



# HDAC Inhibitor Drug Screening Kit (Fluorometric)

(Catalog #K340-100; 100 assays; Store kit at -80°C)

#### I. Introduction:

Inhibition of histone deacetylase (HDAC) has been implicated to modulate transcription, to induce apoptosis or differentiation in cancer cells. However, screening of compounds for HDAC inhibition has been difficult due to the lack of convenient tools for analyzing HDAC activity. The new HDAC Inhibitor Drug Screening Kit provides a fast, fluorescence-based method that eliminates radioactivity, extractions, or chromatography, as used in traditional assays. The new procedure requires only two easy steps, both performed on the same microtiter plate. First, your inhibitor candidates are mixed with HeLa Nuclear Extract and HDAC fluorometric substrate, which comprises an acetylated lysine side chain. Deacetylation of the substrate sensitizes the substrate, so that, in the second step, treatment with the Lysine Developer produces a fluorophore. The fluorophore can be easily analyzed using a fluorescence plate reader or a fluorometer. The assay is well suited for high throughput screening applications.

#### II. Kit Contents:

Component	K340-100	Color Code	Part
	100 assays	Cap Color	Number
HDAC Substrate	500 μl	Amber	K340-100-1
10X HDAC Assay Buffer	1.0 ml	Green	K340-100-2
Lysine Developer	1.0 ml	Orange	K340-100-3
HDAC Inhibitor (Trichostatin A, 1 mM)	10 µl	Blue	K340-100-4
HeLa Nuclear Extract (5 mg/ml)	200 µl	Red	K340-100-5

## III. Storage and Handling:

Store kit at -80°C, protected from light. Warm HDAC Assay Buffer to room temperature (RT) before use. Briefly centrifuge all small vials before opening. Read the entire protocol before performing the assay. Aliquot & store HeLa Nuclear Extract at -80°C to avoid loss of activity. The kit provides sufficient reagents for 100 Positive Control assays with the HeLa Nuclear Extract and 5 Negative Control assays with the HDAC Inhibitor, Trichostatin A.

## IV. HDAC Assay Protocol:

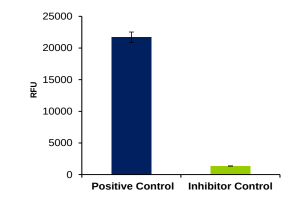
- Screen compounds, Inhibitor Control and Positive Control Preparations: Dissolve candidate inhibitors into proper solvent. Dilute to 2X the desired test concentration with ddH<sub>2</sub>0. Add 50 µl of diluted candidate inhibitor into well(s). For Positive Control, add 50 µl ddH<sub>2</sub>0 only. For Negative Control, add 48 µl of ddH<sub>2</sub>0 and 2 µl of Trichostatin A.
- 2. Reaction Mix Preparation: Mix enough reagents for the number of assays to be performed. For each well, prepare 50 µl Reaction Mix containing:

	Reaction Mix
10X HDAC Assay Buffer	10 µl
HeLa Nuclear Extract	2 μl
HDAC Substrate	5 µl
ddH <sub>2</sub> 0	33 µl

Mix well. Add 50  $\mu l$  of the Reaction Mix into each well. Mix well. Incubate plate at 37°C for 30 min (or longer if desired).

- 3. Stop the reaction by adding 10  $\mu l$  of Lysine Developer and mix well. Incubate the plate at 37°C for 30 min.
- 4. **Measurement:** Read sample in a fluorescence plate reader with Ex. = 350-380 nm and Em. = 440-460 nm. Signal should be stable for several hours at RT.
- 5. **Calculation**: Set the RFU of Positive Control as the 100%, and calculate the relative activity remains with candidate compounds as follow.





## **RELATED PRODUCTS:**

- HDAC Fluorometric Assay kit
- HDAV Drug Screening Kit
- HDAC Inhibitors & Set
- HeLa Nuclear Extract
- HAT Activity Assay Kit
- Histone H2A, H2B, H3 & H4 Antibodies
- HDAC (1-11) Polyclonal Antibodies & Set

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# GENERAL TROUBLESHOOTING GUIDE:

Problems	Cause	Solution	
Assay not working	Use of ice-cold assay buffer	Assay buffer must be at RT	
	Omission of a step in the protocol	Refer and follow the data sheet precisely	
	Plate read at incorrect wavelength	Check the wavelength in the data sheet and the filter settings of the instrument	
	Use of a different 96-well plate	• Fluorescence: Black plates ; 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	
	Samples were not deproteinized (if indicated in datasheet)	Use the 10 kDa spin cut-off filter (BV Cat# 1997) or PCA precipitation as indicated	
	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	
	<ul> <li>Use of old or inappropriately stored samples</li> </ul>	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	
	<ul> <li>Use of expired kit or improperly stored reagents</li> </ul>	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	
	<ul> <li>Incorrect incubation times or temperatures</li> </ul>	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 before preparing the reaction mix	
	Pipetting errors in the reaction mix	Prepare a master reaction mix whenever possible	
	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	
lote: The most probable list of causes is under each problem section. Causes/ Solutions may overlap with other problems.			