# Human Thioredoxin-1 ELISA<sup>PRO</sup> kit

Kit for 10 plates

Product code: 3580-1HP-10

# TABLE OF CONTENTS

# Kit description4Reagents provided4Storage and expiry date5Materials required but not supplied5Safety issues5Procedural notes5Preparations6Assay procedure8Performance of the assay9Limitations of the assay10

#### PAGE

#### **KIT DESCRIPTION**

This ELISA<sup>PRO</sup> kit is a complete kit for the quantification of human Thioredoxin-1 (Trx1) in biological fluids such as serum, plasma and cell culture supernatants. The assay utilizes ELISA strip plates pre-coated with a capture monoclonal antibody (mAb), to which samples are added. Captured Trx1 is detected by adding a biotinylated mAb followed by streptavidin-horseradish peroxidase (SA-HRP). Addition of the enzyme substrate TMB will result in a colored substrate product with an intensity that is directly proportional to the concentration of Trx1 in the sample. The concentration of Trx1 in the sample is determined by comparison to a serial dilution of recombinant Trx1 standard analyzed in parallel.

Component	Quantity	Description/comments
Precoated 96-well strip plate (12 strips x 8 wells)	10 plates	Plate coated with anti-Human Thioredoxin-1. Supplied in foil bag with desiccant.
ELISA standard	1 vial	Lyophilized recombinant Thioredoxin-1. See "Preparations" for reconstitution and dilution.
Detection antibody	1 vial (250 μl)	Biotinylated anti-human Thioredoxin-1 mAb MT13X3 (5 μg/ml). Dilute before use.
SA-HRP	1 vial (130 μl)	Streptavidin-horseradish peroxidase conjugate. Dilute before use.
Standard reconstitution buffer A5 (ready-to-use)	1 vial (1ml)	For reconstitution of lyophilized Thioredoxin-1 standard.
Wash buffer concentrate (20x)	5 bottles (5x120 ml)	For all wash steps. Dilute before use.
ELISA diluent/ Assay buffer (ready-to-use)	3 bottles (3x120 ml)	Protein-containing buffer for: 1) dilution of all samples; 2) serial dilution of standard; 3) dilution of detection antibody.
SA-HRP diluent (ready-to-use)	1 bottle (120 ml)	For dilution of SA-HRP.
TMB substrate (ready-to-use)	1 bottle (120 ml)	3,3',5,5' tetramethylbenzidine (TMB) enzyme substrate solution containing hydrogen peroxide.
Stop solution (ready-to-use)	1 bottle (120 ml)	0.18 M H <sub>2</sub> SO <sub>4</sub>
Adhesive plate covers	30	To cover plates during incubations.

#### **REAGENTS PROVIDED**

To ensure total recovery of stated quantity, bottles and vials are filled with larger volume than indicated.

#### STORAGE AND EXPIRY DATE

Shipped at ambient temperature. On arrival all components of the kit, with the exception of the lyophilized standard, should be stored at 2-8°C. **Please note that the lyophilized stan-dard should be kept at -20°C.** After reconstitution of the lyophilized standard, it should be aliquoted and kept at -20°C. The expiry date for the unopened kit is indicated on the box. We recommend to use opened kit components within one month.

# MATERIALS REQUIRED BUT NOT SUPPLIED

- Microplate reader capable of reading at 450 nm; preferably able of subtracting a reference wavelength between 570-650 nm.

- ELISA plate washer; automatic (adaptable for ELISA strip plates) or manual (e.g. multipipette or squirt bottle).

- Precision pipettes and tips.
- Beakers, flasks and graduated cylinders necessary for reagent preparations.
- Tubes for standard and sample dilutions.
- Distilled or deionized water.

# SAFETY ISSUES

- The stop solution, 0.18 M  $\rm H_2SO_4$  (< 1%), is irritating to eyes and skin and should be handled with care.

- The standard should be handled with care due to unknown effects of exposure.
- Buffers and reagents in solution contain 0.002% of the preservative Kathon CG,
- a potential contact allergen which may cause sensitization by skin contact.
- Human and animal samples should be treated as potentially hazardous biological material.
- All material and samples should be disposed of in accordance with local regulations.

# **PROCEDURAL NOTES**

- Do not combine components from different kit lots or components from other suppliers.

- Dilutions of standard and samples can be prepared in plastic or glass tubes.
- Total sample or standard volume added per well should not exceed 100 µl.

- The use of duplicates for each dilution of the standard, samples and blank is recommended.

- Prior to adding new reagents to the wells, ensure that there is no residual wash buffer remaining in the wells. The wells can be emptied by turning the plate upside down followed by gently tapping the plate against clean tissue paper. Please be careful to avoid that the strips fall out of the frame e.g. grip the plate by the middle.

- Serum/plasma components present in the sample may have an impact on the performance of the assay. For this reason, all samples should be diluted at least 2x in "Assay buffer".

