

Store at 2 – 8 °C

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

# M30 CytoDEATH<sup>™</sup>

# 200 tests (Peviva Prod. No. 10700)

Mouse monoclonal antibody (Clone M30) For the detection of caspase-cleaved keratin 18 neo-epitope M30

# 1. Product Description

 Name:
 M30 CytoDEATH™

 Clone:
 M30

 Isotye:
 IgG2b

 Immunogen:
 Keratin 18 (K18) fragments purified from supernatant from human carcinoma cell line WiDr CCL218.

 Epitope:
 K18 fragment aa284–396

## 1.1 Formulation

Clear solution. 10 µg of M30 CytoDEATH™ monoclonal antibody provided in 200 µl PBS containing 0.1 % BSA, PEG, sucrose and 0.09 % sodium azide.

## **1.2 Specificity**

M30 CytoDEATH<sup>™</sup> antibody is recommended for the detection of the formalin-resistant M30 neo-epitope on human, monkey and bovine caspase-cleaved keratin 18 (K18) cytoskeletal protein. M30 CytoDEATH<sup>™</sup> does not recognise intact K18.

*Note:* The immunoreactivity of the M30 CytoDEATH<sup>™</sup> antibody is confined to the cytoplasma of apoptotic cells. Nonspecific cross-reactivity with nuclear antigens of highly proliferating cells within tissue sections may occasionally occur at high M30 CytoDEATH<sup>™</sup> concentrations.

#### **1.3 Recommended applications**

- Western blot (WB)
- Immunocytochemistry (ICC)
- Flow cytometry (FACS)
- Immunohistochemistry (IHC) including formalin-fixed and paraffin-embedded tissue sections (PS) and cryostat sections (FS)

#### 1.4 Working solution

M30 CytoDEATH<sup>™</sup> is provided in a convenient ready-to-use stock solution. Use a dilution 1:100 in Incubation Buffer (final concentration 0.5 µg/ml).

#### 1.5 Storage and stability

The M30 CytoDEATH<sup>™</sup> antibody is provided in a ready-to-use format and is stable at 2 − 8 °C through the expiration date printed on the label.

Alternatively, it can be stored in aliquots at -20 °C.

The antibody is shipped at ambient temperature.

*Note:* Repeated freezing and thawing should be avoided.

## 1.6 Quality control

The M30 CytoDEATH<sup>™</sup> antibody is function tested using a celluar model: HeLa cells treated with recombinant TRAIL and CHX analysed by immunocytochemistry and flow cytometry.

## M30 CytoDEATH<sup>™</sup> – Key advantages

Benefits	Features
Early and specific detection of apoptosis	Detects caspase-cleaved keratin 18; caspase activity is one of the earli- est and most common markers for apoptosis.
Sustained signal from cells at early to later stages of apoptosis	In contrast to measuring i. e. active caspase-3, where the signal is defined to a limited time window and de- creases at late stages of apoptosis, the caspase-generated K18 neo-epitope can still be detected even after apo- topic cells have disintegrated.
Superior sensitivity	The keratin 18 neo-epitope is an accu- mulating substrate generated by few activated caspase molecules.
Assay is independent of the activation of a single caspase	Keratin 18 is cleaved <i>in vivo</i> by several effector caspases, including caspase-3, 6, 7 and 9.
Clear results	Apoptotic cells are clearly distinguish- able from viable cells or necrotic cells.
Apoptosis specific No false positive results in circumstances of DNA damage	Whereas TUNEL analysis can give rise to false positive results under condi- tions when DNA double-strand breaks occur, detection of the K18 neo- epitope using the M30 CytoDEATH™ antibody shows superior specificity for apoptotic cells compared to TUNEL.
Specificity for epithelial (i.e. carcinoma or liver tissue) apoptosis	Expression pattern of K18 is restricted to cells of epithelial origin. Lymphoid and neuronal cell apoptosis is not de- tected by M30 CytoDEATH™ antibody.
Application versatility	The M30 CytoDEATH <sup>™</sup> antibody has been successfully used in Western blot, immunocytochemistry, flow cy- tometry and immunohistochemistry, including frozen and formalin-fixed, paraffin-embedded tissue sections.
Recommended for formalin- fixed paraffin-embedded tissue	Recommended for routinely fixed tissue samples. Retrospective studies are possible.
Convenience	Easy to use standard protocol. M30 CytoDEATH™ antibody provided in a ready-to-use format.

