

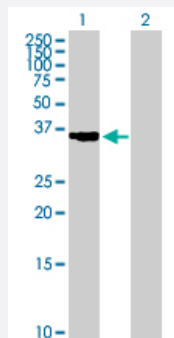
MaxPab®

CTSK purified MaxPab mouse polyclonal antibody (B01P)

Catalog # H00001513-B01P

Size 50 ug

Applications



Western Blot (Transfected lysate)

Western Blot analysis of CTSK expression in transfected 293T cell line ([H00001513-T01](#)) by CTSK MaxPab polyclonal antibody.

Lane 1: CTSK transfected lysate(36.19 KDa).

Lane 2: Non-transfected lysate.

Specification

Product Description	Mouse polyclonal antibody raised against a full-length human CTSK protein.
Immunogen	CTSK (NP_000387.1, 1 a.a. ~ 329 a.a) full-length human protein.
Sequence	MWGLKVL LLPVVSFALYP E EILDTHWELWKKTHR KQYNNKVDEISRRLIWEKNLKYISIHNL EASLG VHTYELAMNHLGDMTSEE VVQKMTGLKVPLSHSR SNTLYIPEWEGRAPDSVDYRKKGYVTPVK NQQQCGSCWAFSSVGALE GQLKKKTGKLLNLSPQNLVDCVSENDGCGGGYMTNAFQYVQKNR GIDSEDAYPYVGQEESCMYNPTGKAAKCRGYREIPEGNEKALKRAVARVGPVSVAIDASLTSFQF YSKGVYDESCNSDNLNHAVLAVGYGIQKGNKHWI IKNWGENWGNKGYILMARNKNNACGIANL ASFPKM
Host	Mouse
Reactivity	Human
Interspecies Antigen Sequence	Mouse (86); Rat (88)
Quality Control Testing	Antibody reactive against mammalian transfected lysate.
Storage Buffer	In 1x PBS, pH 7.4

Storage Instruction

Store at -20°C or lower. Aliquot to avoid repeated freezing and thawing.

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[Protocol Download](#)

Gene Info — CTSK

Entrez GeneID [1513](#)

GeneBank Accession# [NM_000396.2](#)

Protein Accession# [NP_000387.1](#)

Gene Name CTSK

Gene Alias CTS02, CTSO, CTSO1, CTSO2, MGC23107, PKND, PYCD

Gene Description cathepsin K

Omim ID [265800 601105](#)

Gene Ontology [Hyperlink](#)

Gene Summary The protein encoded by this gene is a lysosomal cysteine proteinase involved in bone remodeling and resorption. This protein, which is a member of the peptidase C1 protein family, is predominantly expressed in osteoclasts. However, the encoded protein is also expressed in a significant fraction of human breast cancers, where it could contribute to tumor invasiveness. Mutations in this gene are the cause of pycnodysostosis, an autosomal recessive disease characterized by osteosclerosis and short stature. This gene may be subject to RNA editing. [provided by RefSeq]

Other Designations OTTHUMP00000032938|cathepsin O1|cathepsin O2|cathepsin X

Pathway

- [Lysosome](#)

Disease

- [Alzheimer disease](#)
- [Body Weight](#)
- [Cardiovascular Diseases](#)
- [Diabetes Complications](#)
- [Diabetes Mellitus](#)
- [Edema](#)
- [Genetic Predisposition to Disease](#)
- [Metabolic Syndrome X](#)
- [Neoplasms](#)
- [Osteoporosis](#)
- [Ovarian Neoplasms](#)