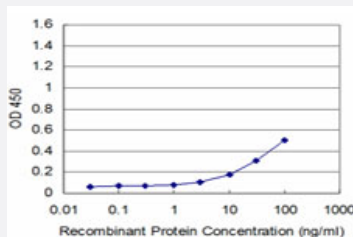


ROR2 monoclonal antibody (M04), clone 4B8

Catalog # H00004920-M04

Size 100 ug

Applications



Sandwich ELISA (Recombinant protein)

Detection limit for recombinant GST tagged ROR2 is approximately 3ng/ml as a capture antibody.

Specification

Product Description	Mouse monoclonal antibody raised against a partial recombinant ROR2.
Immunogen	ROR2 (NP_004551, 34 a.a. ~ 143 a.a) partial recombinant protein with GST tag. MW of the GST tag alone is 26 KDa.
Sequence	EVEVLDPN DPLG PLDGQD GPI TLKGYFLNFLEPVNNITVQGQTALHCKVAGNPPPNVRWLKND APVVQEPRRIIRKTEYGSRLRIQDLDTTDTGYQCVATNGMKT
Host	Mouse
Reactivity	Human
Interspecies Antigen Sequence	Mouse (88); Rat (89)
Isotype	IgG2b Kappa
Quality Control Testing	Antibody Reactive Against Recombinant Protein.
Storage Buffer	In 1x PBS, pH 7.4
Storage Instruction	Store at -20°C or lower. Aliquot to avoid repeated freezing and thawing.

Applications

- Sandwich ELISA (Recombinant protein)

Detection limit for recombinant GST tagged ROR2 is approximately 3ng/ml as a capture antibody.

[Protocol Download](#)

- ELISA

Gene Info — ROR2

Entrez GeneID [4920](#)

GeneBank Accession# [NM_004560](#)

Protein Accession# [NP_004551](#)

Gene Name ROR2

Gene Alias BDB, BDB1, MGC163394, NTRKR2

Gene Description receptor tyrosine kinase-like orphan receptor 2

Omim ID [113000 268310 602337](#)

Gene Ontology [Hyperlink](#)

Gene Summary

The protein encoded by this gene is a receptor protein tyrosine kinase and type I transmembrane protein that belongs to the ROR subfamily of cell surface receptors. The protein may be involved in the early formation of the chondrocytes and may be required for cartilage and growth plate development. Mutations in this gene can cause brachydactyly type B, a skeletal disorder characterized by hypoplasia/aplasia of distal phalanges and nails. In addition, mutations in this gene can cause the autosomal recessive form of Robinow syndrome, which is characterized by skeletal dysplasia with generalized limb bone shortening, segmental defects of the spine, brachydactyly, and a dysmorphic facial appearance. [provided by RefSeq]

Other Designations

OTTHUMP00000021634|OTTHUMP00000063680|neurotrophic tyrosine kinase receptor-related 2|tyrosine-protein kinase transmembrane receptor ROR2

Disease

- [Cerebral Hemorrhage](#)
- [Chronic Disease](#)

- [Cleft Lip](#)
- [Cleft Palate](#)
- [Genetic Predisposition to Disease](#)
- [Hypertension](#)
- [Intracranial Hemorrhages](#)
- [Kidney Diseases](#)
- [Stroke](#)
- [Subarachnoid Hemorrhage](#)