# 2. Background Information

#### 2.1 Caspase substrate K18 and apoptosis in epithelial cells

Apoptosis induced by either death-inducing receptors or other stimuli leads to activation of specific caspases [1, 2]. Subsequently, apoptotic cells are eliminated by an intrinsic suicide program, resulting in DNA fragmentation, nuclear condensation, cytoskeletal reorganisation, plasma membrane blebbing and loss of cell adhesion.

Keratin 18 (K18) is a type I intermediate filament protein and the major component of single-layer and glandular epithelial cells. It is expressed in most types of carcinomas such as lung, liver, prostate, breast and colon, and abundantly present in liver cells, whereas K18 is absent in lymphoid and neuronal cells and tissues. During apoptosis after initiation of effector caspases 3, 6, 7 and 9, K18 is cleaved into proteolytic fragments liberating neo-epitopes (NE) at the cleavage sites [3-6].

## 2.2 M30 CytoDEATH<sup>™</sup> antibody for the specific detection of apoptosis

K18 is cleaved by capases, liberating a neo-epitope (M30) that is specifically recognised by the M30 CytoDEATH™ monoclonal antibody. Specific proteolytic cleavage of K18 is an event taking place before disruption of membrane asymmetry and induction of DNA strand breaks. Numerous studies confirm that M30 CytoDEATH™ antibody detects only apoptotic but not viable or necrotic cells. Reactivity of M30 CytoDEATH™ antibody in immunohistochemistry correlates to apoptosis measured by TUNEL and shows superior reliability in conditions when DNA double-strand breaks occur independent of apoptosis [7].

The capacity of M30 CytoDEATH<sup>™</sup> antibody in flow cytomery and immunohistochemistry studies to distinguish between necrotic and apoptotic epithelial cells has been verified in several disease entities. Consequently, M30 CytoDEATH<sup>™</sup> antibody represents a unique tool for easy and reliable determination of apoptosis from very early until well advanced stages in single cells and tissue sections of epithelial origin [8].

Moreover, there are two M30 CytoDEATH<sup>™</sup> antibody-based ELISAs available:

- M30 CytoDeath<sup>™</sup> ELISA (PEVIVA Prod. No.: 10900) is suggested to serve as a high-throughput assay for functional screening and in vitro characterisation of effective pro-apoptotic drugs using cell culture supernatants, and spheroid or tissue lysates.
- The CE-marked M30 Apoptosense® ELISA (PEVIVA Prod. No: 10010) has been successfully used to determine elevated K18 neo-epitope levels in blood samples from patients as a useful biomarker to monitor response to treatment or disease staging [9–12].

# **3. Procedures and Materials Required**

## 3.1 Procedure for immunofluorescence and flow cytometry

### 3.1.1 Introduction

The following procedure describes the detection of apoptosis with M30 CytoDEATH™ antibody in immunofluorescence and flow cytometry.

Please note: a fluorescein-conjugated M30 CytoDEATH™ antibody is available from PEVIVA (Prod. No: 10800). The M30 CytoDEATH™ Fluorescein antibody is recommended for immunofluorescence and flow cytometry applications as it does not require an anti-mouse IgG-fluorescein secondary antibody. If using other detection methods or sample material, the conditions may vary and have to be adapted.

Additional reagents required

- PBS, Methanol and BSA
- Secondary detection reagents such as anti-mouse IgG-fluorescein (i.e. from DAKO)

#### Preparation of working solutions

Incubation Buffer: PBS containing 1 % BSA Washing Buffer: PBS

## Preparation of M30 CytoDEATH<sup>™</sup> antibody working solution

Dilute the M30 CytoDEATH<sup>™</sup> antibody stock solution 1:100 in Incubation Buffer (final concentration 0.5  $\mu g/ml).$ 

Note: The antibody solutions should be free of precipitate. If necessary, centrifuge the solution at high speed prior to use.

