

Ubiquitin target protein ELISA quantitative assay kit

Cat. No. BGT-OPU-09002

Instruction Manual

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Intended Use

BiogradeTech' Ubiquitin target protein ELISA quantitative assay kit is intended for the relative determination of the concentration of a specific ubiquitylated, target protein in cells. This assay is designed to replace more laborious, semi-quantitative immunoprecipitation and Western blots to examine changes in ubiquitylation.

This kit is for research use only and is not intended for human or animal diagnostic or therapeutic applications.

Principle of the assay

The Ubiquitin target protein ELISA quantitative assay kit is a sandwich ELISA assay in which total ubiquitylated proteins in a cell lysate are captured in the wells of a precoated microtiter plate using a proprietary ubiquitin binding reagent. Unbound protein is removed by washing and then the amount of bound target protein is determined using either an antibody specific to the target protein or to an epitope tag incorporated in to the target protein. After removing unbound antibody, the amount bound is measured using an enzyme linked anti-antibody.

Components

Unless otherwise noted, all components should be stored at 4°-8°C.

Coated plates: One (1) pre-coated 96-well strip plate is provided. The

plate is dried and vacuum packed. If not used within one

(1) month remove plate and store at -80°C.

Strip holder: One (1) empty strip plate frame is provided.

Primary Antibody (PA1): 1 vial of a 200x concentrate of an anti-epitope mAb in PBS

containing 0.02% NaN₃.

Available mAbs include anti-GST, anti-FLAG®, anti-myc,

anti-HA, anti-V5

Blocking concentrate (BC): 12 mL of 5X blocking agent

Detection Antibody (DA1): 1 vial of 200X anti-mouse HRP conjugate

Plastic plate seals: Four (4) provided

Detection reagent 1 (DR1): 0.5 mL vial of Luminol reagent

Detection reagent 2 (DR2): 0.5 mL vial of stabilized peroxide reagent

Components required but not supplied

Phosphate buffered saline (PBS) and PBS containing 0.1% Tween 20 (PBST)

Disposable gloves, pipette tips, reagent reservoirs

Multi-channel pipettors

Automatic plate washer (optional)

A luminescence microplate reader

Short Protocol

*The listed amounts of components below are for a full 96-well plate. Please make the appropriate calculations based on (1) the number of test samples, (2) the number of duplicates, triplicates, etc., (3) the number of dilutions desired (see Detailed protocol for the information on suggested dilutions).

Allow all components to come to room temperature

Prepare Diluent **A** by adding **four (4) parts of** PBS to **one (1)** part of 5X BC.

12 ml of 5X BC provided is sufficient to make 60 mL of Buffer A.

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Dilute test-samples to 25 to 400 µg/mL in PBS in microfuge tubes.

Pipette 100 μL of each in appropriate wells of the <u>UbiQuant™ plate</u> (See detailed protocol).



Incubate at room temperature for 1 hr.



Wash plate four (4) times with PBST.



For full- plate add **60** μ L of PA1 into **12** mL of diluent A and pipette **100** μ L into each well of the microtiter plate.

*Example: add 30µL into 6 mL for half- plate, or make adjustments as necessary Incubate at room temperature for 1 hr.



Wash plate four (4) times with PBST



If using a full-plate, add **60 μL** of DA1 conjugate to **12 mL** of diluent A.

Pipette 100 µL into each well.



Incubate at room temperature for 1 hr.



Wash plate four (4) times with PBST.



If using a whole plate, mix 150 μ L DR1 and 150 μ L DR2 in 10mL of ultra-pure water. Pipette 100 μ L into each well.

Wait two minutes and read the plate

Detailed Protocol

1. We recommend that all samples be carried out in triplicate. Remove kit from refrigerator and allow all components to reach room temperature. All incubations are performed at room temperature (22°-27°C). To avoid cross contamination, do not re-use plate sealers. The amounts listed below should be enough for one 96-well plate or several samples run in triplicate. Please calculate the number of wells based on the number of test samples, the number of triplicates, etc., and how you will perform the dilutions (Step 4b).

