

Human tPA ELISA Kit (Rapid One-Step)

Enzyme Immunoassay for the quantification of Human tPA in serum, plasma (heparin, EDTA, citrate), urine and cell culture supernatants.

Catalog number: ARG83216

Package: 96 wells

For research use only. Not for use in diagnostic procedures.

TABLE OF CONTENTS

SECTION	Page
INTRODUCTION	3
PRINCIPLE OF THE ASSAY	3
MATERIALS PROVIDED & STORAGE INFORMATION	4
MATERIALS REQUIRED BUT NOT PROVIDED	4
TECHNICAL HINTS AND PRECAUTIONS	4
SAMPLE COLLECTION & STORAGE INFORMATION	6
REAGENT PREPARATION	7
ASSAY PROCEDURE	9
CALCULATION OF RESULTS	10
EXAMPLE OF TYPICAL STANDARD CURVE	11
OUALITY ASSURANCE	11

MANUFACTURED BY:

Arigo Biolaboratories Corporation

Address: No. 22, Ln. 227, Gongyuan Rd., Hsinchu City 300, Taiwan

Phone: +886 (3) 562 1738

Fax: +886 (3) 561 3008

Email: info@arigobio.com

INTRODUCTION

This gene encodes tissue-type plasminogen activator, a secreted serine protease which converts the proenzyme plasminogen to plasmin, a fibrinolytic enzyme. Tissue-type plasminogen activator is synthesized as a single chain which is cleaved by plasmin to a two chain disulfide linked protein. This enzyme plays a role in cell migration and tissue remodeling. Increased enzymatic activity causes hyperfibrinolysis, which manifests as excessive bleeding; decreased activity leads to hypofibrinolysis which can result in thrombosis or embolism. Alternative splicing of this gene results in multiple transcript variants encoding different isoforms. [provided by RefSeq, Jul 2008]

PRINCIPLE OF THE ASSAY

This assay employs the quantitative sandwich enzyme immunoassay technique. An antibody specific for tPA has been pre-coated onto a microtiter plate. Standards or samples are pipetted into the wells and any tPA present is bound by the immobilized antibody. Then Antibody conjugated specific for tPA is added to each well and incubate. Following a washing to remove unbound substances, a substrate solution (TMB) is added to the wells and color develops in proportion to the amount of tPA bound in the initial step. The color development is stopped by the addition of acid and the intensity of the color is measured at a wavelength of 450nm ±2nm. The concentration of tPA in the sample is then determined by comparing the O.D of samples to the standard curve.

MATERIALS PROVIDED & STORAGE INFORMATION

Store the unopened kit at 2-8°C. Use the kit before expiration date.

NO	Component	Quantity	Storage information
C1	Antibody-coated microplate	8 X 12 strips	4°C. Unused strips should be sealed tightly in the air-tight pouch.
C2	Standard (Lyophilized)	2 X 10 ng/vails	4°C
C3	Diluent buffer	30 ml (ready to use)	4°C
C4	Antibody conjugated	1 vial (ready to use)	4°C
C5	25X Wash buffer	20 ml	4°C
C6	TMB substrate	10 ml (ready to use)	4°C (Protect from light)
C7	STOP solution	10 ml (ready to use)	4°C
C8	Plate sealer	4 strips	Room temperature

MATERIALS REQUIRED BUT NOT PROVIDED

- Microplate reader capable of measuring absorbance at 450nm
- Pipettes and pipette tips
- Deionized or distilled water
- 37°C oven or incubator
- Automated microplate washer (optional)

TECHNICAL HINTS AND PRECAUTIONS

- Wear protective gloves, clothing, eye, and face protection especially while handling blood or body fluid samples.
- Store the kit at 2-8°C at all times.
- If crystals are observed in the 25X Wash buffer, warm to RT (not more than 50°C) until the crystals are completely dissolved.

Human tPA ELISA kit (Rapid One-Step) ARG83216

- Ensure complete reconstitution and dilution of reagents prior to use.
- All materials should be equilibrated to room temperature (RT, 22-25°C)
 20 min before use.
- All reagents should be mixed by gentle inversion or swirling prior to use.
 Do not induce foaming.
- Before using the kit, spin tubes and bring down all components to the bottom of tubes.
- Mix the contents of the microplate wells thoroughly by microplate shaker for 1 min or gently tap the plate to ensure good test results. Please mix carefully to avoid well-to-well contamination. Do not reuse microwells.
- The TMB Color developing agent should be colorless and transparent before using.
- Use reservoirs only for single reagents. This especially applies to the substrate reservoirs. Using a reservoir for dispensing a substrate solution that had previously been used for the conjugate solution may turn solution colored. Do not pour reagents back into vials as reagent contamination may occur.
- Do not let wells dry during assay; add reagents immediately after completing the rinsing steps.
- Avoid using reagents from different batches.
- It is highly recommended that the standards, samples and controls be assayed in duplicates.
- Change pipette tips between the addition of different reagent or samples.

SAMPLE COLLECTION & STORAGE INFORMATION

The sample collection and storage conditions listed below are intended as general guidelines. Sample stability has not been evaluated.

<u>Cell Culture Supernatants</u> - Remove particulates by centrifugation for 10 min at 1500 x g at 4°C and aliquot & store samples at -20°C up to 1 month or -80°C up to 6 months. Avoid repeated freeze-thaw cycles.

<u>Serum</u>- Use a serum separator tube (SST) and allow samples to clot for 30 minutes before centrifugation for 15 minutes at 1000 x g. Collect serum and assay immediately or aliquot & store samples at -20°C up to 1 month or -80°C up to 6 months. Avoid repeated freeze-thaw cycles.