- Sodium azide and other nucleophilic substances (often used as preservative in various buffers) interfere with the activity of horseradish peroxidase. Thus, avoid the use of other wash buffers or solutions that may contain such interfering substances.

#### PREPARATIONS

#### Plates

Allow the plates to adjust to room temperature (18-25°C) before opening the bags.

Plan the experiment to include a standard curve and an assay background control (9 x 2 wells), blank (2 wells) and sample wells. To the blank wells, only "TMB substrate" and "Stop solution" should be added. Before the analysis of absorbance values, the mean value of the blank wells should be subtracted from the standard, the assay background control and the sample values.

Assemble the required numbers of strips in the plate frame. The strips used for the experiment can be marked e.g. with a marker pen. Store the remaining strips in the foil bag containing the desiccant at 4-8°C.

#### Wash buffer

Prepare the required volume of wash buffer by diluting "Wash buffer concentrate" 20x with distilled or deionized water. For 1 plate, prepare 1000 ml wash buffer by adding 50 ml "Wash buffer concentrate" to 950 ml distilled or deionized water.

#### ELISA standard

Reconstitute the Trx1 standard to a standard stock solution of 0.5  $\mu$ g/ml. Add 1 ml of "Standard reconstitution buffer A5" and allow the standard to dissolve for 5 min. Mix thoroughly and aliquot. Store at -20°C. Avoid repeated freeze-thaw cycles of the standard aliquotes.

#### Preparation of standard curve

The standard curve can be made from a standard stock solution just reconstituted or from thawed aliquotes. Prepare a serial dilution of the standard no more than 30 min prior to the experiment. Duplicate wells for the standard are recommended. Dilute the standard stock solution to create a standard curve ranging from 31.25-4000 pg/ml according to the scheme below. For the assay background control (0 pg/ml), use only "Assay buffer".

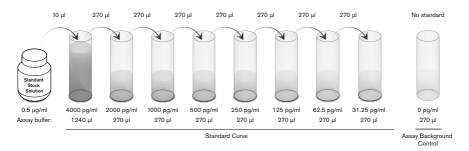


Figure 1. Recommended serial dilution of standard. The volumes indicated are sufficient for duplicates. The last vial should be 0 pg/ml of standard i.e. the analyte should be omitted.

#### Samples

Dilute all samples in "Assay buffer" prior to use. We recommend the use of duplicate wells for all samples. It is recommended that visible precipitate in the sample should be removed. For accurate measurement of Trx1 blood levels, the use of plasma is recommended since serum contains high levels of Trx1 released from platelets. To minimize the platelet content in plasma, an additional centrifugation at 10,000 x g for 10 minutes in connection with the plasma preparation is recommended. The monoclonal antibodies cross-react with bovine Trx1, thus cell culture supernatants must be free of bovine serum for determination of human Trx1. **Important!** All samples should be diluted at least 2x in "Assay buffer". This applies also for samples where low analyte levels are anticipated. Samples containing high levels of Trx1 exceeding the standard range of the assay will require further dilution. Sample dilutions can be made either in tubes or directly in the plate. For tube dilutions, equal volumes of "Assay buffer" and sample should be mixed, e.g. 120  $\mu$ l "Assay buffer" (50  $\mu$ l per well) should be added prior to addition of the sample (50  $\mu$ l per well).

#### **Detection antibody**

Dilute the "Detection antibody" 500x in "Assay buffer" prior to use. For 1 plate, dilute 24  $\mu$ l "Detection antibody" in 12 ml "Assay buffer" which will give a final "Detection antibody" concentration of 0.01  $\mu$ g/ml.

#### SA-HRP

Dilute the "SA-HRP" 1000x in "SA-HRP diluent" prior to use. For 1 plate, dilute 12  $\mu l$  "SA-HRP" in 12 ml "SA-HRP diluent".

## ASSAY PROCEDURE

Throughout the assay all reagents and samples, except the "TMB substrate", should be adjusted to room temperature (18-25°C) prior to use. The "TMB substrate" should preferably be used cold.

1. Assemble the numbers of strips required for the standard curve, assay background control, the blanks and samples in the plate frame. The use of duplicates is recommended.

2. Wash the strips with  $5x300 \,\mu$ /well of diluted wash buffer. The wash buffer should be thoroughly removed in immediate relation to the next step.

3. Add 100  $\mu$ /well of each concentration of the diluted Trx1 standard and assay background control. For the samples, add either 100  $\mu$ /well of pre-diluted sample or add 50  $\mu$ l "Assay buffer" + 50  $\mu$ l sample per well. Mix by tapping the plate.

**Important!** Both the serial dilution of the Trx1 standard and sample dilutions should be made in "Assay buffer". Leave the blank wells empty. Cover the plate with adhesive plate cover and incubate at room temperature for 2 h.