## 3.1.2 Immunofluorescence and flow cytometry protocol

Step	Action
1	Wash cells in PBS.
2	Fix cells in ice-cold pure methanol at -20 °C for 30 min.
3	Wash cells with Washing Buffer twice.
4	Remove Washing Buffer.
5	Incubate with 100 µl M30 CytoDEATH™ antibody for 30 min at 15–25 °C.
6	Wash cells with Washing Buffer twice.
7	Incubate with 100 $\mu$ l fluorescein-conjugated anti-mouse lgG (1 – 2 $\mu$ g/ml) antibody for 30 min at 15 – 25 °C in the dark.
8	Wash cells with Washing Buffer twice.
9	Examine the cells on a slide under the fluorescence microscope, or, dilute cells in 0.5 ml PBS and store samples in the dark until analysis by flow cytometry.

#### 3.2 Procedure for immunohistochemistry

## 3.2.1 Introduction

The following procedure describes the detection of apoptosis with M30 CytoDEATH<sup>™</sup> antibody in a three step method in immunohistochemistry (paraffin-embedded tissue) for maximal sensitivity

Please note: A biotin-conjugated M30 CytoDEATH™ antibody is available from PEVIVA (Prod. No: 10750), which can be used in combination with sensitive amplification reagents. If using other detection methods or sample material, the conditions may vary and have to be adapted.

3.2.2 Recommended reagents

For preparation of samples:

- Xvlol
- Ethanol 96 %
- Ethanol 70 %
- Methanol/H<sub>2</sub>O<sub>2</sub> (3 %) Citric acid
- NaOH, 1 M
- Hematoxylin (i.e. from Merck)
- Mounting medium (i.e. Kaiser's glycerine gelatine from Merck)

For the immunohistochemistry procedure:

- Anti-mouse-IgG biotin (i.e. from DAKO)
- Streptavidin-POD (i.e. from DAKO)
- DAB or AEC substrate (i.e. from Zymed)
- PBS •
- BSA

Preparation of working solutions

The following table lists the working solutions needed to perform the immunohistochemistry staining procedure.

Working Solution	Composition	Stability/ storage	Use
Washing Buffer	PBS	4 weeks at 2 – 8 °C	Washing step
Incubation Buffer	PBS containing 1 % BSA	4 weeks at 2–8 °C	Preperation of antibody work- ing solution
Citric Acid Buffer (0.01 M)	2 g/l citric acid, pH 6.0 adjusted with 1 M NaOH	4 weeks at 2–8 °C	Antigen retrival

Preparation of M30 CytoDEATH<sup>™</sup> antibody working solution

Dilute the M30 CytoDEATHTM antibody stock solution 1:100 in Incubation Buffer (final concentration 0.5  $\mu g/m$ ).

*Note:* The antibody solutions should be free of precipitate. If necessary, centrifuge the solution at high speed prior to use.

Preparation of sample material

Before starting the immunohistochemical protocol, dewax paraffin-embedded tissue sections as described in the following table.

Step	Action
1	Place paraffin-embedded sections into an incubator at 37 $^{\circ}\mathrm{C}$ over night to air-dry.
2	To dewax formalin-fixed, paraffin-embedded tissue sections, process the sections as follows:
	<ul> <li>2 coplin jars of xylol (2 – 5 min),</li> <li>2 coplin jars of ethanol (96 %)</li> <li>1 coplin jar of ethanol (70 %)</li> <li>1 coplin jar of methanol/H<sub>2</sub>O<sub>2</sub> (3 %) for 10 min at 15 – 25 °C.</li> </ul>
3	Rinse 10 min in PBS.

Note: The sections should not be allowed to dry during this procedure.

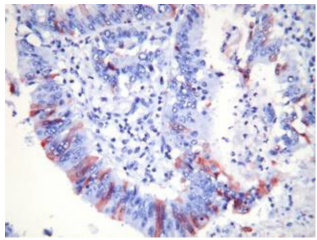
3.2.3 Immunohistochemistry protocol

*Note:* For optimal results it is highly recommended to follow the below mentioned method for antigen retrieval.

