



ClinMax™ Human Soluble TROP-2 ELISA Kit, Pro

Catalog Number: CEA-B032

Assay Tests: 96 tests

CEA-B032-EN03

IMPORTANT: Please carefully read this user guide before performing your experiment.

Product information

This kit is specifically designed for the accurate quantitation of human Soluble Tumor-associated calcium signal

transducer 2 (TROP-2) from cell culture supernates, serum and plasma.

The principle of this assay employs a quantitative sandwich enzyme immunoassay approach. Initially, a microplate

is coated with a capture antibody. Then, samples and biotinylated capture antibody are added to the wells. After

the removal of any unbound materials through washing, streptavidin-HRP (SA-HRP) conjugate is added to the

wells. Streptavidin has a very high affinity for biotin, so it binds to the biotinylated capture antibody that is already

bound to the target antigen. After washing, a substrate specific to HRP is added to the wells. HRP catalyzes a

reaction that converts the substrate into a detectable signal, often a color change or luminescence, depending

on the substrate used. This enzymatic reaction amplifies the signal, allowing for higher sensitivity in detecting the

target analyte. The intensity of the signal is measured using a spectrophotometer.

NOTE:

1. This kit is for research use only and is not for use in diagnostic or therapeutic applications.

2. Please do not use the kit after the expiration date indicated on the kit label.

3. Do not mix or substitute reagents with those from other lots or sources.

Manufactured and distributed by

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Contents

The kit contains sufficient reagents for 96 wells.

Catalog	Contents	Amount
CEA032-C01	Pre-coated Anti-TROP-2 Antibody Microplate	1 plate
CEA032-C02	Human TROP-2 Standard	20 μg×2
CEA032-C03	Biotin-Anti-TROP-2 Antibody Con. Solution	400 μL
CEA032-C04	Biotin-Antibody Dilution Buffer	8 mL
CEA032-C05	Streptavidin-HRP Con. Solution	500 μL
CEA032-C06	Streptavidin-HRP Dilution Buffer	15 mL
CEA032-C07	20× Washing Buffer	50 mL
CEA032-C08	Sample Dilution Buffer	15 mL×2
CEA032-C09	Substrate Solution	12 mL
CEA032-C10	Stop Solution	6 mL

Storage

Keep the unopened kit stored at 2-8 °C. Avoid using the kit beyond its expiration date. For opened kit and reconstituted reagents, with the exception of the two contents listed in following table, others can be stored for up to 30 days at 2-8 °C.

Contents	Storage conditions
Pre-coated Anti-TROP-2 Antibody Microplate	Return unused wells to the foil pouch, reseal along entire edge. May be stored for up to 1 month at 2-8°C.
Human TROP-2 Standard	Aliquot and store for up to 1 month at -70°C in a freezer. Avoid repeated freeze-thaw cycles.

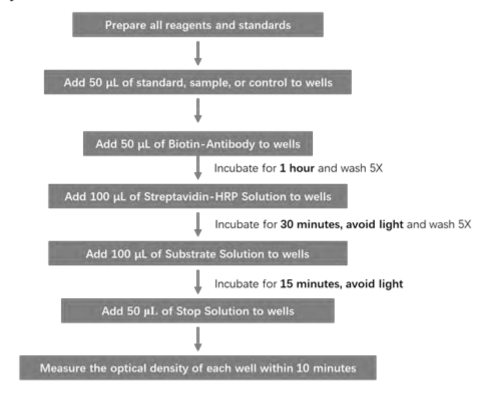
NOTE: Streptavidin-HRP Con. Solution and Substrate Solution should avoid light.

Required materials not supplied.

Instrument	Microplate reader capable of measuring absorbance at 450 nm		
Reagents	Deionized, ultrapure or distilled water		
50 mL and 500 mL graduated cylinders Consumables Pipettes and pipette tips			
			Tubes to prepare standard dilutions.

Workflow

Analyte: Soluble TROP-2



NOTE: Incubation temperature is 18 ℃-25 ℃

Prepare the working buffers and standard dilutions.

