







TECHNOZYM[®] ADAMTS-13 Antigen

For research use only



REF	5450601	TECHNOZYM [®] ADAMTS-13 Antigen	
REF	5450661	TECHNOZYM [®] ADAMTS-13 Antigen Calibrator Set	5 x 0.5 mL
REF	5450663	TECHNOZYM [®] ADAMTS-13 Antigen Control Set	2 x 0.5 mL

Symbols key			
	Manufacturer		Expiry date
	Storage temperature		Consult instructions for use
AQUA	Distilled water		Determinations
BUF	Reaction buffer	LOT	Lot
CAL	Calibrator	MTP	Microtiter plate
CONJ	Conjugate	REF	Catalogue number
CONT	Control	RTU	Ready to use
DIL	Dilute or dissolve in	STOP	Stop solution
INC	Incubation buffer	SUB	Substrate
RUO	For research use only	WASH	Washing solution concentrate



PRODUCT DESCRIPTION

INTENDED USE

The TECHNOZYM® ADAMTS-13 Antigen ELISA is a chromogenic test for the determination of ADAMTS-13 antigen concentration in human plasma. ADAMTS-13 is the enzyme that cleaves vWF under laminar flow conditions. A functional defect of this enzyme leads to the presence of higher molecular weight forms of vWF and thus to increased platelet aggregation, mainly in the microvasculature. This is believed to be the major cause for thrombotic thrombocytopenic Purpura (TTP).

COMPOSITION

- ELISA test strips (12), with 8 wells each, coated with a monoclonal anti ADAMTS-13 antibody, directed against the CUB domain; the drying agent is supplied in an aluminium bag.
- Washing buffer concentrate (PBS; pH 7.3); containing detergent; 0.01% merthiolate; 1 vial, 80 mL.
- Incubation buffer (= sample dilution buffer) (PBS; pH 7.3); contains stabiliser protein; 0.05% proclin; and dye, 1 vial, 90 mL, ready for use.
- Calibrators (Standards) numbered from 1 to 5; lyophilised; 1 vial each; 0.5 mL. **Concentrations are lot-specific; consult the label on the vial.**
- High and Low control plasma; lyophilised; 1 vial each; 0.5 mL. **Concentrations are lot-specific; consult the label on the vial.**
- Conjugate: anti-ADAMTS-13 POX; dyed blue; 1 vial, 0.3 mL.
- Chromogenic substrate TMB (tetramethylbenzidine); 1 vial; 12 mL; ready to use.
- Stopping solution sulphuric acid 0.45 mol/L; 1 vial; 12 mL; ready to use.
- Adhesive film: for ELISA test strips; 2 pieces.

MATERIAL REQUIRED (not supplied with the kit)

- Distilled water
- Measuring cylinder (1000 mL)
- Precision pipettes (50, 100 and 1000 µL)
- Variable pipette (100 and 1000 µL)
- Multichannel and/or dispensing pipettes (100 and 200 µL)
- ELISA washer or multichannel pipette
- ELISA reader with 450 nm filter, with a 620 nm reference filter if available.

WARNING AND PRECAUTIONS

- For research use only
- All human blood or plasma products as well as samples must be considered as potentially infectious. They have to be handled with appropriate care and in strict observance of safety regulations. The rules pertaining to disposal are the same as applied to disposing hospital waste.
- Calibrators and control plasmas are made from human blood and any individual plasma involved in the procedure is HBsAg, HIV 1/2 Ab and HCV-Ab-negative (see labels on the vials). However, all human blood products should be handled as potentially infectious material.
- Stopping solution may irritate the skin. Should acid get into your eyes, wash out immediately with water and consult a doctor.
- The reagents sometimes contain preserving agents (merthiolate). Beware of swallowing! Avoid contact with skin or mucous membranes!

STABILITY AND STORAGE

The expiry date printed on the labels applies to storage of the unopened vial at +2...8°C.

Stability after reconstitution/opening:

Material/Reagent	State	Storage	Stability
Calibrators, control plasmas	after reconstitution	-20 °C	6 months
ELISA test strip	after opening	+2...8 °C with adhesive film in plastic bag with drying agent	expiry date
Washing buffer conc.	after opening	+2...8°C	6 months
Washing buffer	1+11.5 dilution of concentrate	+2...8°C	3 weeks
Incubation buffer (= sample dilution buffer)	after opening	+2...8°C	2 months
Conjugate	after opening	+2...8°C	6 months
	working solution	room temperature (+18...25°C)	60 minutes
Chromogenic substrate TMB	after opening	+2...8°C	expiry date

TEST PROCEDURE

PREPARATION OF THE SAMPLES

Sample material: Human citrated plasma. Samples may be stored for three hours at room temperature. At -20°C they can be stored for several months. Samples may not be frozen and thawed several times.

PREPARATION OF REAGENT

- Before starting the test, all the required components are to be brought to room temperature.
- Preparing the washing buffer: Dilute 1 part by volume washing buffer concentrate with 11.5 parts by volume distilled water (1+11.5). Mix well! (Diluted washing buffer concentrate = washing buffer). There may be crystalline precipitations which will dissolve at +37°C within 10 minutes.
- Reconstituting calibrators and control plasmas: Calibrators and control plasmas are reconstituted with 500 µL distilled water and mixed for 10 seconds after a reconstitution time of 15 minutes (vortex mixer).
Reconstituted components are clear to slightly turbid.
No dilution is necessary for calibrators and controls!
- Samples are used undiluted
- Preparing the conjugate working solution (1+50): Dilute 1 part by volume conjugate with 50 parts by volume incubation buffer.