Step	Action	
1	<ul> <li>Prewarm citric acid buffer (0.01 M, pH 6.0) by incubation in a microwave oven at 750 W until the solution boils.</li> <li>When the solution is boiling, turn the setting of the microwave oven to "keep warm" (about 100 W).</li> <li>Place tissue section slides in a slide rack and put them into the heated citric acid solution (approx. 90 °C).</li> <li>Incubate at this setting for 20 min.</li> <li>Note: For optimal morphology it is recommended to keep the</li> </ul>	
	solution shortly below the boiling point to avoid gas formation under the sections.	
2	Rinse $3 \times$ in PBS and incubate 10 min in a separate jar of PBS to cool down.	
3	Remove Incubation Buffer and add 100 μl M30 CytoDEATH™ antibody working solution. Incubate for 30 min at 15 – 25 °C in a humid chamber.	
4	Wash slides in Washing Buffer (use 3 separate jars and dip 3 $\times$ into each jar).	
5	Cover the section with 100 $\mu$ l of anti-mouse-lgG biotin according to your established optimised procedure for the reagent from your selected supplier or use DAKO reagent at 1:400 dilution.	
	Incubate for 30 min at 37 °C in a humid chamber.	
6	Wash slides in Washing Buffer (use 3 separate jars and dip 3 $\times$ into each jar).	
7	Cover the section with 100 µl of streptavidin-POD according to your established optimised procedure for the reagent from your selected supplier, or use DAKO reagent at 1:600 dilution.	
	Incubate for 30 min at 15 – 25 °C in a humid chamber.	
8	Wash slides in Washing Buffer (use 3 separate jars and dip 3 $\times$ into each jar).	
9	Incubate slides in a freshly prepared substrate solution (i. e. AEC) at $15-25$ °C until a clearly visible colour develops (1-5 min). A negative control should not show any develop- ment of colour during the incubation period.	
10	Stop the reaction by extensive rinsing in double distilled water.	
11	Subsequently, counterstain the preparation with hematoxylin and mount the section (i. e. Kaiser's glycerine gelatine when using AEC).	

# 4. Results

Immunohistochemistry (paraffin-embedded)



**Figure:** Detection of apoptosis in a formalin-fixed and paraffin-embedded tissue section from a human colon cancer showing confined cytoplasmic staining for K18-Asp396-NE (caspase-cleaved K18) using M30 CytoDEATH<sup>TM</sup>. Secondary detection with anti-mouse IgG-biotin, streptavidin-POD and AEC as substrate, counterstained with hematoxylin.

# 5. Appendix

K18 positive cell lines and tissues successfully analyzed with the M30 CytoDEATH  $^{\rm m}$  antibody:

## Human epithelial cell lines:

Breast cancer: MDA-MB-231, MCF-7, HBL100 Colon cancer: WiDr, HCT 116, HT29, SW620 Cervical cancer: HeLa Kidney cancer: ACHN, A498 Head & neck cancer: SCC9, SCC25, FaDu Prostate cancer: PC-3, LNCaP, DU 145 Bladder cancer: RT4, J82

### Human epithelial tissues:

Breast, lung, liver, colon, pancreas, intestine, kidney, salivary gland, trophoblast, endometrium, bladder, oral epithelium.

## **6. PEVIVA Products**

M65 EpiDeath® ELISA

M30 CytoDeath™ ELISA

M30 CytoDEATH<sup>™</sup> antibodies M30 CytoDEATH™ Prod. No. 10700 M30 CytoDEATH™ Biotin M30 CytoDEATH™ Fluorescein Prod. No. 10750 Prod. No. 10800 M30 CytoDEATH<sup>™</sup> Orange Prod. No. 10830 Keratin 18 antibodies M5 Keratin 18 mAb Prod. No. 10600 M6 Keratin 18 mAb Prod. No. 10650 **ELISA kits** M30 Apoptosense® ELISA M65° ĖLISA