2. Prepare reagents:

- a. <u>Diluent A.</u> Prepare Diluent **A** by adding **four (4)** parts of PBS to **one (1)** part of 5X BC and vortex to mix.
- b. <u>Test Samples:</u> Dilute test samples in PBS of choice to give between 25 and 400 μg/mL. The final volume will depend on the number of triplicates you wish to do for each sample. Alternatively, perform a serial two-fold dilution of each test sample.
- c. Dilute **30** μ L of primary antibody **(PA1)** into **6** mL of Diluent **A** (or 60μ L in 12 ml if using a whole plate).
- d. Dilute **30** μ L of detection antibody **(DA1)** into **6** mL of Diluent **A** (or 60μ L in 12 ml if using a whole plate).
- 3. Determine the number of strip wells required and place them into the holder supplied. Return the unused strips to the bag, tape closed, and refrigerate.
- 4. Pipette 100 μ L each of: blank, controls, and test samples into appropriate wells and cover with plastic plate sealer. Incubate 1 hr.
- 5. Wash plate 4 times with \sim 150 μ L/well PBST using a multichannel pipette, a squeeze bottle, or an automatic plate washer. After the last wash, remove the last droplets of buffer by lightly tapping the plate (upside down) on paper towels or other blotting paper. DO NOT ALLOW WELLS TO DRY COMPLETELY.
- 6. After blotting, add 100 μ L of diluted PA1 to each well, cover with plastic plate seal and incubate 1 hr.
- 7. Wash wells as described in step 5.
- 8. Add 100 μ L of diluted DA1 to each well, cover and incubate 1 hr
- 9. Wash wells as described in step 5.
- 10. Just before use, mix 150 μ L of DR1 and 150 μ L of DR2 into 10 mL of ultrapure water (deionized or distilled). Add 100 μ L of this solution to each well, wait 2 min and read in a plate reader optimized for chemiluminescence. Although there is a gradual decrease in the total signal with time, the plate may be read up to 20 min after addition of the luminescence reagent.

Data Reduction

- 1. Calculate the mean counts for blank, controls, and unknowns. Subtract the mean counts for the blank from each sample.
- 2. The data can be plotted as a bar graph or as a percent of the maximum value. If performing a compound dose-response assay, the maximum value should correspond to a zero concentration dose and the data can be fit to a sigmoidal plot to determine the ED_{50} .

Example: Cell Lysis Protocol

- 1. Aspirate the medium completely and rinse cells with ice cold 1X PBS (5ml PBS/10cm dish). Add 5ml of 1X PBS and scrape cells into the PBS using a cell scraper.
- 2. Transfer cells into a 15 mL conical tube and spin at 3000 rpm for 5 min. Freeze the cell pellet at
 - 80°C for long term storage or take out after 30 minutes to continue with lysis.

Note: After rinsing with ice cold PBS, cells can also be lysed by directly adding lysis buffer to the plate and scraping cells into the eppendorf tube.

- 3. Place the frozen cell pellet on ice and add RIPA lysis buffer (5-10 times the volume of the pellet, (i.e. $100 \mu L$ of pellet add $500-1000 \mu L$ lysis buffer). Vortex intermittently for about 10-15 minutes
- 4. Centrifuge at 13,000 rpm for 15-20 min at 4° C.
- 5. Collect the supernatant (lysate) and determine protein concentration using standard methods.
- 6. Proceed with the dilution of the lysate for the assay (2b in the Detailed protocol).

RIPA Lysis Buffer

50 mM Tris-HCl, pH 7.5 150 mM NaCl 1% NP40 1% Sodium deoxycholate 2 mM EDTA 10% Glycerol 1 mM PMSF

Protease inhibitor cocktail (Sigma cat #P8849, 1:500)

Aprotinin (10mg/ml stock) 20 µL/10 mL

50 uM PR619 (cat.no.BGT-CRS-09001) non-selective DUB inhibitor - recommended to protect ubiquitylated proteins from degradation by DUBs

5 mM 1,10-phenanthroline (oPA, cat.no. BGT-CRS-09002) – metalloproteinase inhibitor recommended to protect K63-linked polyubiquitin chains from degradation by JAMM-type DUBs