

EKA51981 - 96 Tests TGFbeta-R2 Colorimetric Sandwich ELISA Kit Species Reactivity: Human

FOR RESEARCH USE ONLY. NOT FOR USE IN DIAGNOSTIC PROCEDURES

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INSTRUCTION MANUAL

This manual may be updated as a result of continuous improvements. Please always refer to the hard copy manual included in the kit for your experiment.

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INTRODUCTION

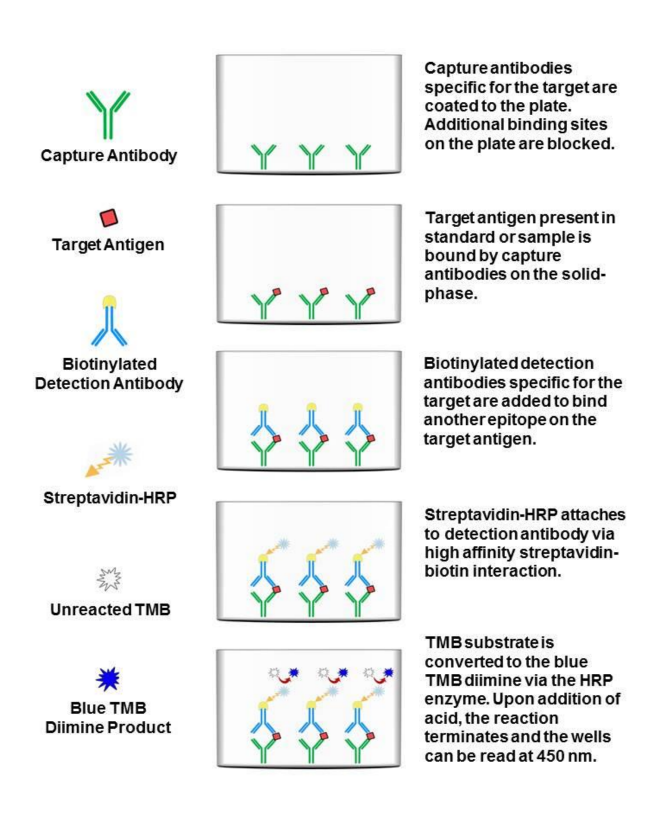
TGF β -R2 is a transmembrane serine/threonine kinase that forms with the TGF-beta type I serine/threonine kinase receptor, TGFBR1, the nonpromiscuous receptor for the TGF-beta cytokines TGFB1, TGFB2 and TGFB3. It transduces the TGFB1, TGFB2 and TGFB3 signal from the cell surface to the cytoplasm and is thus regulating a plethora of physiological and pathological processes including cell cycle arrest in epithelial and hematopoietic cells, control of mesenchymal cell proliferation and extracellular differentiation. wound healing, matrix production. immunosuppression and carcinogenesis. The formation of the receptor complex composed of 2 TGFBR1 and 2 TGFBR2 molecules symmetrically bound to the cytokine dimer results in the phosphorylation and the activation of TGFRB1 by the constitutively active TGFBR2. Activated TGF_β-R1 phosphorylates SMAD2 which dissociates from the receptor and interacts with SMAD4. The SMAD2-SMAD4 complex is subsequently translocated to the nucleus where it modulates the transcription of the TGFbeta-regulated genes. This constitutes the canonical SMAD-dependent TGF-beta signaling cascade. This receptor is also involved in noncanonical, SMAD-independent TGF-beta signaling pathways. TGFβ-R2 is a homodimer that interacts with TGFB1, TGFB2 and TGFB3 homodimeric ligands which assemble a functional receptor composed of two TGFBR1 and TGFBR2 heterodimers to form a ligand-receptor heterohexamer. The respective affinity of TGFRB1 and TGFRB2 for the ligands may modulate the kinetics of assembly of the receptor and may explain the different biological activities of TGFB1, TGFB2 and TGFB3. TGFβ-R2 also interacts with DAXX, TCTEX1D4, and ZFYVE9. ZFYVE9 recruits SMAD2 and SMAD3 to the TGF-beta receptor. The receptor is phosphorylated on a Ser/Thr residue in the cytoplasmic domain. Defects in TGF_β-R2 are the cause of hereditary non-polyposis colorectal cancer type 6 (HNPCC6), esophageal cancer (ESCR), Loeys-Dietz syndrome type 1B (LDS1B), Loeys-Dietz syndrome type 2B, and aortic aneurysm familial thoracic type 3 (AAT3).

