

# **Proteasome Activity Assay Kit**

(Catalog #K245-100; 100 assays; Store kit at -20°C)

#### I. Introduction:

Proteasomes are very large (20S, 26S) protein assemblies found in both the nucleus and cytoplasm of all eukaryotes (and in some prokaryotes). They are responsible for the degradation and recycling of proteins which have been previously tagged with ubiquitin. Such tagged proteins are degraded into peptides approximately 7 - 8 amino acids long which are subsequently further degraded. The 20S assembly is the functional protease structure with chymotrypsin-like, trypsin-like and caspase-like protease activities. BioVision's Proteasome Activity Assay takes advantage of the chymotrypsin-like activity, utilizing an AMC-tagged peptide substrate which releases free, highly fluorescent AMC in the presence of proteolytic activity. The kit also includes a positive control (Jurkat Cell lysate with significant proteasome activity) and a specific proteasome inhibitor MG-132 which suppresses all proteolytic activity due to proteasomes. This permits differentiation of proteasome activity from other protease activity which may be present in samples.

#### II. Kit Contents:

Components	100 assays	Cap Code	Part Number
Proteasome Assay Buffer Proteasome Substrate (Succ-LLVY-AMC in DMSO) Proteasome Inhibitor (MG-132 in DMSO) AMC Standard (1 mM in DMSO) Positive Control	25 ml	WM	K245-100-1
	100 μl	Red	K245-100-2
	100 μl	Blue	K245-100-3
	100 μl	Yellow	K245-100-4
	lyophilized	Green	K245-100-5

#### III. Storage and Handling:

Store the kit at -20 °C, protect from light. Read the entire protocol before performing assay. Avoid repeated freeze/thaw cycles. All samples and the Positive Control should be assayed in duplicate, (once in the absence and once in the presence of the Proteasome Inhibitor). An opaque white microwell plate is recommended. This protocol is designed for use in a 96 well plate. 384-well plates may be used but all reagent amounts should be reduced 5-fold (diluted if necessary). Do not use protease inhibitors during cell lysate preparation.

**Proteasome Substrate, Proteasome Inhibitor, AMC Standard:** Ready to use as supplied. These DMSO solutions must be warmed to room temperature prior to use to melt frozen DMSO. We recommend warming in a 37 °C water bath, pipetting up and down to ensure they are completely melted and mixed before use. Store at -20 °C in the dark when not in use.

**Positive Control:** Reconstitute with 100  $\mu$ l dH<sub>2</sub>O. If kit will be used multiple times over an extended period of time, aliquot portions and store at -80 °C. Keep on ice while in use. Avoid repeated freeze/thaw cycles.

### IV. Assay Protocol:

- AMC Standard Preparation: Dilute AMC Standard 100-fold (10 μl + 990 μl dH<sub>2</sub>O) then add 0, 2, 4, 6, 8, 10 μl of AMC standard to a series of microplate wells. Adjust volume to 100 μl/well with Assay Buffer to generate 0, 20, 40, 60, 80 and 100 pmol per well AMC Standard.
- 2. **Positive Control Preparation:** Add 10 µl of the positive control to paired wells. Bring volume to total 100 µl by adding 90 µl of Assay Buffer to each well.
- 3. **Samples:** Prepare by homogenizing cells with 0.5 % NP-40 in dH<sub>2</sub>O or PBS. Add up to 50 μl of each cell extract or other proteasome sample to be tested to paired wells. Bring the volume of each well to 100 μl with Assay Buffer.
- 4. **Inhibitor**: Add 1 μI of the Proteasome inhibitor to one of the paired wells, 1 μI of Assay Buffer to the other well, mix.
- 5. Substrate: Add 1 ul of Proteasome Substrate to all wells, mix, protected from light, mix.

6. Read: Measure kinetics of fluorescence development at Ex/Em = 350/440 nm in a microplate reader at 37°C for 30 - 60 min. There is a slight lag and nonlinearity to the kinetics due to the time it takes for the reaction mix to warm up to 37°C. Measurement of the wells which do not contain Proteasome Inhibitor will show total proteolytic activity RFU₁ and the wells containing Proteasome Inhibitor will show non-proteasome activity iRFU₁ at T₁. Measure RFU₂ and iRFU₂ at T₂ after 30 min (or longer time if the sample activity is low). The RFU generated by proteasome activity is ΔRFU = (RFU₂ – iRFU₂) – (RFU₁ – iRFU₁).

**Note:** It is essential to read RFU<sub>1</sub>, iRFU<sub>2</sub> and iRFU<sub>2</sub> in the linear reaction range. It will be more accurate if you monitor the reaction kinetics as shown in Fig. 1B, then choose  $T_1$  and  $T_2$  in the appropriate linear range. From our experience, initial readings RFU<sub>1</sub> and iRFU<sub>1</sub> should be measured after ~ 20 - 25 min.