Prod. No. 10011 Prod. No. 10020 Prod. No. 10040 Prod. No. 10900

# 7. References

1. Creagh EM, Conroy H, Martin SJ. (2003) Caspase-activation pathways in apoptosis and immunity. Immunol Rev. 193:10-21

2. Creagh EM, Martin SJ. (2001) Caspases: cellular demolition experts. Biochem Soc Trans. 29: 696-702.

3. Caulin C, Salvesen GS, Oshima G. (1997) Caspase Cleavage of keratin 18 and reorganization of intermediate filaments during epithelial cell Apoptosis. J Cell Biol. 138: 1379-1394.

4. Leers MP, Kölgen W, Björklund V, Bergman T, Tribbick G, Persson B, Björklund P, Ramaekers FCS, Björklund B, Nap M, Jörnvall H, Schutte B. (1999) Immunocytochemical detection and mapping of a Cytokeratin 18 neoepitope exposed during early apoptosis. J Pathol. 187: 567-572.

5. Schutte B, Henfling M, Kolgen W, Bouman M, Meex S, Leers MP, Nap M, Bjorklund V, Bjorklund P, Bjorklund B, Lane EB, Omary MB, Jornvall H, Ramaekers FCS. (2004) Keratin 8/18 breakdown and reorganization during apoptosis. Exp Cell Res. 297:11-26.

6. Dinsdale D, Lee JC, Dewson G, Cohen GM, Peter ME. (2004) Intermediate filaments control the intracellular distribution of caspases during apoptosis. Am J Pathol. 164: 395-407.

7. Walker JA and Quirke P. (2001) Viewing apoptosis through a "TUNEL". J Pathol. 95: 275-276.

8. Grassi A, Susca M, Ferri S, Gabusi E, D'Errico A, Farina G, Maccariello S, Zauli D, Bianchi FB, Ballardini G. (2004) Detection of the M30 neoepitope as a new tool to quantify liver apoptosis: timing and patterns of positivity on frozen and paraffin-embedded sections. Am J Clin Pathol. 121: 211-219.

9. Bivén K, Erdal H, Hägg M, Ueno T, Zhou R, Lynch M, Rowley B, Wood J, Zhang C, Toi M, Shoshan MC, Linder S. (2003). A novel assay for discovery and characterization of pro-apoptotic drugs and for monitoring apoptosis in patient sera. Apoptosis 8: 263-268.

10. Ueno T, Toi M, Bivén K, Bando H, Ogawa T, Linder S. (2003) Measurement of an apoptosis product in the sera of breast cancer patients. Eur J Cancer 39: 769-774.

11. Kramer G, Erdal H, Mertens H, Nap M, Mauermann J, Steiner G, Marberger M, Bivén K, Shoshan MC, Linder S. (2004) Differentiation between cell death modes using measurements of different soluble forms of extracellular cytokeratin 18. Cancer Res. 64: 751-1756.

12. Linder S, Havelka AM, Ueno T, Shoshan MC. (2004) Determining tumor apoptosis and necrosis in patient serum using cytokeratin 18 as a biomarker. Cancer Lett. 214: 1-9.

# 8. M30 CytoDEATH<sup>™</sup> Application References

## Immunocytochemistry (ICC)

MacFarlane M, Merrison W, Dinsdale D, Cohen GM. (2000) Active caspases and cleaved cytokeratins are sequestered into cytoplasmic inclusions in TRAIL-induced apoptosis. J Cell Biol. 148: 1239-1254.

Barrett KL, Willingham JM, Garvin AJ, Willingham MC. (2001) Advances in cyto-chemical methods for detection of apoptosis. J Histochem Cytochem 49: 821-32.

Lee JC, Schickling O, Stegh AH, Oshima RG, Dinsdale D, Cohen GM, Peter ME. (2002) DEDD regulates degradation of intermediate filaments during apoptosis. J Cell Biology 158: 1051-1066.

Dinsdale D, Lee JC, Dewson G, Cohen GM, Peter ME. (2004) Intermediate filaments control the intracellular distribution of caspases during apoptosis. Am J Pathol. 164: 395-407.

