/ml	500 μl of the std.(500 pg/ml)	500.0 µl of the Standard/Sample Dilution Buffer	
125 pg/ml	500 μ l of the std.(250 pg/ml)	500.0 µl of the Standard/Sample Dilution Buffer	
62.5 pg/ml	500 μ l of the std.(125 pg/ml)	500.0 µl of the Standard/Sample Dilution Buffer	
31.25 pg/ml	500 μl of the std.(62.5 pg/ml)	500.0 µl of the Standard/Sample Dilution Buffer	
15.63 pg/ml	500 μl of the std.(31.25 pg/ml)	500.0 µl of the Standard/Sample Dilution Buffer	
7.81 pg/ml	500 μl of the std.(15.63 pg/ml)	500.0 µl of the Standard/Sample Dilution Buffer	
0 pg/ml	1.0 ml of the Standard/Sample Dilution Buffer		

2) Secondary Antibody

100X secondary antibody solution can be made by adding 150 µl Secondary antibody/ Streptavidin HRP dilution buffer in the vial.

1. Equilibrate to room temperature, mix gently.

2. Mix 20 μl Secondary Antibody concentrated solution (100X) + 1.98 ml Secondary antibody/ Streptavidin HRP dilution buffer. (Sufficient for one 16-well strip, prepare more if necessary)

Label as "Working Secondary antibody Solution".

3. Return the unused Secondary Antibody concentrated solution to the refrigerator.

3) Streptavidin HRP(X100)

- 1. Equilibrate to room temperature, mix gently.
- 2. Mix 20 μl *Streptavidin HRP concentrated solution* (100X) + 1.98 ml *Secondary antibody/ Streptavidin HRP dilution buffer*. (Sufficient for one 16-well strip, prepare more if needed)

Label as "Working Streptavidin HRP Solution".

3. Return the unused Streptavidin HRP concentrated solution to the refrigerator.

4) Washing buffer

- 1. Equilibrate to room temperature, mix to re-dissolve any precipitated salt.
- 2. Mix 0.5 volume *Wash buffer concentrate solution* (20X) + 9.5 volumes of deionized water. Label as "Working Washing Solution".
- 3. Store both the concentrated and the Working Washing Solution in the refrigerator.

* Directions for washing

1. Fill the wells with 300 µl of "Working Washing Buffer".

Let soak for 1 to 3 minutes and then all residual wash-liquid must be drained from the wells by aspiration (taking care not to scratch the inside of the well) or decantation, followed by forceful tapping of the plate on absorbent paper. Never insert absorbent paper directly into the wells.

If using an automated washer, the operating instructions for washing equipment should be carefully followed.

- 2. Incomplete washing will adversary affects the assay and renders false results.
- 3. It is recommended to use laboratory tape to hold plate strips to the plate frame while performing the plate washing to avoid strips coming free of the frame.

5) Sample preparation

Blood should be collected by venipuncture. For plasma samples, blood may be drawn into tubes containing sodium citrate or heparin, EDTA. The serum or plasma should be separated from the coagulated or packed cells by centrifugation. Samples may be shipped at room temperature and then stored refrigerated at 2-8°C if testing is to take place within one week after collection. If testing is to take place later than one week, samples should be stored at -20°C. Avoid repeated freeze/thawing.

9. Assay Procedure

- Allow all reagents to reach room temperature before use. Gently mix all liquid reagents prior to use.
- All standards, controls and samples should be run in duplicate for confirmation of reproducibility.
- A standard curve must be run with each assay.
- If particulate matter is present in the analyte, centrifuge or filter prior to analysis.
- Maintain a consistent order of components and reagents addition from well to well. This ensures equal incubation times for all wells.
- Determine the number of 16-well strips needed for assay. Insert these in the flame(s) for current use (Re-bag extra strips and frame. Refrigerate for further use).
- For the standard curve, add 100 μl of the standard to the appropriate microtiter wells. Add 100 μl of the *Standard/Sample/secondary antibody Dilution Buffer* to zero wells.
- 3) Serum and plasma require **at least 4 fold dilution** in the *Standard/Sample Dilution Buffer*. And add 100 μl of samples to each wells.
- 4) Cover the plate with the plate cover and incubate for 2 hours at 37°C.
- 5) Thoroughly aspirate or decant the solution from the wells. Wash the wells 3 times (See "Directions for washing").
- 6) Pipette 100 µl of "Working Secondary Antibody Solution" into each well.
- 7) Cover the plate with the plate cover and incubate for 1 hour at 37°C.
- Thoroughly aspirate or decant the solution from the wells. Wash the wells 3 times (See "Directions for washing").
- 9) Add 100 µl "Working Streptavidin HRP Solution" to each well.
- 10) Cover the plate with the plate cover and incubate for 30 minutes at 37°C.
- 11) Thoroughly aspirate or decant the solution from the wells. Wash the wells 3 times (See "Directions for washing").
- 12) Add 100 µl of Substrate to each well. The liquid in the wells should begin to turn blue.
- 13) Incubate the plate at room temperature.
- Do not cover the plate with aluminum foil, or color may develop.
 - The incubation time for chromogen substrate is often determined by the microtiter plate reader used. O.D. values should be monitored and the substrate reaction stopped before O.D. of the positive wells exceeds the limits of the instrument. O.D. values at 450nm can only be read after the Stop Solution has been added to each well.
- Because the *Substrate* is light sensitive, avoid the remained *Substrate* solution prolonged exposure to light.

