GE Healthcare

Amersham
High Sensitivity Tumor
Necrosis Factor-Alpha ((h)
TNFα) Human,
Biotrak ELISA System

Product Booklet

Code: RPN2788



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1. Legal

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Previously published 1995

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2. Handling

2.1. Safety warnings and precautions

Warning: For research use only. Not recommended or intended for diagnosis of disease in humans or animals. Do not use internally or externally in humans or animals

All chemicals should be considered as potentially hazardous. We therefore recommend that this product is handled only by those persons who have been trained in laboratory techniques and that it is used in accordance with the principles of good laboratory practice. Wear suitable protective clothing such as laboratory overalls, safety glasses and gloves. Care should be taken to avoid contact with skin or eves. In the case of contact with skin or eyes wash immediately with water. See material safety data sheet(s) and/or safety statement(s) for specific advice.

2.2. Storage

Store at 2-8°C.

2.3. Expiry

The expiry date is stated on the package and will normally be at least 4 weeks from the date of despatch.

3. Components of the assay system

This pack contains the following assay components, sufficient material for 96 wells.

All reagents are stored refrigerated at 2–8°C. Refer to the expiry date on the kit box.

(h)TNF α microplate – 96 well polystyrene microplate (12 strips of 8 wells) coated with an antibody against (h)TNF α .

Biotinylated antibody reagent – antibody against (h)TNF $\!\alpha$ conjugated to biotin, 11 ml.

(h)TNF α standard – 2 vials of recombinant human (h)TNF α , lyophilized.

Amdex[™] amplification reagent – lyophilized.

Standard diluent - 12 ml, 2 vials.

Wash buffer concentrate – 30-fold concentrated solution, 50 ml.

Pre-mixed TMB substrate reagent - 12-13 ml.

Stop solution – <1% sulfuric acid, 13-15 ml.

Plate covers - 6 adhesive strips.



4. Description

The BiotrakTM high sensitivity human tumor necrosis factor-alpha ELISA system from GE Healthcare provides a simple, specific, reliable and precise quantitative determination of (h)TNF α in cell culture supernatants, plasma and serum.

The assay system is based on a solid phase ELISA, which utilizes an antibody for (h)TNF α bound to the wells of a microplate together with a biotinylated antibody to (h)TNF α and Amdex amplification reagent. Although the Biotrak (h)TNF α immunoassay contains recombinant (h)TNF α and antibodies raised against recombinant (h) TNF α , it has been shown to quantitate accurately both natural (h) TNF α and recombinant (h)TNF α .

(h)TNF α can be measured in the range 0.31–10 pg/ml (0.031–1 pg/well) in less than 4 hours using the protocol provided with the kit. Each pack contains sufficient material for 96 wells. If one standard curve is constructed, 41 unknowns can be measured in duplicate.

- High sensitivity 0.1 pg/ml
- · Same day protocol
- · Pre-coated plate
- Specific for (h)TNFα

4.1. Summary of the assay

This assay employs the quantitative 'sandwich' enzyme immunoassay technique. An antibody specific for (h)TNF α antibody has been coated on the microplate provided in the kit. Samples are pipetted into the wells and the (h)TNF α , if present, is bound by the TNF α immobilized antibody. A biotinylated antibody reagent is added to the wells and allowed to bind to any TNF α bound by the immobilized antibody in the first incubation. After washing away

any unbound biotinylated antibody an Amdex amplification reagent is added, any TNF α which was bound by both the immobilized and the biotinylated antibody will be bound by the amplification reagent. Following a wash to remove unbound amplification reagent, a substrate solution is added to the wells and color develops in proportion to the amount of (h)TNF α bound in the initial step.

The Amdex amplification reagent is a high performance conjugate based on a novel chemistry that utilizes a hydrophilic straight chain dextran backbone to which many hundreds of horseradish peroxidase molecules are covalently coupled, together with, on average, ten streptavidin molecules. The result is a multifunctional conjugate with a significantly enhanced activity and with well controlled non-specific binding properties.

In addition to the samples to be tested, a series of wells is prepared using known concentrations of the human TNF α standard. A curve, plotting the optical density versus the concentration of the standard well, is prepared. By comparing the optical density of the samples to this curve, the concentration of the (h)TNF α in the unknown samples is then determined.

