For life science research only. Not for use in diagnostic procedures. FOR IN VITRO USE ONLY.

Annexin-V-FLUOS Staining Kit

Kit for the detection and quantification of apoptosis and differentiation from necrosis at single cell level, based on Annexin-V-labeling

Cat. No.	1 858 777
Cat. No.	1 988 549

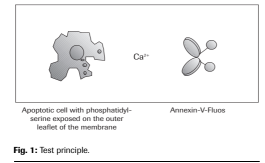
50 tests 250 tests

1. Product overview

Vial/	Label	Content/Cat.No.		Use	
Cap		1 858 777	1 988 549	Use	
1 green	Annexin-V- Fluorescein	110 µl	500 μl	Ready-to-use	
2 red	Propidium iodide	150 μl	500 μl	 Ready-to-use For the preparation of the Annexin-V-Fluores cein labeling solution 	
3 blue	Incubation buffer	50 ml HEPES buffer	4 × 50 ml HEPES buffer	 Ready-to-use For the dilution of the Annexin-V-Fluorescein solution 	
ntrodu		cell surface (1, alterations is th (PS) from the in outer layer, by nal surface of i macrophages surface of lym ptosis (2). The totic cells and	2, 3). One of ne translocati nner part of th which PS bee the cell (4). Fa specifically re obocytes duri recognition a bodies protectilular compoti	osis, changes occur at the these plasma membrane on of phosphatidylserine ne plasma membrane to the comes exposed at the exter adok et al. showed that ecognize PS exposed on the ng the development of apo- curd phagocytosis of apop- cts organisms from the unds leading to inflamma- mice negregie	

Assay principle The analysis of phosphatidylserine on the outer leaflet of apoptotic cell-membranes is performed by using Annexin-V-Fluorescein and Proidium iodid (PI) for the differentiation from necrotic cells or labeling with a cell surface marker for cell characterization. The procedure involves:

Stage	Description
1	Washing the cells in PBS.
2	Incubation of cells with Annexin-V-Fluorescein in a Hepes buffer containing Pl or labeling reagent for cell surfaces (<i>e.g.</i> , CD-marker).
3	Analysis of the samples under a fluorescence microscope or on a flow cytometer.



Application Annexin V is a Ca2+ -dependent phospholipid-binding protein with high affinity for phosphatidylserine (4). This protein can hence be used as a sensitive probe for PS exposure upon the outer leaflet of the cell membrane and is therefore suited to detect apoptotic cells (4, 5, 6, 7) in cell populations but not on tissue sections. Since necrotic cells also expose PS according to the loss of membrane integrity, apoptotic cells have to be differentiated from these necrotic cells. The simultaneous application of a DNA stain which is used for dye exclusion tests allows the discrimination of necrotic cells from the Annexin V positively stained cell cluster. Any other secondary labeling should be possible, *e.g.*, membrane surface staining with a phycoerythrin or TRITC-labeled monoclonal antibody for further cellular characterization (8). Sample material Cell lines Freshly isolated cells Number of tests For 50 tests (Cat. No. 1 858 777) For 250 tests (Cat. No. 1 988 549) Preparation Recombinant Annexin- V is produced in E. coli (strain NB42). The GST-tagged protein is purified by standard purification protocols. Annexin-V-Fluorescein and propidium iodide show the Fluorescence

Version 3, March 2003

488-540 nm

Store at 2-8°C

following fluorescence characteristics: characteristics **Propidium Iodide** Fluorescein 488 nm

	Exolution	400 1111	400 040 1111	
	Emission	518 nm	617 nm	
Specificity	Annexin-V-Fluos binds in a Ca ²⁺ -dependent manner negatively charged phospholipid surfaces and shows high specificity to phosphatidylserine. Therefore, it stains apoptotic as well as necrotic cells. Propidium iodide stains DNA of leaky necrotic cells only.			
Storage/Stability	Stable at 2–8° label.	C until the expira	ation date printed on the	

2. Procedures and required material

Excitation

Additional solutions required	PBS		
Preparation of Annexin-V-FLUOS labeling solution	Predilute 20 μ l Annexin-V-Fluos labeling reagent (vial 1) in 1 ml Incubation buffer (bottle 3) and add 20 μ l Propidium iodide solution (vial 2). Note : 1 ml is enough for 10 samples.		
Staining of cell suspensions	In the following table please find the staining proce- dure for cell suspensions.		
	Step Action		
	1	Wash 10 ⁶ cells with PBS and centrifuge cells at	
		200 × g for 5 min	
	2	Resuspend the cell pellet in 100 μl of Annexin-V- FLUOS labeling solution. Incubate 10–15 min at 15–25°C.	
	3	Analyze by fluorescence microscopy or on a flow cytometer (see 3. Analysis).	

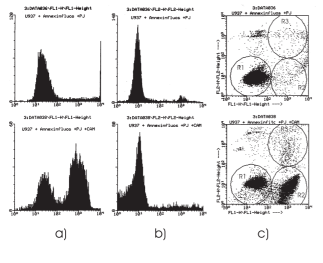


Staining of adherent cells In the following table please find the staining procedure for adherent cells.

