

Sumoylated-HDAC1 ELISA Kit

Catalog #: PTE-SHDAC1

User Manual Last Revised: March 21, 2024

Introduction

The RayBio® Sumoylated-HDAC1 ELISA Kit is an in vitro enzyme-linked immunosorbent assay for the measurement of human and mouse sumoylated histone deacetylase 1. An antipan HDAC1 antibody has been coated onto a 96-well plate. Samples are pipetted into the wells and HDAC1 present in a sample is bound to the wells by the immobilized antibody. The wells are washed and biotinylated anti-sumoylation antibody is used to detect sumoylated HDAC1. After washing away unbound antibody, HRP-conjugated streptavidin is pipetted to the wells. The wells are again washed, a TMB substrate solution is added to the wells and color develops in proportion to the amount of sumoylated HDAC1 bound. The Stop Solution changes the color from blue to yellow, and the intensity of the color is measured at 450 nm.

Storage / Stability

The entire kit may be stored at -20°C for up to 6 months from the date of shipment. Avoid repeated freeze-thaw cycles. For prepared reagent storage, see kit contents on the next page.



Kit Components

Name	Catalog #	Size / Qty	Description	Storage / Stability After Preparation
Anti-Pan-HDAC1 Microplate	PTE- SHDAC1-A	96 wells	Microplate coated with antipan-HDAC1 antibody.	1 month at -20°C*
Positive Control	PTE- SHDAC1-PC	1 vial	Jurkat cell lysate	1 week at -20°C
Anti-Sumoylation Detection Antibody	PTE- SHDAC1-C1	2 vials	Biotinylated anti-Sumoylation antibody. 1 vial is enough to assay half the microplate.	5 days at 4°C
HRP-Streptavidin	EL-HRP	200 µl	100x concentrated HRP- conjugated streptavidin	Do not store and reuse.
Wash Buffer	EL-ITEMB	25 ml	20X concentrated wash buffer	1 month at 4°C
Assay Diluent B	EL-ITEME	15 ml	5X concentrated assay diluent	1 month at 4°C
Lysis Buffer	EL-Lysis	5 ml	2X cell lysate buffer	1 month at 4°C
TMB One-Step Substrate Reagent	EL-TMB	12 ml	3,3,5,5'-tetramethylbenzidine (TMB) in buffer solution	N/A
Stop Solution	EL-STOP	8 ml	0.2 M sulfuric acid	N/A

*Return unused wells to the pouch containing desiccant pack, reseal along entire edge.

Additional Materials Required

- Microplate reader capable of measuring absorbance at 450 nm
- Protease and Phosphatase inhibitors.
- Precision pipettes to deliver 2 µl to 1 ml volumes
- Adjustable 1-25 ml pipettes for reagent preparation
- 100 ml and 1 liter graduated cylinders
- Log-log graph paper or computer and software for ELISA data analysis.
- Absorbent paper
- Tubes to prepare positive control or sample dilutions
- Distilled or deionized water
- Shaker



Reagent Preparation

- 1. Bring all reagents and samples to room temperature (18 25°C) before use.
- 2. Assay Diluent B should be diluted 5-fold with deionized or distilled water before use.
- 3. Lysis Buffer should be diluted 2-fold with deionized or distilled water (for cell lysate and tissue lysate). We also recommend the addition of protease and phosphatase inhibitors (not included) to the lysis buffer prior to use.
- Preparation of Positive Control: Briefly spin the Positive Control Vial. Add 400 µl of prepared 1X Assay Diluent to prepare Positive Control solution. Gently mix the powder to allow it to dissolve thoroughly.
- 5. If the Wash Buffer concentrate contains visible crystals, warm to room temperature and mix gently until dissolved. Dilute 20 ml of Wash Buffer concentrate into deionized or distilled water to yield 400 ml of 1X Wash Buffer.
- 6. Preparation of Detection Antibody: Briefly spin the vial of biotinylated detection antibody. Add 100 μl of 1X Assay Diluent into the vial to prepare a detection antibody concentrate. Pipette up and down to mix gently (the concentrate can be stored at 4°C for 5 days or at -80°C for one month). The concentrate should then be diluted 55- fold with 1X Assay Diluent and used in step 4 of the Assay Procedure.
- 7. Preparation of HRP-Streptavidin: Briefly spin the vial of HRP-Streptavidin concentrate before use. HRP-Streptavidin should be diluted 100-fold with 1X Assay Diluent and used in step 7 of the Assay Procedure.