5. Critical parameters

- Allow samples and all reagents to reach room temperature prior to performing the assays. Do not use water baths to thaw samples or reagents.
- 2. Mix samples and all reagents thoroughly before use.
- Avoid excessive foaming of reagents. Also avoid exposure of reagents to excessive heat or light during storage and incubation.
- 4. Avoid handling the tops of the wells both before and after filling.
- 5. Standards and samples should be assayed in duplicate.
- 6. Run a separate standard curve for each assay.
- The total dispensing time for each plate should not exceed 20 minutes.
- **8.** Use only coated wells from the same reagent batch for each assay. Also do not mix reagents from different kit lots.
- 9. It is important that all the wells are washed thoroughly and uniformly. If using an automatic washer, check operation of heads before starting. If washing by hand ensure wells are completely filled at each wash.
- Timings in this assay are critical and should be adhered to strictly. Failure to do so could alter optical densities significantly.
- 11. A small amount of precipitate may be present in some vials. It will not affect assay performance and should be ignored.

6. Additional materials and equipment required

The following materials and equipment are required:

- Pipettes or pipetting equipment with disposable tips (20 μ l, 100 μ l, 400 μ l, 500 μ l, 1.00 ml and 5.00 ml)
- Disposable polypropylene test tubes do not use polystyrene, polycarbonate or glass
- Measuring cylinder, 2 L
- Distilled or deionized water
- Plate reader capable of reading at 450 nm
- Plate shaker

Optional equipment

Assays may be performed with commercially available microplate washers to aid convenience and assay throughput.

7. Sample preparation

Serum, plasma and cell culture supernatants

Serum, plasma and culture supernatant samples that are to be assayed within 24 hours should be stored at 2–8°C. Specimens to be stored for longer periods of time should be frozen at -70°C to avoid loss of activity. Avoid freezing and thawing samples more than once. Test samples should be assayed in duplicate each time the ELISA is performed, $100 \, \mu l$ of sample per well is required in this assay.

The measurement of cytokines in serum and plasma has been reported to be affected by non-specific matrix effects which may vary between samples from different individuals (1–3). Dilution of such samples in the diluent supplied may help to reduce these interference effects.

Dilution of test samples

If it is suspected that the (h)TNF α concentration of a sample exceeds the highest point of the standard curve, prepare one or more five-fold dilutions of the test sample. Mix thoroughly between dilutions and before assaying.

It remains the investigators responsibility to validate the chosen sample dilution.

8. Assay procedure

8.1. Reagent preparation

Wash buffer concentrate

Any precipitate formed during storage will redissolve upon dilution. Dilute the contents of the bottle to 1500 ml using distilled water. Store at 2–8°C until the expiry date of the kit. Do not use wash buffer if it becomes visibly contaminated on storage.

Amdex amplification reagent

Reconstitute the Amdex reagent in 11 ml of distilled or deionized water approximately 15 minutes before use. Reconstituted reagent may be stored at -15° C to -30° C for up to 1 week.

(h)TNFα standard

Reconstitute 1 vial of (h)TNF α standard with distilled or deionized water. Reconstitution volume is shown on the vial label. Mix by gently inverting the vial.

It is important that the diluent selected for reconstitution and dilution of the standard reflects the environment of the samples being measured. Standard diluent will be suitable for serum or plasma measurements. If your samples are cell culture supernatants, the culture media will be suitable for preparation of the standard curve.

Testing of RPMI with different lots and concentrations of fetal bovine serum has shown that this ELISA is not adversely affected by culture medium. Therefore when culture supernatants, serum and plasma samples are assayed on the same plate, standards prepared in standard diluent may be used.

8.2. Preparation of standard curve

Preparation of working stock solution

1. Pipette 400 μl of standard diluent into a polypropylene tube.

- 2. Add to this tube 100 μ l of reconstituted (h)TNF α standard and mix thoroughly.
- **3.** This is the 500 pg/ml stock solution from which the working standards are prepared.