4. Wash the wells as in step 2.

5. Add 100  $\mu$ l per well of "Detection antibody", diluted 500x to a final concentration of 0.01  $\mu$ g/ml in "Assay buffer". Leave the blank wells empty. Cover the plate with adhesive plate cover and incubate at room temp for 60 min.

6. Wash the wells as in step 2.

7. Add 100  $\mu$ l/well of "SA-HRP" diluted 1000x in "SA-HRP diluent". Leave the blank wells empty. Cover the plate and incubate at room temp for 60 min.

8. Wash the wells as in step 2.

9. Develop by adding 100  $\mu l$  of "TMB substrate" to all wells (including the blank wells) and incubate at room temp in the dark for 15 min.

10. Stop the color development by adding 100  $\mu l$  of "Stop solution" to all wells (including the blank wells).

11. Measure the absorbance at 450 nm in a microplate reader within 15 min of the addition of the "Stop solution". If possible, use a reader capable of subtracting a reference wavelength between 570-650 nm.

12. The use of ELISA software utilizing e.g. a 4-parameter curve fitting program is recommended for the data analysis. Subtract the mean absorbance value of the blank from the standard, the assay background control and the sample values prior to creating the standard curve and determining the Trx1 concentrations in the samples. Note that Trx1 values obtained should be multiplied with the dilution factor used for each sample.

#### PERFORMANCE OF THE ASSAY

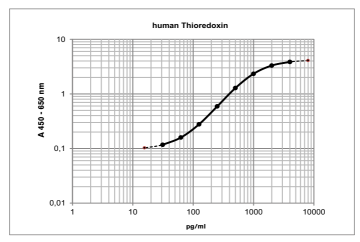


Figure 2. Display of the standard curve.

#### Standard range: 31.25-4000 pg/ml.

The standard range is the range in which determinations of analyte concentration can be done with precision, accuracy and linearity.

**Sensitivity:** The limit of detection of this assay is 15 pg/ml. It is the lowest concentration that is possible to detect but not necessarily quantify with precision and accuracy.

Accuracy: No international standard exists for calibration.

**Dilution recovery:** Dilution of plasma samples give a mid-curve recovery of 98% in repeated experiments.

Precision: The intraassay variation is 3.7%(CV). The interassay variation is 3.8%(CV).

#### LIMITATIONS OF THE ASSAY

#### Analysis of samples with high Trx1 content

The standard curve should not be extrapolated beyond the recommended standard range as these parts of the standard curve are non-linear. Samples yielding absorbance values exceeding the highest point of the standard range should be re-analyzed at a higher dilution.

#### Aberrant sera and plasma

The use of strongly hemolyzed and hyperlipemic serum and plasma samples may result in erroneous determinations of Trx1 concentrations.

#### Heterophilic antibodies in human serum and plasma

Heterophilic antibodies found in human serum/plasma are capable of binding to both the capture and detection antibodies used in capture ELISA. Heterophilic antibodies are found in a majority of human individuals and can, by cross-linking the assay antibodies used, result in false positive signals in capture ELISA. The "Assay buffer" provided and used for dilution of samples prevents the heterophilic antibodies from cross-linking the capture and detection mAbs. The Trx1 content of serum/ plasma samples can therefore be measured without interference by heterophilic antibodies. The lack of interference by heterophilic antibodies in this MABTECH kit has been validated using plasma/serum samples from normal healthy human blood donors. Please note that heterophilic antibody interference in samples from human subjects with various diseases or other conditions have not been assessed.

#### MABTECH AB Augustendalstorget 9 Box 1233, SE-131 28 Nacka Strand Sweden Tel: +46 8 716 27 00 Fax: +46 8 716 27 01 E-mail: mabtech@mabtech.com

MABTECH Inc M.E.B. 220, 3814 West Street Cincinnati, OH 45227 USA Toll free: +1 866 ELI-SPOT Tel: +1 513 871 4500 Fax: +1 513 871 7353 E-mail: mabtech.usa@mabtech.com

MABTECH Australia Pty Ltd Australia Tel: +61 3 9470 4704 Fax: +61 3 8678 3216 E-mail: mabtech.au@mabtech.com

MABTECH AB Büro Deutschland Germany Tel: +49 40 4135 7935 Fax: +49 40 4135 7945 E-mail: mabtech.de@mabtech.com

MABTECH AB Bureau de liaison France Tel: +33 (0)4 92 38 80 70 Fax:+33 (0)4 92 38 80 71 E-mail: mabtech.fr@mabtech.com



## NOTE; for research use only.

MABTECH shall not be liable for the use or handling of the product or for consequential, special, indirect or incidental damages therefrom.



Developed and manufactured by MA-BTECH AB, Sweden, whose quality management system complies

with the above standards.

For more information about Mabtech services and our products, visit

www.mabtech.com

Updated on 2019-04-24