Source: Entrez Gene; Swiss-Prot

ASSAY PRINCIPLES

The Human TGFbeta-R2 ELISA Kit contains the components necessary for quantitative determination of natural or recombinant Human TGFbeta-R2 concentrations within any experimental sample including cell lysates, serum and plasma. This particular immunoassay utilizes the quantitative technique of a "Sandwich" Enzyme-Linked Immunosorbent Assay (ELISA) where the target protein (antigen) isbound in a "sandwich" format by the primary capture antibodies coated to each well-bottom and the secondary detection antibodies added subsequently by the investigator. The capture antibodies coated to the bottom of each well are specific for a particular epitope on Human TGFbeta-R2 while the user-added detection antibodies bind to epitopes on the captured target protein. Amid each step of the procedure, a series of wash steps must be performed to ensure the elimination of non-specific binding between proteins to other proteins or to the solid phase. After incubation and "sandwiching" of the target antigen, a peroxidase enzyme is conjugated to the constant heavy chain of the secondary antibody (either covalently or via Avidin/Streptavidin-Biotin interactions), allowing for a colorimetric reaction to ensue upon substrate addition. When the substrate TMB (3, 3', 5, 5'-Tetramethylbenzidine) is added, the reaction catalyzed by peroxidase yields a blue color that is representative of the antigen concentration. Upon sufficient color development, the reaction can be terminated through addition of Stop Solution (2 N Sulfuric Acid) where the color of the solution will turn yellow. The absorbance of each well can then be read by a spectrophotometer, allowing for generation of a standard curve and subsequent determination of protein concentration.







ASSAY RESTRICTIONS

- This ELISA kit is intended for research purposes only, NOT diagnostic or clinical procedures of any kind.
- Materials included in this kit should NOT be used past the expiration date on the kit label.
- Reagents or substrates included in this kit should NOT be mixed or substituted with reagents or substrates from any other kits.
- Variations in pipetting technique, washing technique, operator laboratory technique, kit age, incubation time or temperature may cause differences in binding affinity of the materials provided.
- The assay is designed to eliminate interference and background by other cellular macromolecules or factors present within any biological samples. However, the possibility of background noise cannot be fully excluded until all factors have been tested using the assay kit.
- Individual results may vary due to differences in technique, plasticware and water sources.

Component	Quantity Per Plate	Container -	
Microstrips Coated w/ Capture Antibody	12 x 8-Well Microstrips		
Protein Standard	Lyophilized	Red	
Biotinylated Detection Antibody	Lyophilized	Yellow	
400x Streptavidin-HRP	30 µl	Blue	
Wash Buffer (15x)	50 ml	Clear	
Assay Diluent	50 ml	Clear	
Ready-to-Use Substrate	12 ml	Brown	
Stop Solution	12 ml	Clear	
Adhesive Plate Sealers	2 Sheets	-	
Technical Manual	1 Manual	-	

MATERIALS INCLUDED

ADDITIONAL MATERIALS REQUIRED

The following materials and/or equipment are NOT provided in this kit but are necessary to successfully conduct the experiment:

- Microplate reader able to measure absorbance at 450 nm (with correction wavelength set to 540 nm or 570 nm)
- Micropipettes with capability of measuring volumes ranging from 1 μI to 1 ml
- Distilled, deionized, and or purified water (recommended TOC 1-50 ppb, M Ω -cm 18.0)
- Squirt bottle, manifold dispenser, multichannel pipette reservoir or automated microplate washer
- Graph paper or computer software capable of generating or displaying logarithmic functions
- Absorbent paper or vacuum aspirator
- Test tubes or microfuge tubes capable of storing ≥1 ml
- Bench-top centrifuge (optional)
- Bench-top vortex (optional)
- Orbital shaker (optional)

HEALTH AND SAFETY PRECAUTIONS

- Reagents provided in this kit may be harmful if ingested, inhaled or absorbed through the skin. Please carefully review the MSDS for each reagent before conducting the experiment.
- Stop Solution contains 2 N Sulfuric Acid (H₂SO₄) and is an extremely corrosive agent. Please wear proper eye, hand and face protection when handling this material. When the experiment is finished, be sure to rinse the plate with copious amounts of running water to dilute the Stop Solution prior to disposing the plate.

STORAGE INFORMATION

Note: If used frequently, reagents may be stored at 4°C.

Unopened Kits: Store at 4°C for 6 months.

