**Gentaur Europe BVBA** Voortstraat 49, 1910 Kampenhout BELGIUM Tel 0032 16 58 90 45 <u>info@gentaur.com</u> For research use only

## CaspGLOW<sup>™</sup> Fluorescein Active Caspase-3 Staining Kit

(Catalog# K183-25, -100; Store kit at -20° C)

#### I. Introduction:

Activation of caspases plays a central role in apoptosis. The CaspGLOW<sup>TM</sup> Fluorescein Active Caspase-3 Staining Kit provides a convenient means for sensitive detection of activated caspase-3 in living cells. The assay utilizes the caspase-3 inhibitor, DEVD-FMK, conjugated to FITC (FITC-DEVD-FMK) as a marker. FITC-DEVD-FMK is cell permeable, nontoxic, and irreversibly binds to activated caspase-3 in apoptotic cells. The FITC label allows for direct detection of activated caspases in apoptotic cells by fluorescence microscopy, flow cytometry, or fluorescence plate reader.

#### II. Kit Contents:

	K183-25	K183-100	
Component	25 assays	100 assays	Part Number
FITC-DEVD-FMK Wash Buffer Z-VAD-FMK	25 μl 50 ml 10 μl	100 μl 2 x 100 ml 10 μl	K183-xx(x)-1 K183-xx(x)-2 K183-xx(x)-3

#### III. Caspase-3 Assay Procedure:

#### A. Staining Procedure:

- Induce apoptosis in cells (1 x 10<sup>6</sup>/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-3 activation.
- 2. Aliquot 300 µl each of the induced and control cultures into eppendorf tubes.
- Add 1 μI of FITC-DEVD-FMK into each tube and incubate for 0.5 1 hour at 37° C incubator with 5 % CO<sub>2</sub>.
- 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  $\mu$ I of Wash buffer. Keep samples on ice. Analyzing samples by flow cytometry using the FL-1 channel.

#### C. Detection by Fluorescence Microscopy:

For fluorescence microscopic analysis, resuspend cells in 100 µl 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.

D. Analysis by Fluorescence Plate Reader: For analysis with fluorescence plate reader, resuspend cells in 100 μl Wash Buffer and then transfer the cell suspension to each well in the black microtiter plate. Measure the fluorescence intensity at Ex/Em = 485/535 nm. For control, use wells containing unlabeled cells.

#### **RELATED PRODUCTS:**

- CaspGLOW Fluorescein Active Caspase Staining Kit (Cat. No. K180-25, 100)
- CaspGLOW Fluorescein Active Caspase-12 Staining Kit (Cat. No. K172-25, 100)
- CaspGLOW Fluorescein Active Caspase-2 Staining Kit (Cat. No. K182-25, 100)
- CaspGLOW Fluorescein Active Caspase-8 Staining Kit (Cat. No. K188-25, 100)
- CaspGLOW Fluorescein Active Caspase-9 Staining Kit (Cat. No. K189-25, 100)
- CaspGLOW Red Active Caspase Staining Kit (Cat. No. K190-25, 100)
- CaspGLOW Red Active Caspase-3 Staining Kit (Cat. No. K193-25, 100)

CaspGLOW Red Active Caspase-8 Staining Kit (Cat. No. K198-25, 100)

CaspGLOW Red Active Caspase-9 Staining Kit (Cat. No. K199-25, 100)

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# **BioVision**



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### GENERAL TROUBLESHOOTING GUIDE FOR CaspGLOW BASED ASSAYS:

Problems	Cause	Solution	
High background	Cell density is higher than recommended	Refer to datasheet 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 datasheet	
	Cells were Incubated for extended period of time	Refer to datasheets for proper incubation time	
	Use of extremely confluent cells	Perform assay when cells are at 70-95% confluency	
	Cells were contaminated	Check for bacteria/ yeast/ mycoplasma contamination	
Lower signal level	Cells did not initiate apoptosis	Determine the optimal time and dose for apoptosis induction (time-course experiment)	
	Very few cells were used for analysis	Refer to data sheet for appropriate cell number	
	Incorrect setting of the equipment or wavelength 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	Old (unhealthy) cells used	Seed healthy cells and make sure cells are healthy prior to induction of apoptosis	
	Adherent cells were dislodged and washed away prior to assaying	Collect all cells (both attached and dislodged) after induction for accurate results	
	Incorrect incubation times or temperatures	Refer to datasheet & verify correct incubation times and temperatures	
	Incorrect volumes used	Use calibrated pipettes and aliquot correctly	
Note: The most probable cause is listed under each section. Causes may overlap with other sections.			