

ab65611

pan-Caspase (active) FITC Staining Kit

Instructions for Use

For the rapid, sensitive and accurate detection of active Caspases in living cells

This product is for research use only and is not intended for diagnostic use.

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1. Overview

Activation of Caspases plays a central role in apoptosis. Abcam's pan-Caspase (active) FITCStaining Kit provides a convenient and sensitive means for detecting activated Caspases in living cells. The assay utilizes the Caspase family inhibitor VAD-FMK conjugated to FITC (FITC-VAD-FMK) as a marker. FITC-VAD-FMK is cell permeable, nontoxic, and irreversibly binds to activated Caspases in apoptotic cells.

2. Protocol Summary

Induce Apoptosis in Cell Samples

Add FITC-VAD-FMK

Analyze by Flow Cytometry

OR

Detect by Fluorescence Microscopy

OR

Measure Fluorescence in Microplate Reader

3. Components and Storage

A. Kit Components

Item	Quantity
FITC-VAD-FMK	100 µL
Wash Buffer	2 x 100 mL
Z-VAD-FMK	10 µL

^{*} Store kit at -20°C.

B. Additional Materials Required

- Microcentrifuge
- Pipettes and pipette tips
- Fluorescent microplate reader or microscope
- Flow cytometer
- Black microtiter plate
- Orbital shaker

4. Assay Protocol

1. Sample Preparation:

a) Induce apoptosis in cells (1 x 10⁶/ml) by desired method.
Concurrently incubate a control culture without induction.

An additional **negative control** can be prepared by adding the Caspase inhibitor Z-VAD-FMK at 1 µl/ml to an induced culture to inhibit Caspase activation.

Notes:

This product detects proteolytic activity. Do not use protease inhibitors in the sample preparation step as it might interfere with the assay.

- **b)** Aliquot 300 μl each of the induced and control cultures into eppendorf tubes.
- c) Add 1 μI of FITC-VAD-FMK into each tube and incubate for 0.5-1 hour at 37°C incubator with 5% CO₂.
- **d)** Centrifuge cells at 3000 rpm for 5 minutes and remove supernatant.
- **e)** Re-suspend cells in 0.5 ml of Wash Buffer, and centrifuge again.
- f) Repeat Step e.

g) Proceed to Step 2, 3 or 4 depending on methods of analysis.

2. Quantification by Flow Cytometry:

For flow cytometric analysis, re-suspend cells in 300 μ l of Wash Buffer. Keep samples on ice. Analyze samples by flow cytometry using the FL-1 channel.

3. Detection by Fluorescence Microscopy:

For fluorescence microscopic analysis, re-suspend cells in 100 μ l Wash Buffer. Put one drop of the cell suspension onto a microslide and cover with a coverslip.

Observe cells under a fluorescence microscope using FITC filter. Caspase positive cells appear to have brighter green signals, whereas Caspase negative control cells show much weaker signal.

4. Analysis by Fluorescence Plate Reader:

For analysis with fluorescence plate reader, re-suspend cells in $100~\mu l$ Wash Buffer and then transfer the cell suspension to each well in the black microtiter plate.

Measure the fluorescence intensity at Excitation = 485 nm and Emission = 535 nm. For control, use wells containing unlabeled cells.

5. Troubleshooting

Problem	Reason	Solution
	Cell density is	Refer to datasheet and use the
High Background	higher than	suggested cell number
	recommended	
	Increased volumes of components added	Use calibrated pipettes accurately
	Incubation of cell samples for extended periods	Refer to datasheets and incubate for exact times
	Use of extremely confluent cells	Perform assay when cells are at 80-95% confluency
	Contaminated cells	Check for bacteria/ yeast/ mycoplasma contamination
Lower signal levels	Cells did not initiate apoptosis	Determine the time-point for initiation of apoptosis after induction (time-course experiment)
	Very few cells used for analysis	Refer to data sheet for appropriate cell number
	Incorrect setting of the equipment used to read samples	Refer to datasheet and use the recommended filter setting
	Use of expired kit or improperly stored reagents	Always check the expiry date and store the components appropriately

Erratic results	Uneven number of cells seeded in the wells	Seed only healthy cells (correct passage number)
	Adherent cells dislodged at the time of experiment	Perform experiment gently and in duplicates or triplicates for each treatment
	Incorrect incubation times or temperatures	Refer to datasheet & verify correct incubation times and temperatures
	Incorrect volumes used	Use calibrated pipettes and aliquot correctly

For further technical questions please do not hesitate to contact us by email (technical@abcam.com) or phone (select "contact us" on www.abcam.com for the phone number for your region).



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