IMPORTANT: Bring all reagents to room temperature before use. If crystals have formed in buffer solution, place the buffer solution in an 37°C incubator until the crystals have completely dissolved and bring the solution back to room temperature before use.

Prepare the working buffers.

- 1. 1×Washing Buffer: Dilute 50 mL 20×Washing Buffer with deionized or distilled water to 1000 mL.
- 2. Biotin-Anti-TROP-2 Antibody Solution: Add 240 μL of Biotin-Anti-TROP-2 Antibody Con. Solution to 6 mL Biotin-Antibody Dilution Buffer, thoroughly mix. The solution was freshly prepared just before use.
- 3. TROP-2 Streptavidin-HRP Solution: Add 150 μ L of TROP-2 Streptavidin-HRP Con. Solution to 12 mL of Streptavidin-HRP Dilution Buffer, thoroughly mix. The solution was freshly prepared just before use.
- 4. Sample Dilution Buffer:

Prepare the reconstituted standard.

Add 1000 μ L ultrapure water to the provided lyophilized product (Catalog: CEA032-C02) , dissolve at room temperature for 15-30 minutes, and mix by gently pipetting. The concentration of reconstituted human TROP-2 Standard is 20 μ g /mL.

NOTE: Avoiding vigorous shaking or vortexing. The reconstituted solution should be stored at -70°C. The freeze-thaw cycle should not exceed 1 time, and the size of the aliquot should not be less than 10 µg.

Prepare the standard serial dilutions.

- 1. Label a tube "Cm". Add 50 μ L of the reconstituted human TROP-2 Standard and 950 μ L of Sample Dilution Buffer to tube Cm, gently mix well.
- 2. Label 5 tubes, one for each standard point: Std.-1, Std.-2, Std.-3, Std.-4, Std.-5.
- 3. Add 12 μ L of the liquid from **Cm** and 988 μ L of Sample Dilution Buffer to tube Std.-1, thoroughly mix (Std.-1 =12000 pg/mL).
- 4. Prepare 1:3 serial dilutions for the standard curve as follows: Add 300 μL of Sample Dilution Buffer to each tube (Std.-2, Std.-3, Std.-4, Std.-5).
- 5. Transfer 100 μ L of liquid from Std.-1 to the tube Std.-2, and thoroughly mix (Std.-2 = 3000 pg/mL).
- 6. Continue to transfer 100 μ L of liquid from previous dilution tube to the next dilution tube until add liquid to tube Std.-5.
- 7. Sample Dilution Buffer serves as zero standard (blank).

PROCEDURE OF ASSAY

- 1. Add 50 μL of TROP-2 Standard, sample, or control to wells.
- 2. Add 50 μL Biotin-Anti-TROP-2 Antibody Solution to each well, Seal the plate with microplate sealing film. Incubate at room temperature (18-25 °C) for **1 hours**.
- 3. Aspirate each well and add 300 μ L of 1×Washing Buffer to each well, gently tap the plate for **1 minute**. Remove any remaining Washing Buffer by aspirating or decanting. Invert the plate and blot it against clean paper towels. Repeat the wash process four times for a total of five washes.
- 4. Add 100 μ L of TROP-2 Streptavidin-HRP Solution to each well. Seal the plate with microplate sealing film. Incubate at room temperature (18-25 °C) for 30 minutes, avoid light.
- 5. Repeat step 3.
- 6. Add 100 μ L of Substrate Solution to each well. Seal the plate with microplate sealing film and incubate at room temperature (18-25 °C) for **15 minutes, avoid light**.
- 7. Add 50 μL of Stop Solution to each well. Tap the plate gently to ensure thorough mixing.

 *Note: the color in the wells should change from blue to yellow.
- 8. Read the absorbance at 450nm and 630nm using Microplate reader within 10minutes.

 *Note: To reduce the background noise, subtract the readings at 630nm from the readings at 450nm.

