

Caspase 7 (Cleaved-Asp198) Antibody
Purified Rabbit Polyclonal Antibody (Pab)
Catalog # AP50700

Specification

Caspase 7 (Cleaved-Asp198) Antibody - Product Information

Application	WB, IHC
Primary Accession	P55210
Reactivity	Human, Mouse, Rat
Host	Rabbit
Clonality	Polyclonal
Calculated MW	32 34 28 38 KDa
Antigen Region	185-209

Caspase 7 (Cleaved-Asp198) Antibody - Additional Information

Gene ID 840

Other Names

Caspase-7, CASP-7, Apoptotic protease Mch-3, CMH-1, ICE-like apoptotic protease 3, ICE-LAP3, Caspase-7 subunit p20, Caspase-7 subunit p11, CASP7, MCH3

Dilution

WB~~1:1000
IHC~~1:50-100

Format

Rabbit IgG in phosphate buffered saline (without Mg²⁺ and Ca²⁺), pH 7.4, 150mM NaCl, 0.09% (W/V) sodium azide and 50% glycerol.

Storage Conditions

-20°C

Caspase 7 (Cleaved-Asp198) Antibody - Protein Information

Name CASP7 {ECO:0000303|PubMed:9070923, ECO:0000312|HGNC:HGNC:1508}

Function

Thiol protease involved in different programmed cell death processes, such as apoptosis, pyroptosis or granzyme-mediated programmed cell death, by proteolytically cleaving target proteins (PubMed:11257230, PubMed:11257231, PubMed:11701129, PubMed:15314233, PubMed:16916640, PubMed:17646170, PubMed:18723680, PubMed:<a href="http://www.uniprot.org/citations/19581639"

target="_blank">>19581639, PubMed:>8521391, PubMed:>8567622, PubMed:>8576161, PubMed:>9070923). Has a marked preference for Asp-Glu-Val-Asp (DEVD) consensus sequences, with some plasticity for alternate non-canonical sequences (PubMed:>12824163, PubMed:>15314233, PubMed:>17697120, PubMed:>19581639, PubMed:>20566630, PubMed:>23650375, PubMed:>23897474, PubMed:>27032039). Its involvement in the different programmed cell death processes is probably determined by upstream proteases that activate CASP7 (By similarity). Acts as an effector caspase involved in the execution phase of apoptosis: following cleavage and activation by initiator caspases (CASP8, CASP9 and/or CASP10), mediates execution of apoptosis by catalyzing cleavage of proteins, such as CLSPN, PARP1, PTGES3 and YY1 (PubMed:>10497198, PubMed:>16123041, PubMed:>16374543, PubMed:>16916640, PubMed:>18723680, PubMed:>20566630, PubMed:>21555521, PubMed:>22184066, PubMed:>22451931, PubMed:>27889207, PubMed:>28863261, PubMed:>31586028, PubMed:>34156061, PubMed:>35338844, PubMed:>35446120). Compared to CASP3, acts as a minor executioner caspase and cleaves a limited set of target proteins (PubMed:>18723680). Acts as a key regulator of the inflammatory response in response to bacterial infection by catalyzing cleavage and activation of the sphingomyelin phosphodiesterase SMPD1 in the extracellular milieu, thereby promoting membrane repair (PubMed:>21157428). Regulates pyroptosis in intestinal epithelial cells: cleaved and activated by CASP1 in response to S.typhimurium infection, promoting its secretion to the extracellular milieu, where it catalyzes activation of SMPD1, generating ceramides that repair membranes and counteract the action of gasdermin-D (GSDMD) pores (By similarity). Regulates granzyme-mediated programmed cell death in hepatocytes: cleaved and activated by granzyme B (GZMB) in response to bacterial infection, promoting its secretion to the extracellular milieu, where it catalyzes activation of SMPD1, generating ceramides that repair membranes and counteract the action of perforin (PRF1) pores (By similarity). Following cleavage by CASP1 in response to inflamasome activation, catalyzes processing and inactivation of PARP1, alleviating the transcription repressor activity of PARP1 (PubMed:>22464733). Acts as an inhibitor of type I interferon production during virus-induced apoptosis by mediating cleavage of antiviral proteins CGAS, IRF3 and MAVS, thereby preventing cytokine overproduction (By similarity). Cleaves and activates sterol regulatory element binding proteins (SREBPs) (PubMed:>8643593). Cleaves phospholipid scramblase proteins XKR4, XKR8 and XKR9 (By similarity). In case of infection, catalyzes cleavage of Kaposi sarcoma-associated herpesvirus protein ORF57, thereby preventing expression of viral lytic genes (PubMed:>20159985).