 Calculation: Plot the AMC Standard Curve. Apply the ΔRFU to the AMC Standard Curve to get B pmol of AMC (amount generated between T<sub>1</sub> and T<sub>2</sub> in the reaction wells specifically by proteasome activity).

Proteasome Activity = 
$$\frac{B}{(T2-T1)\times V}$$
 × Sample Dilution Factor = nmol/min/ml = U/ml

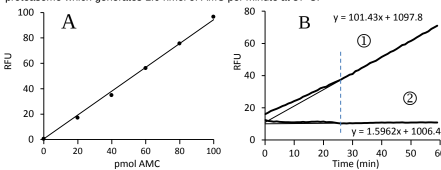
Where: **B** is the AMC amount from AMC Standard Curve (in pmol)

T<sub>1</sub> is the time of the first reading (RFU<sub>1</sub> and iRFU<sub>1</sub>) (in min)

T<sub>2</sub> is the time of the second reading (RFU<sub>2</sub> and iRFU<sub>2</sub>) (in min)

V is the pretreated sample volume added into the reaction well (in μl)

**Proteosome Unit Definition:** One unit of proteasome activity is defined as the amount of proteasome which generates 1.0 nmol of AMC per minute at 37°C.



**Fig. 1. AMC Standard Curve and Proteasome Activity assay using the kit protocol: A:** AMC standard curve 0-100 pmole; **B:** Kinetics of Proteasome Activity assay in the absence ① and presence ② of MG-132 Proteasome inhibitor. Equations represent best fit of lines during the linear portion of the reaction (after ~ 25 min in this case).

#### **RELATED PRODUCTS:**

Jurkat Cell Extract
Caspase and Cathepsin Inhibitors
Caspases: Substrates and Assay Kits
Cathepsins: Substrates and Assay Kits
EZBlock™ Protease Inhibitor Cocktails

Caspase, Cathepsin, Calpain active proteins Proteasome/Calpain Substrate and Assay Kits MG-132 Proteasome Inhibitor Protease Inhibitor Cocktails Cell Fractionation Kits

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## **GENERAL TROUBLESHOOTING GUIDE:**

Problems	Cause	Solution			
Assay not working	Use of ice-cold assay buffer	Assay buffer must be at room temperature			
	Omission of a step in the protocol	Refer and follow the data sheet precisely			
	Use of a different 96-well plate	• Fluorescence: Black plates (clear bottoms) ; Luminescence: White plates ; Colorimeters: Clear plates			
Samples with erratic readings	Use of an incompatible sample type	Refer data sheet for details about incompatible samples			
	Samples prepared in a different buffer	Use the assay buffer provided in the kit or refer data sheet for instructions			
	Cell/ tissue samples were not completely homogenized	Use Dounce homogenizer (increase the number of strokes); observe for lysis under microscope			
	Samples used after multiple free-thaw cycles	Aliquot and freeze samples if needed to use multiple times			
	Presence of interfering substance in the sample	Troubleshoot if needed, deproteinize samples			
	Use of old or inappropriately stored samples	Use fresh samples or store at correct temperatures till use			
Lower/ Higher readings in Samples and Standards	Improperly thawed components	Thaw all components completely and mix gently before use			
	Use of expired kit or improperly stored reagents	Always check the expiry date and store the components appropriately			
	Allowing the reagents to sit for extended times on ice	Always thaw and prepare fresh reaction mix before use			
	Incorrect incubation times or temperatures	Refer datasheet & verify correct incubation times and temperatures			
	Incorrect volumes used	Use calibrated pipettes and aliquot correctly			
Readings do not follow a linear pattern for Standard curve	Use of partially thawed components	Thaw and resuspend all components			
	Pipetting errors	Avoid pipetting small volumes			
	T1 readings too early	Usuually wait ~ 25 min before reading T1 (to get past lag phase)			
	Air bubbles formed in well	Pipette gently against the wall of the tubes			
	Standard stock is at an incorrect concentration	Always refer the dilutions in the data sheet			
	Calculation errors	Recheck calculations after referring the data sheet			
	Substituting reagents from older kits/ lots	Use fresh components from the same kit			
Unanticipated results	Measured at incorrect wavelength	Check the equipment and the filter setting			
	Samples contain interfering substances	Troubleshoot if it interferes with the kit			
	Use of incompatible sample type	• Refer data sheet to check if sample is compatible with the kit or optimization is needed			
	Sample readings above/below the linear range	Concentrate/ Dilute sample so as to be in the linear range			
Note: The most probable list of c	ote: The most probable list of causes is under each problem section. Causes/ Solutions may overlap with other problems.				