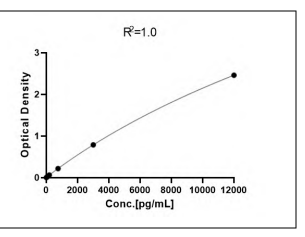
CALCULATION OF RESULTS

- 1. Compute the average of the duplicated readings for every standard, control, and sample. Then, subtract the average optical density (O.D.) of the zero standard(blank).
- 2. Establish a standard curve by processing the data using computer software capable of executing a four-parameter logistic (4-PL) curve fitting.
- 3. Normal range of Standard curve: $R^2 \ge 0.9900$.
- 4. If the OD value of the sample to be tested is higher than the highest standard, the sample shall be diluted with dilution buffer and assay repeated.

Typical data

Note: For each experiment, a standard curve needs to be set for each microplate, and the specific OD value may vary depending on different laboratories, testers, or equipment. The following example data is for reference only. The sample concentration was calculated based on the results of the standard curve.

TROP-2 Standard	OD
(pg/mL)	OD _{450nm-630nm}
12000	2.464
3000	0.791
750	0.222
187.5	0.065
46.875	0.022
Blank	0.007



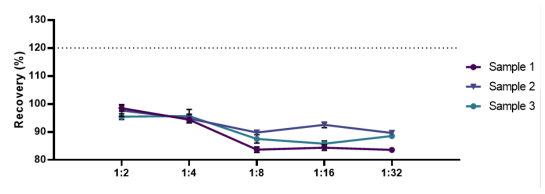
PERFORMANCE CHARACTERISTICS

1. Sensitivity

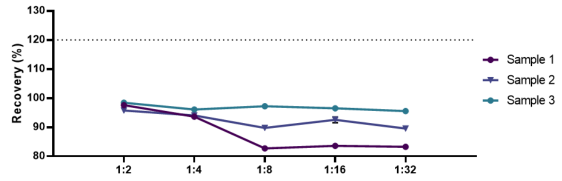
The minimum detectable concentration (MDC) of TROP-2 is typically less than 46.875 pg/mL. The MDC was determined by adding two standard deviations to the mean optical density value of twenty zero standard replicates and calculating the corresponding concentration.

2. Linearity

Three samples (Serum) spiked with high concentrations of TROP-2 were serially diluted with dilution buffer to produce samples with values within the dynamic range of the assay and then assayed. The average recovery of TROP-2 for serum samples is 90.8%.



Three samples (EDTA plasma) spiked with high concentrations of TROP-2 were serially diluted with dilution buffer to produce samples with values within the dynamic range of the assay and then assayed. The average recovery of TROP-2 for serum samples is 92.4%.



3. Intra-Assay Precision

Ten replicates of each of 4 samples containing different TROP-2 concentrations were tested in one assay. Acceptable criteria: CV < 10%.

Sample Concentration (pg/mL)	Mean (pg /mL)	SD	Numbers	CV
12000	11741.1	620.4	10	5.3%
9000	8954.6	413.9	10	4.6%
6000	5562.7	297.0	10	5.3%
187.5	173.1	10.4	10	6.0%

4. Inter-Assay Precision

Five samples containing different concentrations of TROP-2 were tested in independent assays. Acceptable criteria: CV<15%.

Sample Concentration (pg/mL)	Mean (pg/mL))	SD	Numbers	CV
12000	11692.9	610.8	9	5.2%
9000	8735.1	719.3	9	8.2%
6000	5487.1	267.0	9	4.9%
750	680.8	50.5	9	7.4%
187.5	169.6	14.0	9	8.3%

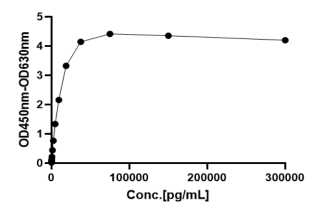
5. Recovery

Recombinant TROP-2 was spiked into 3 human serum samples, and then analyzed. The average recovery of TROP-2 for serum samples is 92.4%.

