

Product Information

Red Active Caspase-9 Staining Kit

I. Kit Contents:

Component	K2055-25	K2055-100	Part Number
	25 assays	100 assays	
Red-LEHD-FMK	25 µl	100 µl	K2055-C-1
Wash Buffer	50 ml	2 x 100 ml	K2055-C-2
Z-VAD-FMK	10 µl	10 µl	A1902

II. Introduction:

Caspases activation is curial in apoptosis. Red Caspase-9 Staining Kit is an easy and sensitive way of detecting activated caspases in living cells. This assay uses the Caspase-9 inhibitor, LEHD-FMK, coupled to sulfo-rhodamine (Red-IEHD-FMK) as a marker. In apoptotic cells, the cell permeable and nontoxic Red-LETD-FMK irreversibly binds to activated Caspase-9.

III. Caspase-9 Assay Protocol:

A. Staining Procedure:

1. Induce apoptosis in cells (1 x 10^6 /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.

2. Aliquot 300 μ l each of the induced and control cultures into eppendorf tubes.

3. Add 1 µl of Red-LEHD-FMK into each tube and incubate for 0.5 - 1 hour at 37 °C incubator with 5% CO2.

4. Centrifuge cells at 3000 rpm for 5 minutes and remove supernatant.

5. Resuspend cells in 0.5 ml of Wash Buffer, and centrifuge again.

6. Repeat Step 5.

Proceed to B, C, or D depending on methods of analysis.

B. Quantification by Flow Cytometry:

For flow cytometric analysis, resuspend cells in 300 µl of Wash buffer. Put samples on ice. Analyzing samples by flow cytometry using the FL-2 channel.

C. Detection by Fluorescence Microscopy:

For fluorescence microscopic analysis, resuspend cells in $100 \ \mu$ 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 rhodamine filter. Caspase-9 positive cells appear to have brighter red signals, whereas Caspase-9 negative control cells show much weaker signal.

D. Analysis by Fluorescence Plate Reader:

For analysis with fluorescence plate reader, resuspend cells in 100 μ l Wash Buffer and then transfer the cell suspension to each well of the black microtiter plate. Measure the fluorescence intensity at Ex/Em = 540/570 nm (Note: Ex/Em=488/570 nm will also work, although it's not an optimal wavelength). For control, use wells containing unlabeled cells.



General Troubleshooting Guide:

Problems	Cause	Solution	
High Background	• Cell density is higher than recommended	• Refer to data sheet and use the suggested cell number	
	• Cells were not washed well with wash buffer after staining	• Use the wash buffer provided, and as instructed in the	
	• Cells were Incubated for extended period of time	datasheet	
	• Use of extremely confluent cells	• Refer to data sheets and incubate for exact times	
	Contaminated cells	• Perform assay when cells are at 70-95% confluency	
		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)	
	• Incorrect setting of the equipment used to read samples	• Refer to data sheet for appropriate cell number	
	• Use of expired kit or improperly stored reagents	• Refer to data sheet and use the recommended filter setting	
		• Always check the expiry date and store the components	
		appropriately	
Erratic results	• Old (unhealthy) cells used	• Seed healthy cells and make sure cells are healthy prior to	
	• Adherent cells were dislodged and washed away prior to	induction of apoptosis	
	assaying	• Collect all cells (both attached and dislodged) after	
	• Incorrect incubation times or temperatures	induction for accurate results	
	• Incorrect volumes used	• Refer to datasheet & verify correct incubation times and	
		temperatures	
		• Use calibrated pipettes and aliquot correctly	
Note: The most probable cause is listed under each section. Causes may overlap with other sections			

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Our promise

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