Assay Procedure

- 1. Bring all reagents and samples to room temperature (18 25°C) before use. It is recommended to run all positive control and samples in at least duplicate.
- 2. Label removable 8-well strips as appropriate for your experiment.
- Add 100 µI sample and Positive Control (see Reagent Preparation step 4) into appropriate wells. Cover the wells and incubate for 2.5 hours at room temperature or overnight at 4°C with gentle shaking.
- 4. Discard the solution and wash 4 times with 1X Wash Solution. Wash by filling each well with Wash Buffer (300 µl) using a multi-channel pipette or auto-washer. Complete removal of liquid at each step is essential for good performance. After the last wash, remove any remaining Wash Buffer by aspirating or decanting. Invert the plate and blot it against clean paper towels.



- Add 100 µl of prepared 1X detection antibody, anti-Sumoylation (see Reagent Preparation step 6) to each well. Incubate for 1 hour at room temperature with gentle shaking.
- 6. Discard the solution. Repeat the wash as in step 4.
- 7. Add 100 µl of prepared HRP-Streptavidin solution (see Reagent Preparation step 7) to each well. Incubate for 1 hour at room temperature with gentle shaking.
- 8. Discard the solution. Repeat the wash as in step 4.
- 9. Add 100 µl of TMB One-Step Substrate Reagent to each well. Incubate for 30 minutes at room temperature in the dark with gentle shaking.
- 10. Add 50 µl of Stop Solution to each well. Read at 450 nm immediately.

Assay Procedure Summary

- 1. Prepare all reagents, samples and Positive Control as instructed.
- 2. Add 100 μl sample and Positive Control to each well. Incubate 2.5 hours at room temperature or overnight at 4°C with gentle shaking.
- 3. Add 100 µl prepared detection antibody to each well. Incubate for 1 hour at room temperature with gentle shaking.
- 4. Add 100 µl prepared HRP-Streptavidin solution. Incubate for 1 hour at room temperature with gentle shaking.
- 5. Add 100 µl TMB One-Step Substrate Reagent to each well. Incubate 30 minutes at room temperature.
- 6. Add 50 µl Stop Solution to each well. Read at 450 nm immediately.

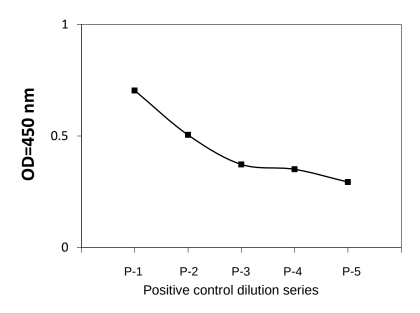


Typical Data

Calculate the mean absorbance for each set of duplicate positive controls, and samples.

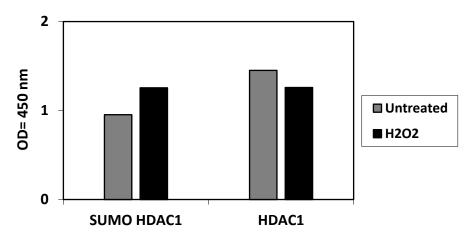
A. Positive Control

Jurkat cells were treated with Hydrogen Peroxide (H₂O₂). Solubilize cells at 4 x 10^7 cells/ml in Cell Lysate Buffer. Three fold serial dilutions of lysates were analyzed in this ELISA.



B. H₂O₂ Stimulation of Jurkat Cell Line

Jurkat cells were treated or untreated with H₂O₂. Cell lysates were analyzed using this ELISA.





Troubleshooting Guide

Problem	Cause	Solution	
Low signal in samples	 Sample concentration is too low Improper preparation of detection antibody Too brief incubation times Inadequate reagent volumes or improper dilution 	 Increase sample concentration Briefly spin down vials before opening. Dissolve the powder thoroughly. Ensure sufficient incubation time; assay procedure step 3 may be done overnight Check pipettes and ensure correct preparation 	
High signal in samples	Sample concentration is too high	Reduce sample concentration	
Large CV	 Inaccurate pipetting Air bubbles in wells 	Check pipettesRemove bubbles in wells	
High background	Plate is insufficiently washedContaminated wash buffer	 Review the manual for proper wash. If using a plate washer, ensure that all ports are unobstructed. Make fresh wash buffer 	
Low sensitivity	 Improper storage of the ELISA kit Stop solution Improper primary or secondary antibody dilution 	 Store your positive control at <- 70°C after reconstitution, others at 4°C. Keep substrate solution protected from light. Add stop solution to each well before reading plate Ensure correct dilution 	