Preparation of working standards

- **1.** Label 6 polypropylene tubes, 0.31, 0.63, 1.25, 2.5, 5 and 10 pg/ml.
- 2. Pipette 980 µl of standard diluent into the 10 pg/ml tube.
- 3. Pipette 500 µl of standard diluent into all remaining tubes.
- 4. Into the 10 pg/ml tube pipette 20 μ l of working stock solution of (h) TNF α and mix thoroughly.
- 5. Transfer 500 μ l from the 10 pg/ml tube to the 5 pg/ml tube and mix thoroughly.
- 6. Repeat this doubling dilution successively with the remaining tubes.
- 7. 100 μ l aliquots from each serial dilution will give rise to 6 standard levels of (h)TNF α ranging from 0.31 to 10 pg/ml.

NOTE: Working standards should be freshly prepared before each assay, and not re-used.

Running partial plates

This ELISA provides the flexibility to run two partial plates on separate occasions. Decide the number of strips you wish to run, leaving the strips to be used in the frame. Remove the unnecessary strips and store them in the foil pouch with the desiccant provided at 2–8°C, making sure the foil pouch is sealed tightly.

When adding the TMB substrate reagent, pour out from the bottle ONLY the amount needed to run the first half plate. Do not combine left over substrate with that reserved for the second half of the plate. Care must be taken to ensure that the remaining TMB substrate reagent is not contaminated. If the substrate reagent is bright blue prior to use, it has been contaminated. DO NOT USE.

8.3. Assay protocol

- Prepare assay reagents and working standards as described in the previous sections.
- 2. Set up the microplate with sufficient wells to enable the running of all standards and samples as required (see figure 1).
- Remove excess microplate strips from the frame and store in the resealable foil bag.
- 4. Pipette 100 μl of standard diluent or cell culture medium (see section on reagent preparation) into NSB wells.
- 5. Pipette 100 µl of standard into the appropriate wells.
- 6. Pipette 100 µl of sample into the appropriate wells.
- 7. Cover the plate with the adhesive strip provided and incubate for 1 hour at room temperature (20–25°C) with continuous shaking.
- 8. Aspirate or decant each well and wash, repeating the process three times for a total of four washes. Wash thoroughly by completely filling each well with wash buffer using a washbottle, or manifold dispenser. Complete removal of liquid at each step is essential for good performance. After the last wash, remove any remaining wash buffer by inverting the plate and blotting it against clean paper towelling.
- 9. Pipette 100 µl of the biotinylated antibody reagent into all wells. Cover with a new adhesive strip and incubate for 1 hour at room temperature (20–25°C) with continuous shaking.
- 10. Repeat the aspiration/wash step as in step 8.
- 11. Pipette 100 µl of Amdex amplification reagent to all wells. Cover with a new adhesive strip and incubate for 30 minutes at room temperature (20–25°C) with continuous shaking.
- 12. Repeat the aspiration/wash step as in step 8.

- 13. Pipette 100 µl of TMB substrate solution into all wells, incubate for 1 hour at room temperature (20–25°C) with continuous shaking. If the substrate reagent is bright blue prior to use, do not use. Do not cover the plate with aluminium foil or an adhesive strip.
- **14.** Pipette 100 µl of stop solution into all wells.
- **15.** Determine the optical density of each well within 30 minutes, using a spectrophotometer set to 450 nm.

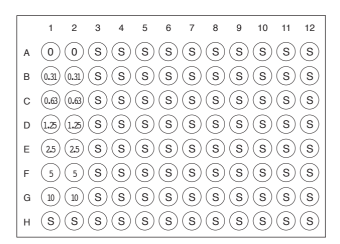


Figure 1. Recommended positioning of standard (0.31–10 pg/ml) and sample (S) wells.