Schutte B, Henfling M, Kolgen W, Bouman M, Meex S, Leers MP, Nap M, Bjorklund V, Bjorklund P, Bjorklund B, Lane EB, Omary MB, Jornvall H, Ramaekers FC. (2004) Keratin 8/18 breakdown and reorganization during apoptosis. Exp Cell Res. 297:11-26.

## Flow cytometry (FACS)

RupaJD,DeBruineAP,GerbersAJ,LeersMP,NapM,KesselsAG,SchutteB,Arends JW.(2003)Simultaneousdetectionofapoptosisandproliferationincolourectal carcinoma by multiparameter flow cytometry allows separation of high and low-turnover tumors with distinct clinical outcome. Cancer 97: 2404-2411.

Morsi HM, Leers MP, Radespiel-Troger M, Björklund V, Kabarity HE, Nap M, Jäger W. (2000) Apoptosis, bcl-2 expression, and proliferation in benign and malignant endometrial epithelium: An approach using multiparameter flow cytometry. Gynecol. Oncol. 77, 11-7.

#### Immunohistochemistry (IHC)

Morsi HM, Leers MP, Jäger W, Björklund V, Radespiel-Troger M, el Kabarity H, Nap M and Lang N. (2000) The patterns of expression of an apoptosis-related CK18 neoepitope, the bcl-2 proto-oncogene, and the Ki67 proliferation marker in normal, hyperplastic, and malignant endometrium. Int J. Gynecol. Pathol. 19: 118-126.

Bantel H, Ruck P, Gregor M, Schulze-Osthoff K. (2001) Detection of elevated caspase activation and early apoptosis in liver diseases. Eur J Cell Biol. 80: 230-239

Fukuda M, Tanaka A, Kitada M, Fukuda F, Suzuki S, Jiang Y, Kebusa Y, Ohno J, Yamamoto Y, Minato H, Nonomura A, Sakashita H, Kusama K. (2001) Immunohistochemical detection of cytokeratin 18 and its neo-epitope in Warthin's tumor (adenolymphoma) of the parotid glands. Anticancer Res 21:109-12.

Walker JA and Quirke P. (2001) Viewing apoptosis through a "TUNEL". J. Pathol 195: 275-276

Leers MP, Björklund V, Björklund B, Jörnvall H, Nap M. (2002) An immunohistochemical study of the clearance of apoptotic cellular fragments. Cell Mol Life Sci. 59: 1358-1365.

Grassi A, Susca M, Ferri S, Gabusi E, D'Errico A, Farina G, Maccariello S, Zauli D, Bianchi FB, Ballardini G. (2004) Detection of the M30 neoepitope as a new tool to quantify liver apoptosis: timing and patterns of positivity on frozen and paraffin-embedded sections. Am J Clin Pathol. 121: 211-219.

Koornstra JJ, Rijcken FE, De Jong S, Hollema H, De Vries E, Kleibeuker JH. (2004) Assessment of apoptosis by M30 immunoreactivity and the correlation with morphological criteria in normal colourectal mucosa, adenomas and carcinomas. Histopathol. 44: 9-17.

Takada M, Kataoka A, Toi M, Bando H, Toyama K, Horiguchi S, Ueno T, Linder S, Saji S, Hayashi Y, Funata N, Kinoshita J, Murakami S, Ohono S. (2004) A close association between alteration in growth kinetics by neoadjuvant chemotherapy and survival outcome in primary breast cancer. Int J Oncol. 25: 397-405.

Simopoulos C, Tsaroucha AK, Asimakopoulos B, Giatromanolaki A, Gavriilidis P, Polychronidis A, Karayiannakis A. (2008) Total and caspase-cleaved cytokeratin 18 in chronic cholecystitis: a prospective study. BMC Gasteroenterol. 8:14–18.

For further, up-to-date information and to order, please visit www.peviva.com.



VIVALAVIDA AB, Hästholmsvägen 32, 131 30 Nacka, Sweden Website: www.vlvbio.com • E-mail: info@vlvbio.com Phone: +46 8 122 053 00