Sample ID	Conc Measured (pg/mL)	Conc Added (pg/mL)	Conc Recovered (pg/mL)	Recovery	
	8844.5	9000	8777.8	97.6%	
1	5681.8	6000	5615.1	93.7%	
	2788.1	3000	2721.4	90.9%	
	8175.6	9000	8108.9	90.2%	
2	5459.2	6000	5392.5	90.0%	
	2797.1	3000	2730.4	91.2%	
	8676.6	9000	8592.9	95.6%	
3	5657.0	6000	5573.3	93.0%	
	2767.7	3000	2684.1	89.7%	

6. Hook Effect

Not be affected by the concentration of TROP-2 up to 300,000 pg/mL.



Interference Effect

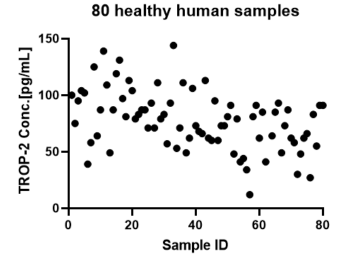
To evaluate the hemolysis matrix effect and high-dose triglyceride matrix effect of assay, serum samples spiked with high concentrations of hemoglobin (2%), or triglyceride (3 mg/mL) were tested. Results shown that all spiked analytes had recoveries between 86% and 112%, no hemolysis matrix effect and high-dose triglyceride matrix effect was observed in assay.

Spiked Material	ID	Conc-1(pg/mL)	Conc-2(pg/mL)	Mean(pg/mL)	Recovery
	Sample 1	183.31	179.02	181.17	00%
	Spiked Sample1	179.33	178.32	178.83	99%
0% 11	Sample 2	143.6	134.5	139.05	92%
	Spiked Sample 2	125.01	131.06	128.04	92%
2% Hemoglobin (v/v)	Sample 3	100.49	95.04	97.77	109%
	Spiked Sample 3	105.31	108.06	106.69	109%
	Sample 4	49.36	46.93	48.15	06%
	Spiked Sample 4	42.08	40.27	41.18	86%

Spiked material	ID	Conc-1 (pg/mL)	Conc-2 (pg/mL)	Mean (pg/mL)	Recovery
	Sample 1	183.31	179.02	181.17	112%
	Spiked Sample1	207.12	199.36	203.244	11290
	Sample 2	143.6	134.5	139.05	94%
Triglyceride (3 mg/mL)	Spiked Sample 2	135.01	127.06	131.04	94%
	Sample 3	100.49	95.34	97.92	107%
	Spiked Sample 3	106.31	104.16	105.24	10790
	Sample 4	49.36	46.93	48.15	110%
	Spiked Sample 4	52.08	54.27	53.18	110%

7. Sample Values

80 healthy serum samples were evaluated for the concentrations of human TROP-2 in assay.



8. Specificity

No cross-reactivity was observed when this kit was used to analyze the following recombinant cytokines at up to 1 μ g/mL.

Human	IL-1β, IL-2, IL-4, IL-5, IL-6, IL-7, IL-8, IL-12 p70, IL-10, IL-13, IL-15, IL-17,
Tiulilali	GM-CSF, TNF-α

TROUBLESHOOTING GUIDE

Problem	Cause	Solution
Poor standard curve	* Inaccurate pipetting	* Check pipettes
Large CV	* Inaccurate pipetting * Air bubbles in wells	* Check pipettes * Remove bubbles in wells
High background	* Plate is insufficiently washed* Contaminated wash buffer	* Review the manual for proper wash. * Make fresh wash buffer
Very low readings across the plate	* Incorrect wavelengths * Insufficient development time	* Check filters/reader * Increase development time
Samples are reading too high, but standard curve looks fine	* Samples contain cytokine levels above assay range	* Dilute samples and run again
Drift	* Interrupted assay set-up * Reagents not at room temperature	* Assay set-up should be continuous - have all standards and samples prepared appropriately before commencement of the assay * Ensure that all reagents are at room temperature before pipetting into the wells unless otherwise instructed in the antibody inserts