

Product Information

Caspase-8 Fluorometric Assay Kit

I. Kit Contents:

Component	K2012-25	K2012-100	K2012-200	K2012-400	Part Number
	25 assays	100 assays	200 assays	400 assays	
Cell Lysis Buffer	25 ml	100 ml	100 ml	100 ml	K2012-C-1
2X Reaction Buffer	2 ml	4 x 2 ml	16 ml	32 ml	K2012-C-2
IETD-AFC (1 mM)	125 µl	500 µl	2 x 0.5 ml	2 x 1 ml	K2012-C-3
DTT (1 M)	100 µ1	400 µ1	400 µ1	400 µ1	K2012-C-4

II. Introduction:

Cysteine-dependent aspartate-directed proteases (Caspases) are a family of cysteine proteases that play important roles in apoptosis, necrosis, and inflammation. Sequential activation of caspases plays an important role in cell apoptosis. Caspase-8 is a caspase protein that cleaves and activates Caspase-3 and is involved in the programmed cell death induced by Fas and different apoptotic stimuli. Caspase-8 was detected in the insoluble fraction of the affected brain region in Huntington disease patients, which suggested it plays an important role in neurodegenerative diseases.

Caspase-8 Fluorometric Assay Kit provides a convenient and simple way for detecting the IETD-dependent caspase activity. IETD-AFC (AFC: 7-amino-4-trifluoromethyl coumarin) emits blue light (λ max = 400 nm); while cleavage of IETD-AFC by Caspase-8 or related caspases, free AFC emits a yellow-green fluorescence (λ max = 505 nm), which can be quantified by using a fluorecence microtiter plate reader or a fluorometer. Comparison of the fluorescence of AFC from an apoptotic sample with an uninduced control determines the fold increase in Caspase-8 activity.

III. Caspase-8 Assay Protocol:

A. General Considerations

Aliquot enough 2X Reaction Buffer for the number of assays to be performed. Add DTT to the 2X Reaction Buffer immediately before use (10 mM final concentration: add 10 µl of 1.0 M DTT stock per 1 ml of 2X Reaction Buffer).

After thawing, store the Cell Lysis Buffer and 2X Reaction Buffer at 4 0

Protect IETD-AFC from light.

B. Assay Procedure

1. Induce apoptosis in cells by desired method. Concurrently incubate a control culture without induction.

2. Count cells and pellet 1-5 x 10⁶ cells or use 50-200 µg cell lysates if protein concentration has been measured.

3. Resuspend cells in 50 µl of chilled Cell Lysis Buffer. Incubate cells on ice for 10 minutes.

4. Add 50 μ l of 2X Reaction Buffer (containing 10 mM DTT) to each sample. Add 5 μ l of the 1 mM IETD-AFC substrate (50 μ M final concentration). Incubate at 37 for 1-2 hours.

5. Read samples in a fluorometer equipped with a 400-nm excitation filter and 505-nm emission filter. For a plate-reading set-up, transfer the samples to a 96-well plate. You may also perform the entire assay directly in a 96-well plate.

Fold-increase in FLICE activity can be determined by comparing these results with the level of the uninduced control.

IV. Storage and Stability:



Store kit at -20 (Store Cell Lysis Buffer and 2X Reaction Buffer at 4° C after opening). All reagents are stable for 6 months under proper storage conditions.

Problems	Cause	Solution	
Assay not working	• Cells did not lyse completely	• Resuspend the cell pellet in the lysis buffer and incubate as	
	• Experiment was not performed at optimal time after	described in the datasheet	
	apoptosis induction	• Perform a time-course induction experiment for apoptosis	
	Plate read at incorrect wavelength	• Check the wavelength listed in the datasheet and the filter	
	• Old DTT used	settings of the instrument	
		• Always use freshly thawed DTT in the cell lysis buffer	
High Background	• Increased amount of cell lysate used	• Refer to datasheet and use the suggested cell number to	
	• Increased amounts of components added due to incorrect	prepare lysates	
	pipetting	• Use calibrated pipettes	
	• Incubation of cell samples for extended periods	• Refer to datasheet and incubate for exact times	
	• Use of expired kit or improperly stored reagents	· Always check the expiry date and store the individual	
	Contaminated cells	components appropriately	
		Check for bacteria/ yeast/ mycoplasma contamination	
Lower signal	Cells did not initiate apoptosis	• Determine the time-point for initiation of apoptosis after	
levels	• Very few cells used for analysis	induction (time-course experiment)	
	• Use of samples stored for a long time	• Refer to datasheet for appropriate cell number	
	• Incorrect setting of the equipment used to read samples	• Use fresh samples or aliquot and store and use within one	
	• Allowing the reagents to sit for extended times on ice	month for the assay	
		• Refer to datasheet and use the recommended filter setting	
		• Always thaw and prepare fresh reaction mix before use	
Samples with	• Uneven number of cells seeded in the wells	• Seed only equal number of healthy cells (correct passage	
erratic readings	Samples prepared in a different buffer	number)	
	• Adherent cells dislodged and lost at the time of experiment	• Use the cell lysis buffer provided in the kit	
	Cell/ tissue samples were not completely homogenized	• Perform experiment gently and in duplicates/triplicates;	
	Samples used after multiple freeze-thaw cycles	apoptotic cells may become floaters	
	• Presence of interfering substance in the sample	• Use Dounce homogenizer (increase the number of strokes);	
	• Use of old or inappropriately stored samples	observe efficiency of lysis under microscope	
		• Aliquot and freeze samples, if needed to use multiple times	
		Troubleshoot as needed	
		• Use fresh samples or store at correct temperatures until use	
Unanticipated	Measured at incorrect wavelength	• Check the equipment and the filter setting	
results	Cell samples contain interfering substances	• Troubleshoot if it interferes with the kit (run proper	
		controls)	
General issues	Improperly thawed components	• Thaw all components completely and mix gently before use	
	• Incorrect incubation times or temperatures	• Refer to datasheet & verify the correct incubation times and	
	• Incorrect volumes used	temperatures	
	• Air bubbles formed in the well/tube	• Use calibrated pipettes and aliquot correctly	

General Troubleshooting	g Guide for Cas	pase Colorimetric and I	Fluorometric Kits:
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Substituting reagents from older kits/ lots	• Pipette gently against the wall of the well/tubes		
• Use of a different 96-well plate	• Use fresh components from the same kit		
	• Fluorescence: Black plates; Absorbance: Clear plates		
Note# The most probable cause is listed under each section. Causes may overlap with other sections.			

For research use only! Not to be used in humans.

Our promise

If the product does not perform as described on this datasheet, we will offer a refund or replacement. For more details, please visit <u>http://www.apexbt.com/</u> or contact our technical team.

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