Table 1. Assay protocol (all volumes are in microlitres)

	Zero	Standards	Samples
	standard		
Standard	-	100	-
Standard diluent or cell			
culture media*	100	-	-
Sample	-	-	100
Cover plate, incubate at s	room tempera naker for 1 hou		on a plate
Aspirate/decant and wash	thoroughly all buffer	wells four times	s with wash
Biotinylated			
antibody reagent	100	100	100
Cover plate, incubate at I sh	room tempera naker for 1 hou		on a plate
Aspirate/decant and wash	thoroughly all buffer	wells four times	s with wash
Amdex amplification reager	nt 100	100	100
Cover plate, incubate at a	room tempera ker for 30 minu		on a plate
Aspirate/decant and wash	thoroughly all buffer	wells four times	s with wash
Substrate	100	100	100
Incubate at room temperat	ure (20-25°C)	on a plate shake	er for 1 hour
Stop solution	100	100	100
Determine optical de	nsity at 450 n	m within 30 mir	nutes.
* Use 100 ul of cell sulture m			

 $^{^{\}star}$ Use 100 μl of cell culture media if your standard curve is diluted in cell culture media. See section on reagent preparation.

9. Data processing

9.1. Calculation of results

Average the duplicate readings for each standard and sample and subtract the zero standard optical density.

Plot the optical density for the standards versus the concentration of the standards and draw the best curve. The data can be linearized by using a log/log plot and regression analysis can be applied to the log transformation.

Figure 2 shows such a plot of the data from table 2. The standard curve is provided for illustration only. A standard curve should be generated for each set of samples to be assayed. This allows for the measurement of 41 unknowns in duplicate.

9.2. Typical assay data

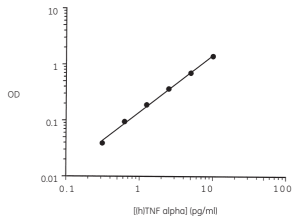


Figure 2. Standard curve

The following data (table 2) were obtained for a standard curve using the protocol provided.

Table 2. Typical assay data

Tube	Optical density	Zero standard subtracted
Zero standard	0.075	-
0.31 pg/ml standard	0.114	0.039
0.63 pg/ml standard	0.169	0.094
1.25 pg/ml standard	0.264	0.189
2.5 pg/ml standard	0.439	0.364
5 pg/ml standard	0.775	0.700
10 pg/ml standard	1.465	1.390

When running these high sensitivity assays some variation in OD values may be observed. However control and sample values will not be affected

10. Additional information

10.1. Specificity

This assay recognizes both natural and recombinant (h)TNF α . It does not cross react with human IL-1 α , IL-1 β , IL-2, IL-3, IL-4, IL-6, IL-7, IL-8, TNF β , IFN γ , IFN α or mouse TNF α .

10.2. Calibration

The standards in this ELISA are calibrated to the WHO reference lot 87/650.

1 pg of Biotrak standard = 3 pg of WHO standard = ~ 0.12 WHO unit.

10.3. Reproducibility

Within-assay precision

The within-assay coefficient of variation of the ELISA has been determined to be <10%

Between-assay precision

The between-assay coefficient of variation of the ELISA has been determined to be <10%.

Precision profile

A precision profile was generated by preparing replicates of each of the standards and calculating the standard deviation and coefficient of variation at each concentration

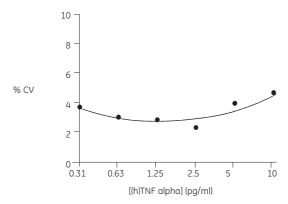


Figure 3. Precision profile

10.4. Sensitivity

The minimum detectable dose of (h)TNF α was determined to be 0.1 pg/ml (0.01 pg/well), by adding two standard deviations to the optical density value of zero and calculating the corresponding concentration from the standard curve.

10.5. Parallelism

The linearity of dilution was determined by serially diluting six different positive samples. The dilutions were run in the ELISA and 'observed' doses were plotted against 'expected' doses.

10.6. Expected values

Normal levels of (h)TNF α in serum samples are in the range 0.9–4.9 pg/ml.

Normal levels of (h)TNF α in plasma samples are in the range 0–1.8 pg/ml.

10.7. Background and references

The tumor necrosis factor family contains two related cytokines, TNF α and TNF β , that demonstrate overlapping activities on a variety of biological assay systems. TNF α is an endogenous pyrogen and cytotoxic factor that plays a central role in the regulation of inflammation, immunity and host defence. The rapid production of this potent cytokine in response to microbial infections can lead to both beneficial effects locally and lethal effects systemically, highlighting the dual nature of TNF α .