PERFORMANCE OF THE TEST

SAMPLE INCUBATION (reference 1,2)	Pipette calibrators, control plasmas, samples into test wells; cover test strips with film	50 µL
	Incubate at room temperature	120 minutes
WASHING (reference 1,3,4)	Washing buffer	4 x 250 µL
	Incubate at room temperature	60 minutes
CONJUGATE REACTION (reference 1,2)	Pipette conjugate working solution into wells, cover test strip with film	50 µL
	Incubate at room temperature	60 minutes
WASHING (reference 1,3,4)	Washing buffer	4 x 250 µL
	Incubate at room temperature	15 minutes
STOPPING (reference 1,2)	Pipette stopping solution into wells	50 µL
	ELISA-Reader, 450 nm	shake 10 sec., measure within 10 min.

References

- Reagents of different lots must not be combined
- Precision and performance, among others, essentially depend on the following factors:
 - Thorough mixing of all substances used for dilution, 10 sec. with Vortex Mixer
 - Test calibrators, controls and samples in duplicates
 - Incubate at indicated temperature (RT: room temperature, +18...25°C)
 - Strict observance of the order of pipetting and of the time element as indicated
 - The time for sample incubation, conjugate and substrate reaction as indicated starts after pipetting the last sample. Incubation times shall not vary by more than ± 5%.
 - During sample incubation and conjugate reaction, the time for pipetting calibrators/control plasmas/samples and/or conjugate solutions must not exceed 60 seconds per ELISA test strip (8 wells).
 - During substrate reaction and at stopping, the time needed for pipetting the substrate and/or the stopping solution must not exceed 10 seconds per ELISA test strip. Short pipetting times may be secured by using multichannel- and dispensing pipettes.
- Label/number strips with a water resistant pen in case the strips accidentally fall out of the frame during testing.
- After the last washing, wells must be aspirated thoroughly, turned upside down and positioned on a blotting paper, by gentle tapping, the last remnants must be removed.
- By measuring the difference in wave lengths at 450 and 620 nm the precision of the test is increased.
- A calibration curve has to be created for every assay

LIMITATION OF THE TEST

It can not be excluded that certain forms of ADAMTS-13 (with mutations in the CUB domains) are not equivalently measured due to reduced binding to the capture antibody on the plate. Thrombin is reported to degrade ADAMTS-13. Therefore serum samples should be avoided.

ANALYSES RESULTS

CALCULATION OF THE RESULTS

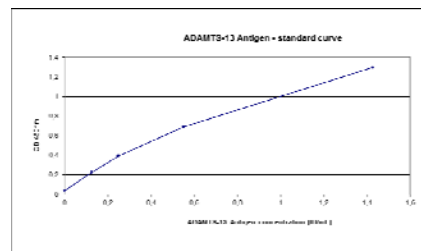
Setting up a reference curve: X axis: concentration ADAMTS-13 antigen [IU/mL]
Y axis: Extinction at 450 nm

Graph plot is linear-linear with a best fit.

Assessment of reference curve

The validity of the test may be checked on the basis of the calculated control values.

Example of standard curve



Measuring concentration of samples

- Read off the concentration from the reference curve.
- If there are samples with extinction coefficients higher than that of the highest point on the curve, they have to be pre-diluted with incubation buffer (1+1, or 1+3). The measured concentration then has to be multiplied with the dilution factor 2 or 4, respectively.

INTERPRETATION OF RESULTS / REFERENCE RANGE

Normal range for ADAMTS-13 Antigen concentration: 0.41 – 1.41 IU/mL (n=188)
Normal range can vary depending on local population. It is recommended that individual laboratories establish their own normal. When interpreting the serological results the history of the sample has to be taken into account.

STANDARDISATION

Standards and controls are produced from a pool of normal donor plasma and calibrated against the WHO International Standard for ADAMTS-13.

PERFORMANCE CHARACTERISTICS

Performance data are given below. Results obtained in individual laboratories may differ.

PRECISION

Reproducibility was determined with different samples (in series and day to day). The following results were obtained.

Sample	Intra assay variation		Inter assay variation	
	Sample 1	Sample 2	Sample 3	Sample 4
N	10	10	3	3
Mean (IU/mL)	0.55	0.11	0.49	0.10
SD (IU/mL)	0.04	0.01	0.05	0.01
CV (%)	6.36	5.71	9.97	6.62

ASSAY RANGE

0 IU/mL – 1.0 IU/mL

(or up to the actual value of calibrator 1)

DETECTION LIMIT

0.012 IU/mL

CORRELATION

Correlation with antigen in TECHNOZYM® ADAMTS-13 fluorogenic method is $r^2 = 0.9$ for normal samples and $r^2 = 0.91$ for TTP samples.

LITERATURE

Please contact Technoclone or your local distributor.

For 8 test wells:
Mix 10 µL conjugate with 500 µL incubation (= sample dilution) buffer