- Typically, reaction is stopped 5~10 minutes after treatment of Substrate, but this time can be adjusted as the user desires.
- 14) Add 100 μl of *Stop Solution* to each well. The solution in the wells should change from blue to yellow.
- 15) Read the absorbance of each well at 450nm. Read the plate within 20 minutes of adding the *Stop Solution*.
- 16) Plot on graph paper the absorbance of the standard against the standard concentration (Optimally, the background absorbance can be subtracted from all data points, including standards, unknowns and controls, prior to plotting.). Draw a smooth curve through these points to construct the standard curve.
- 17) Read the mouse TNF-α concentrations for the unknown samples and controls from the standard curve plotted in step 18. Multiply value(s) obtained for the unknown sample by the dilution factor (Samples producing signals greater than that of the highest standard should be further diluted in the Standard/Sample Dilution Buffer).

10. Characteristics

1) Typical result

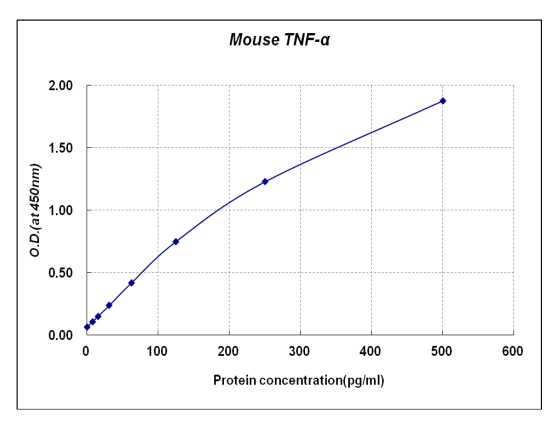
The standard curve below is for illustration only and **should not be used** to calculate results in your assay.

Standard	Optical Density
Mouse TNF-α (pg/ml)	(at 450nm)
0	0.061
7.8125	0.107
15.625	0.149
31.25	0.236
62.5	0.417
125	0.748
250	1.228
500	1.878

A standard curve must be run with each assay.

< Limitations >

- Do not extrapolate the standard curve beyond the 500 pg/ml standard point.
- Other buffers and matrices have not been investigated.
- The rate of degradation of native mouse TNF- α in various matrices has not been investigated.



(TMB reaction incubate at room temperature for 5 min)

2) Sensitivity

The minimal detectable dose of mouse TNF- α was calculated to be 2.535pg/ml, by subtracting three standard deviations from the mean of 12 zero standard replicates (ELISA buffer, S0) and intersecting this value with the standard curve obtained in the same calculation.

N	1	2	3	4	5	6	7	8	9	10	11	12	
ZERO	0.062	0.057	0.058	0.059	0.057	0.059	0.063	0.063	0.061	0.062	0.069	0.065	

Average	SD	LLD	LLD mean(pg/ml)
0.061	0.004	0.072	2.535

3) Specificity

The following substances were tested and found to have no cross-reactivity: Human TNF- α , Rat TNF- α

4) Precision

1 Within-Run (Intra-Assay)

_		(n=	=12)
Mean (pg/ml)	SD	CV (%)	
62.012	2.468	3.981	
126.053	5.491	4.356	
252.043	14.593	5.790	
503.300	18.466	3.669	

2 Between-Run (Inter-Assay)

(n=12)

Mean (pg/ml)	SD	CV (%)
61.933	2.545	4.108
127.518	4.183	3.280
248.248	12.376	4.985
499.216	23.268	4.661

5) Recovery

Recovery on addition is 93.764~101.178% (mean 97.758%)

Added Analyte (pg/ml)	Serum(1/4)+added analyte (450nm)	Serum(1/4)(450nm) +added analyte(450nm)	Recovery (%)
62.869	77.821	76.790	98.676
122.651	137.603	134.047	97.416
247.322	262.274	245.918	93.764
456.690	471.642	477.196	101.178

11. Troubleshooting

Problem	Possible Cause	Solution
	Insufficient washing	• Increase number of washes
		 Increase time of soaking
		between in wash
High signal and background	• Too much streptavidin-HRP	Check dilution, titration
in all wells	• Incubation time too long	• Reduce incubation time
	• Development time too long	• Decrease the incubation time
		before the stop solution is
		added
	Reagent added in incorrect	• Review protocol
	order, or incorrectly prepared	
	• Standard has gone bad	• Check the condition of stored
No signal	(If there is a signal in the	standard
	sample wells)	
	• Assay was conducted from an	• Reagents allows to come to
	incorrect starting point	20~30°C before performing assay
	 Insufficient washing 	• Increase number of washes
	–unbound streptavidin-HRP	carefully
Too much signal – whole	remaining	
plate turned uniformly blue	• Too much streptavidin-HRP	Check dilution
place turned uniformity blue	• Plate sealer or reservoir	• Use fresh plate sealer and
	reused, resulting in presence	reagent reservoir for each
	of residual streptavidin -HRP	step
Standard curve achieved but	• Plate not developed long	• Increase substrate solution
poor discrimination between	enough	incubation time
point	• Improper calculation of	Check dilution, make new
point	standard curve dilution	standard curve
No signal when a signal is	• Sample matrix is masking	 More diluted sample
expected, but standard curve	detection	recommended
looks fine		
Samples are reading too high,	• Samples contain protein levels	• Dilute samples and run
but standard curve is fine	above assay range	again
	• Uneven temperature around	 Avoid incubating plate in
Edge effect	work surface	areas where environmental
Luge effect		conditions vary
		• Use plate sealer

12. Reference

- 1) Berry M, Brightling C, Pavord I, and Wardlaw A., 2007, Curr Opin Pharmacol. 7(3):279-282.
- 2) Anderson GM, Nakada MT, and DeWitte M, 2004, Curr Opin Pharmacol. 4(4):314-320.
- 3) Palladino MA, Bahjat FR, Theodorakis EA, and Moldawer LL., 2003, Nature Reviews Drug Discovery. 2:736-746

Ordering Information

For orders, please contact :

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