TNF α is a 157 amino acid unglycosylated protein with a molecular weight of 17 kDa (determined by SDS PAGE). TNF α is synthesized as a 26 kDa peptide and is processed by the removal of a 76 amino acid signal peptide. Biologically active TNF α occurs as a trimer in solution. TNF α is produced by numerous cell types including: astrocytes, B and T cells, endothelial cells, epidermal cells, fibroblasts, Kupffer cells, mast cells, monocytes and myeloid cell lines, NK cells, smooth muscle cells and several tumor lines.

Two distinct TNF receptors have been identified; type I (55–60 kDa) and type II (75–80 kDa) which despite the lack of homology extracellularly, both TNFR-I and TNFR-II can bind TNF α and TNF β with affinities ranging from 1–10 x 10⁻¹⁰ M. TNFR are expressed by virtually all mammalian cells examined to date, with the exception of erythrocytes. Cell surface receptor densities range from 2000–50 000 sites per cell. TNF α can be induced by a wide range of agents including: bacteria, viruses, protozoa, LPs, S.aureus, IL-1, IL-2, TNF, IFN, GM-CSF, PAF, C5 α , substance P, anti-TCR antibodies, tumor cells, phorbol esters, indomethacin and zymosan. Agents that suppress TNF α production include: dexamethasone, cyclosporin A, PGE $_{2}$, IL-4, IL-10 TGF β , lipoxygenase inhibitors, PAF receptor antagonists and vitamin D $_{2}$.

More than any other cytokine, with the exception of IL-1, TNF has shown to be operative in the pathophysiological processes of numerous chronic and acute diseases. The development of TNF antagonists, including soluble TNFR and anti-TNF antibodies, may therefore lead to more effective therapies for the management of septic shock and other inflammatory diseases. Since TNF α appears to be beneficial at the local level in a variety of systems, recombinant TNF α may also be of value pending the development of targeted delivery systems.

- 1. Heney, D. and Whicher, J.T., Ann. Clin. Biochem. 32, 358 (1995).
- 2. de Kossodo, S. et al., J. Immunuol. Methods 182, 107 (1995).
- 3. Wood, W.G., Methods of Immunological Analysis, Massayeff, R.F. et al., (ED) Vol. 1, 604 (1992).

10.8. Related products

Biotrak range of human cytokine ELISA systems	
Interleukin- 1α [(h)IL- 1α]	RPN2750
Interleukin-1 β [(h)IL-1 β]	RPN2751
Interleukin-2 [(h)IL-2]	RPN2752
Interleukin-6 [(h)IL-6]	RPN2754
Interleukin-10 [(h)IL-10]	RPN2755
Granulocyte-macrophage colony stimulating factor	
[(h)GM-CSF]	RPN2756
Interferon-gamma [(h)IFNγ]	RPN2757
Tumor necrosis factor-alpha [(h)TNFα]	RPN2758
Interferon-alpha [(h)IFNα]	RPN2759
Interleukin-8 [(h)IL-8]	RPN2764
Interleukin-12, p40/70 [(h)IL-12] p40/70	RPN2765

Biotrak range of high sensitivity human cytokine ELISA systems Interleukin- 1α [(h)IL- 1α] RPN2780 Interleukin-4 [(h)IL-4] RPN2783

Interleukin-6 [(h)IL-6]	RPN2784
Interleukin-10 [(h)IL-10]	RPN2785
Interferon-gamma [(h)IFNγ]	RPN2787
Interferon-alpha [(h)IFNα]	RPN2789
Biotrak range of mouse cytokine ELISA systems	
Interleukin-1 β [(m)IL-1 β]	RPN2720
Interleukin-4 [(m)IL-4]	RPN2712
Tumor necrosis factor-alpha [(m)TNFα]	RPN2718
Range of unlabelled and radiolabelled growth factors and cytokines	
Cell proliferation assay system and reagents	
Cell proliferation kit (for immunocytochemical/	RPN20
immunohistochemical measurement)	
Monoclonal anti-bromodeoxyuridine	RPN202
Cell proliferation labelling reagent	RPN201

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