Component	Storage Time	Storage Information	
Microstrips Coated w/ Capture Antibody			
400x Streptavidin-HRP	InformationInformationied w/ odyInformationin-HRP6 Months15x)6 Months15x)6 Monthsint4int4int4ardLyophilized: 6 MonthsardReconstituted: 1 Month		
Wash Buffer (15x)	6 Months	hs 4°C	
Assay Diluent	Lyophilized: 6 Months		
Ready-to-Use Substrate	te		
Stop Solution			
Protein Standard	Lyophilizad: 6 Months		
Biotinylated Detection Antibody	otinylated Detection Reconstituted: 1 Month 4°C		
Adhesive Plate Sealers	-	-	
Technical Manual	-	-	

SAMPLE STORAGE AND PREPARATION

If samples are to be used within 24 hours, aliquot and store at 4°C. If samples are to be used over a long period of time, aliquot and store between -20°C and -80°C, depending on the duration of storage.

Note: Samples containing a visible precipitate or pellet must be clarified prior to use in the assay.

Caution: Avoid repeated freeze/thaw cycles to prevent loss of biological activity of proteins in experimental samples.

Cell Lysate and Supernatants

Remove large cell components via centrifugation and perform the assay. Cell lysates and supernatants require a dilution using Assay Diluent. A serial dilution may be performed to determine a suitable dilution factor for the sample.

Serum

Allow samples to clot in a serum separator tube (SST) for 30 minutes. After sufficient clotting, centrifuge at 1000 x g for 15 minutes and remove serum from SST in preparation for the assay. A serial dilution may be performed to determine a suitable dilution factor for the sample. For serum sample dilutions refer to Serum and Plasma Sample Dilution Protocol.

Plasma

Use heparin, citrate or EDTA as an anticoagulant to gather plasma from original biological sample. After collection of the plasma, centrifuge for 15 minutes at 1000 x g. This step must be performed within 30 minutes of plasma collection. A serial dilution may be performed to determine a suitable dilution factor for the sample. For plasma sample dilutions refer to Serum and Plasma Sample Dilution Protocol.



Serum and Plasma Sample Dilution Protocol

- a. Dilute the serum or plasma samples with PBS supplemented with 10-50% animal serum (Serum/Plasma Diluent).
- b. Reconstitute and dilute the Protein Standards using the Serum/Plasma Diluent, instead of Assay Diluent, so it reflects the environment of the samples being measured.
- c. Reconstitute the Biotin-Conjugated Detection Antibody in Assay Diluent and dilute the Streptavidin-HRP in Assay Diluent. Do not use the Serum/Plasma Diluent to reconstitute or dilute the Detection Antibody or Streptavidin-HRP.

	1	2	3	4	5	6
Α	Standard (High Point)	Standard (High Point)	Standard (High Point)	Sample	Sample	Sample
в	Standard (1:2)	Standard (1:2)	Standard (1:2)	Sample	Sample	Sample
С	Standard (1:4)	Standard (1:4)	Standard (1:4)	Sample	Sample	Sample
D	Standard (1:8)	Standard (1:8)	Standard (1:8)	Sample	Sample	Sample
Ε	Standard (1:16)	Standard (1:16)	Standard (1:16)	Sample	Sample	Sample
F	Standard (1:32)	Standard (1:32)	Standard (1:32)	Sample	Sample	Sample
G	Standard (1:64)	Standard (1:64)	Standard (1:64)	Sample	Sample	Sample
Н	Negative Control	Negative Control	Negative Control	Sample	Sample	Sample

SAMPLE EXPERIMENT LAYOUT



IMMUNOASSAY PROTOCOL

Note: Spin down the Protein Standard, Biotin-Conjugated Detection Antibody vials before opening. If possible, all incubation steps should be performed on an orbital shaker to equilibrate solutions when added to the microplate wells. Also, all provided solutions should be at ambient temperature prior to use.

Reconstitution of Provided Materials

- **1.** Reconstitute the Protein Standard in 83µl of Assay Diluent for a concentration of 110ng/ml. **Note:** If working with serum or plasma, see page 9 prior to reconstitution.
- 2. Reconstitute the Biotin-Conjugated Detection Antibody in 55µl of Assay Diluent for a concentration of 36ug/ml.
- **3.** Dilute the 15x Wash Buffer to 1x Wash Buffer using 14 volumes of ddH₂O and 1 volume of 15x Wash Buffer.

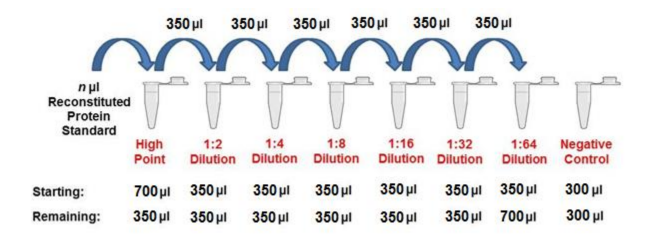
Addition of Known Standard and Unknown Sample to Immunoassay

The Human TGFbeta-R2 ELISA Kit allows for the detection and quantification of endogenous levels of natural and/or recombinant Human TGFbeta-R2 proteins within the range of 16-1000 pg/ml.