Step	Action
1	Remove chambers and silicon borders of cells grown on chamberslides.
2	Remove medium and cover slides with Annexin-V-FLUOS labeling solution (100 μ l/chamber).
3	Put coverslips on slides and incubate for 10–15 min at 15–25°C.
4	Analyze by fluorescence microscopy or on a flow cytometer (see 3. Analysis). Note: We do not recommend to analyze adher- ent cells by flow cytometry, because trypsiniza- tion or scraping for monodispertion of the cells results in false positive staining and analysis of non-dispersed cell clusters.

3. Analysis

Fluorescence- microscopy	For evaluation by fluorescence microscopy use an exci- tation wavelength in the range of 450–500 nm (<i>e.g.</i> , 488 nm) and detection in the range of 515–565 nm (green).		
Flowcytometry	Add 0.5 ml Incubation buffer (bottle 3) per 10 ⁶ cells ar analyze on a flow cytometer using 488 nm excitation and a 515 nm bandpass filter for fluorescein detection and a filter > 600 nm for Pl detection. Electronic com pensation of the instrument is required to exclude ov lapping of the two emission spectra. Typical histogram of apoptotic versus non-apoptotic and necrotic cells are shown in figure 2.		
Figure 2	FACS analysis of apoptotic U937 cells after staining with Annexin-V-FLUOS and propidium iodide. Cultiva- tion for 4 h in the presence (lower row) or absence (upper row) of 4 μ g/ml camptothecin: a) single parameter Annexin-V-FLUOS, b) single parameter propidium iodide and c) dual parameter (FL1 = Annexin-V-FLUOS; FL2 = propidium iodide); Cluster R1 =living cells, R2 = apop- totic cells and R3 = necrotic cells.		



4. References

- Andree, H.A.M. et al. (1990), *J. Biol. Chem.* **265**, 4923. Fadok, V. et al. (1992) *J. Immunology* **148**, 2207. 1
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- Creutz, I.S. (1992) *Science* **258**, 924. Vermes, I. et al. (1995) *J. Immunol. Methods* **184**, 39. Koopman, G. et al (1994) *Blood* **84**, 1415. 5
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- Homburg, C.H.E. et al (1995) *Bloot* **85**, 532. Verhoven, B. et al (1995) *J. Exp. Med.* **182**, 1597. van Engeland, M. et al. (1996) *Cytometrie* **24**, 131. 8

5. Related products

This table only shows a selection of the most important products related to the product described in this pack insert.

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Apoptosis- specific physiologi- cal change	Detection mode/ Product	Pack size	Cat. No.
DNA frag-	Gel Electrophoresis	•	
mentation	Apoptotic DNA-Ladder Kit	20 tests	1 835 246
	In situ assay		
	In Situ Cell Death Detection Kit, TMR red	1 kit (50 tests)	2 156 792
	In Situ Cell Death Detection Kit, Fluorescein	1 kit (50 tests)	1 684 795
	In Situ Cell Death Detection Kit, AP	1 kit (50 tests)	1 684 809
	In Situ Cell Death Detection Kit, POD		1 684 817
	Single reagents for TUNEL and su		
	TUNEL AP	70 tests (3.5 ml)	
	TUNEL POD	70 tests (3.5 ml)	1 772 465
	TUNEL Enzyme	2× 50 μl (20 tests)	1 767 305
	TUNEL Label	3× 550 μl (30 tests)	1 767 291
	TUNEL Dilution Buffer	20 ml	1 966 006
	ELISA		
	Cell Death Detection ELISA	1 kit	1 544 675
	Cell Death Detection ELISAPLUS	1 kit (96 tests)	1 774 425
	Cell Death Detection ELISAPLUS, 10×	1 kit	1 920 685
	Cellular DNA Fragmentation ELISA	1 kit (500 tests)	1 585 045
Cell	Microscopy or FACS	•	
membrane	Annexin-V-Alexa 568	250 tests	1 985 485
alterations	Annexin-V-Biotin	250 tests	1 828 690
	Annexin-V-FLUOS	250 tests	1 828 681
	Annexin V FLUOS Staining Kit	50 tests 250 tests	1 858 777 1 988 549
Enzymatic	Western Blot		
activity	Anti-Poly (ADP-Ribose) Polymerase FIENA	100 µl	1 835 238
	Caspase 3 Activity Assay	1 kit	2 012 952
	Fluorimetric microplate Assay		
	Homogeneous Caspase Assay,	100 tests	3 005 372
	fluorometric	1000 tests	2 236 869
	<i>In situ</i> Assay	•	
	M30 CytoDEATH (formalin grade)	50 tests 250 tests	2 140 322 2 140 349
	M30 CytoDEATH, Fluorescein	250 tests	2 156 857
Expres-	Apoptosis Induction		
sion of	Anti-Fas (CD95/APO-1)	1000 tests	1 922 459
apoptosis-	In situ Assay/Western Blot		
related proteins	Anti-bcl-2 Oncoprotein	1 ml	1 624 989
proteins	Anti-p53-Protein pan (BMG 1B1)	200 µg	1 810 928
]	Anti-p53-POD	50 U	1 810 944
	ELISA	1	L
	p53 pan ELISA	1 kit	1 828 789

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