target="_blank">20159985). Cleaves BIRC6 following inhibition of BIRC6-caspase binding by DIABLO/SMAC (PubMed:36758104, PubMed:36758106).

Cellular Location

Cytoplasm, cytosol. Nucleus. Secreted, extracellular space {ECO:0000250|UniProtKB:P97864}. Note=Following cleavage and activation by CASP1 or granzyme B (GZMB), secreted into the extracellular milieu by passing through the gasdermin-D (GSDMD) pores or perforin (PRF1) pore, respectively {ECO:0000250|UniProtKB:P97864}

Tissue Location

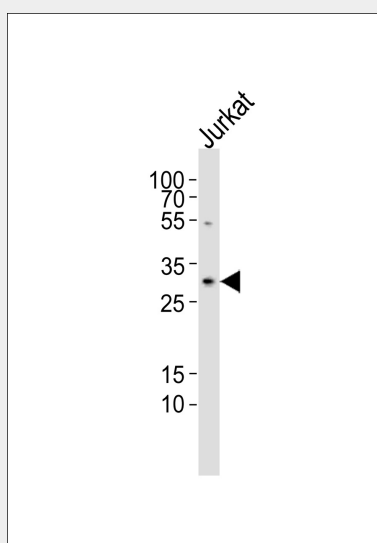
Highly expressed in lung, skeletal muscle, liver, kidney, spleen and heart, and moderately in testis. No expression in the brain.

Caspase 7 (Cleaved-Asp198) Antibody - Protocols

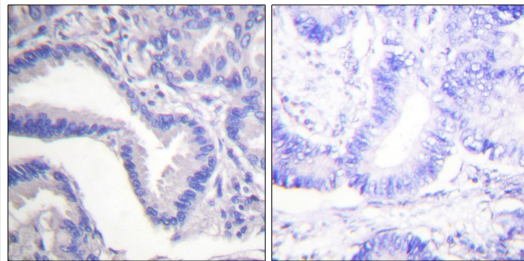
Provided below are standard protocols that you may find useful for product applications.

- [Western Blot](#)
- [Blocking Peptides](#)
- [Dot Blot](#)
- [Immunohistochemistry](#)
- [Immunofluorescence](#)
- [Immunoprecipitation](#)
- [Flow Cytometry](#)
- [Cell Culture](#)

Caspase 7 (Cleaved-Asp198) Antibody - Images



Western blot analysis of lysate from Jurkat cell line, using Caspase 7 (Cleaved-Asp198) Antibody(AP50700). AP50700 was diluted at 1:1000. A goat anti-rabbit IgG H&L(HRP) at 1:5000 dilution was used as the secondary antibody. Lysate at 35ug.



Immunohistochemical analysis of paraffin-embedded human lung carcinoma tissue using Caspase 7 (Cleaved-Asp198) Antibody .

Caspase 7 (Cleaved-Asp198) Antibody - Background

Involved in the activation cascade of caspases responsible for apoptosis execution. Cleaves and activates sterol regulatory element binding proteins (SREBPs). Proteolytically cleaves poly(ADP-ribose) polymerase (PARP) at a '216-Asp-| -Gly- 217' bond. Overexpression promotes programmed cell death.

Caspase 7 (Cleaved-Asp198) Antibody - References

- Fernandes-Alnemri T.,et al.Cancer Res. 55:6045-6052(1995).
Duan H.,et al.J. Biol. Chem. 271:1621-1625(1996).
Lippke J.A.,et al.J. Biol. Chem. 271:1825-1828(1996).
Juan T.S.-C.,et al.Genomics 40:86-93(1997).
Kalnine N.,et al.Submitted (MAY-2003) to the EMBL/GenBank/DDBJ databases.