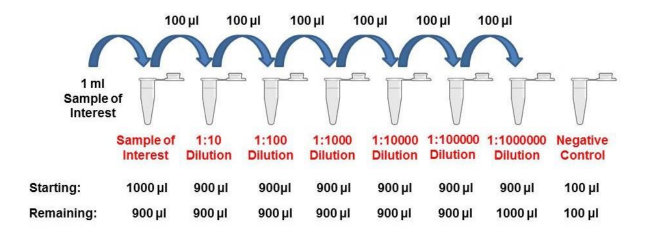
1. Prepare the appropriate diluent for the sample type. Ensure that the Protein Standard is reconstituted and diluted with the same diluent as the sample. Dilute Protein Standard within the range of 1000 pg/ml to 16 pg/ml in a series of microfuge tubes. Mix each tube thoroughly by inverting several times or by vortexing lightly to ensure proper equilibration. Add 100 µl of each serial dilution step into the wells of a specified row or column of the 96-well microtiter plate in duplicate or triplicate and incubate at room temperature for 2 hours. Unknown Samples of Interest can be serial diluted with the appropriate diluent to the concentrations within the detection range of this assay kit and added to the plate at 100 µl per well. See next page for illustration. Blank Control is defined as 100 ul of diluent used to dilute samples and standard per well. Seal the plate.

STANDARD AND SAMPLE SERIAL DILUTION

To obtain serial dilution high point, dilute reconstituted Protein Standard to the maximum concentration for serial dilution by adding $n \mu$ I reconstituted Protein Standard to serial dilution high point tube and then raising the volume to 700 μ I. Shown below is a diagram illustrating an example 2-fold serial dilution on a given reconstituted Protein Standard.



For samples of unknown protein concentrations, serial dilute the experimental sample using Assay Diluent to determine range of detection and acceptable dilutions. Shown below is a diagram illustrating a 10-fold serial dilution on a given Sample of Interest.





Addition of Detection Antibody to Capture Antibody-Bound Samples

- 2. Aspirate the protein standard solution out of the microplate wells. If your lab does not have a vacuum-based aspirator, you may dump the solutions from the microplate into a waste container and blot 3-4 times on a stack of paper towels until most or all of the liquid is removed from the wells. Dilute the 15x Wash Buffer to 1x using ddH₂O. Add 300-400 µl of 1x Wash Buffer to each well and gently shake for 5-7 minutes on an orbital shaker. Perform this wash step 4 times consecutively.
- **3.** After the 4th wash step, dilute the detection antibody solution 1:180 in Assay Diluent to 0.21ug/ml. Mix the test tube either by inverting several times or vortexing to ensure proper equilibration. Ensure that there is enough detection antibody solution for all wells. Add 100 μl of the diluted detection antibody solution into each well, seal the plate and incubate at room temperature for 2 hours.

Conjugation of Streptavidin-HRP to Biotinylated Detection Antibody

- **4.** Remove the detection antibody solution out of the microplate wells by either vacuum-based aspirator or paper towel blotting. Perform 4 consecutive wash steps with gentle shaking between each wash.
- **5.** Dilute the 400x Streptavidin-HRP by 1:400 using Assay Diluent to a 1x Streptavidin-HRP solution.
- **6.** After the 4th wash step, add 100 µl of 1x Streptavidin-HRP solution into each well and incubate at room temperature for 30 minutes. Avoid placing the plate in direct light.

Application of Liquid Substrate for Colorimetric Reaction

7. Remove the 1x Streptavidin-HRP solution out of the microplate wells by either vacuum-based aspirator or paper towel blotting. Prepare the Ready-to-Use Substrate by bringing it to room temperature without exposure to fluorescent or UV light as these may degrade the substrate. Perform 4 consecutive wash steps with gentle shaking between each wash.



- 8. After the 4th wash step, add 100 µl of Ready-to-Use Substrate solution into each well and incubate at room temperature for approximately 15-20 mins. Avoid placing the plate in direct light. Closely monitor the color development as some wells may turn blue very quickly depending on analyte and/or detection antibody-HRP concentrations. The color in the wells should immediately change from blue to yellow.
- **9.** The microplate is now ready to be read by a microplate reader. Within 30 minutes of adding the Stop Solution, determine the optical density (absorbance) of each well by reading the plate with the microplate reader set to 450 nm. If wavelength correction is available, set to 540 nm or 570 nm. If wavelength correction is not available, subtract readings at 540 nm or 570 nm from the readings at 450 nm.