<u>Plasma</u>- Collect plasma using EDTA or heparin as an anticoagulant. Centrifuge for 15 minutes at $1000 \times g$. within 30 minutes of collection. Collect the supernatants and assay immediately or aliquot and store samples at -20° C up to 1 month or -80° C up to 6 months. Avoid repeated freeze-thaw cycles.

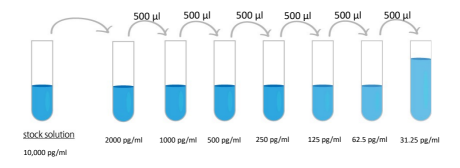
<u>Urine</u> - Collect the urine by micturating directly into a sterile container. Remove impurities by centrifugation at 10,000 x g for 1 min. Collect the supernatants and assay immediately or aliquot and store samples at-20°C up to 1 month or -80°C up to 6 months.

Note:

- a) Do not use haemolytic, icteric or lipaemic specimens.
- b) Samples containing sodium azide should not be used in the assay.

REAGENT PREPARATION

- 1X Wash buffer: Dilute 25X Wash buffer (#C5) into distilled water to yield 1X Wash buffer. (E.g. 50 ml of 25X Wash buffer + 1200 ml of distilled water)
 The diluted Wash buffer is stable for 4 weeks at 2°C to 8°C.
- Sample: If the initial assay found samples contain tPA higher than the
 highest standard, the samples can be diluted with diluent buffer (#C3) and
 then re-assay the samples. For the calculation of the concentrations this
 dilution factor has to be taken into account.
 - (It is recommended to do pre-test to determine the suitable dilution factor).
- Standards: Reconstitute the standard (#C2) with 1 ml diluent buffer (#C3) to yield a stock concentration of 10,000 pg/ml. Keep the buffer in the vail for at least 15 min at RT to make sure the standard is dissolved completely before making serial dilutions. The diluent buffer serves as zero standard (0 pg/ml), and the rest of the standard serial dilution can be diluted as according to the suggested concentration below: 2000 pg/ml, 1000 pg/ml, 500 pg/ml, 250 pg/ml, 125 pg/ml, 62.5 pg/ml, 31.25 pg/ml. DO NOT reuse the reconstituted standard.



Dilute tPA standard as according to the table below:

Standard	tPA Conc.	μl of Standard diluent	μl of standard
			200
S7	2000 pg/ml	800	(10,000 pg/ml
			Stock)
S6	1000 pg/ml	500	500 (S7)
S5	500 pg/ml	500	500 (S6)
S4	250 pg/ml	500	500 (S5)
S3	125 pg/ml	500	500 (S4)
S2	62.5 pg/ml	500	500 (S3)
S1	31.25 pg/ml	500	500 (S2)
S0	0	500	0

ASSAY PROCEDURE

All materials should be equilibrated to room temperature (RT) 20 min before use. Standards, samples and controls should be assayed in duplicates.

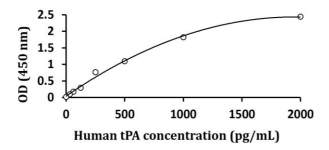
- 1. Remove excess microplate strips from the plate frame, return them to the foil pouch containing the desiccant pack, and reseal it.
- 2. Add 50 µl of standards, samples and zero controls (diluent buffer) into wells.
- 3. Add **50 μl** <u>1X Antibody conjugated</u> into each well, gently tap the plate to mix well. Cover wells and incubate for **1 hour at Room Temperature**.
- 4. Aspirate each well and wash, repeating the process three times for a total four washes. Wash by filling each well with 1× Wash Buffer (300 μl) using a squirt bottle, manifold dispenser, or autowasher, keep the Wash Buffer in the wells for 30 sec before remove. Complete removal of liquid at each is essential to good performance. After the last wash, remove any remaining Wash Buffer by aspirating, decanting or blotting against clean paper towels.
- 5. Add **90 \muI** of <u>TMB Reagent</u> (#C6) to each well, gently tap the plate to mix well. Incubate for **15-25 minutes at Room Temperature** in dark.
- 6. Add **100** μ l of Stop Solution (#C7) to each well, gently tap the plate to mix well. The color of the solution should change from blue to yellow.
- Read the OD with a microplate reader at 450 nm immediately. It is recommended read the absorbance within 3 min after adding STOP solution.

CALCULATION OF RESULTS

- 1. Calculate the average absorbance values for each set of standards, controls and patient samples.
- 2. Using linear graph paper, construct a standard curve by plotting the mean absorbance obtained from each standard against its concentration with absorbance value on the vertical (Y) axis and concentration on the horizontal (X) axis.
- 3. Using the mean absorbance value for each sample determine the corresponding concentration from the standard curve.
- 4. Automated method: The results in the IFU have been calculated automatically using a 4 PL (4 Parameter Logistics) curve fit. 4 Parameter Logistics is the preferred method. Other data reduction functions may give slightly different results.
- 5. arigo provides GainData®, an in-house development ELISA data calculator, for ELISA data result analysis. Please refer our GainData® website for details. (https://www.arigobio.com/elisa-analysis)
- 6. If the samples have been diluted, the concentration read from the standard curve must be further converted by the appropriate dilution factor according to the sample preparation procedure as described above.

EXAMPLE OF TYPICAL STANDARD CURVE

The following data is for demonstration only and cannot be used in place of data generations at the time of assay.



QUALITY ASSURANCE

Sensitivity

The minimum detectable dose (MDD) of Human tPA ranged from 31.2 pg/ml-2,000 pg/ml. The mean MDD was 15 pg/ml.

Specificity

This assay recognizes natural and recombinant Human tPA.

Intra-assay and Inter-assay precision

The CV values of both intra and inter precision fall below 10%.