Caution: Readings made directly at 450 nm without correction may be higher and less accurate.

Generation of Standard Curve and Interpretation of Data

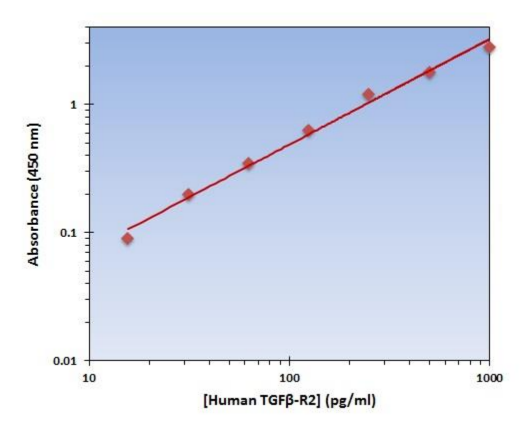
- **10.** Average the duplicate or triplicate readings for each standard, control and sample and subtract the average zero standard optical density.
- 11. Generate a standard curve by using Microsoft Excel or other computer software capable of establishing a 4-Parameter Logistic (4-PL) curve fit. If using Excel or an alternative graphing tool, plot the average optical density values in absorbance units (y-axis) against the known standard concentrations in pg/ml (x-axis).

Note: Only use the values in which a noticeable gradient can be established. Afterwards, generate a best fit curve or "trend-line" through the plotted points via regression analysis.

Note: Shown on the next page is an example of typical data produced by analysis of the standard sample.



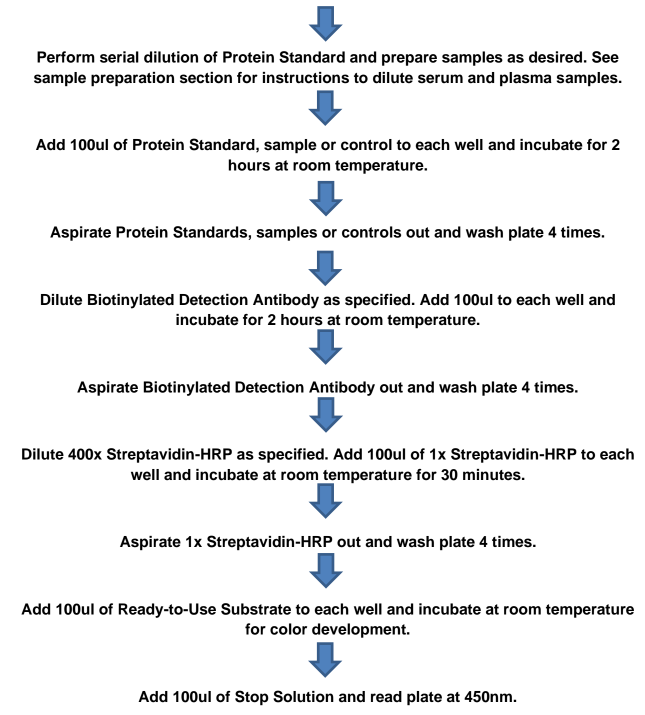
The data and subsequent graph was obtained after performing a cytokine ELISA for Human TGFbeta-R2. Each known sample concentration was assayed in triplicate.





SUMMARIZED PROTOCOL

Reconstitute Biotinylated Detection Antibody and Protein Standard and dilute the 15x Wash Buffer as specified.





SENSITIVITY

The Human TGFbeta-R2 ELISA Kit allows for the detection and quantification of endogenous levels of natural and/or recombinant Human TGFbeta-R2 proteins within the range of 16-1000 pg/ml.

CROSS REACTIVITY AND SPECIFICITY

The Human TGFbeta-R2 ELISA is capable of recognizing both recombinant and naturally produced Human TGFbeta-R2 proteins. The antigens listed below were tested at 50 ng/ml and did not exhibit significant cross reactivity or interference.

- Human: LAP, Latent TGF-beta1, TGFbeta-R1, TGFbeta-R3, TGF-beta1, TGF-beta1.2, TGF-beta2, TGF-beta3
- Murine: TGFbeta-R2/Fc Chimera



TECHNICAL SUPPORT

For troubleshooting information or assistance, please contact us at:

Tel: (519) 489-7195, (800) 836-8089 Fax: 519) 231-0140, (877) 221-3515 Email: product@biomatik.com



ELISA PLATE TEMPLATE

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