# antibodies -online.com





## anti-PEX5 antibody (AA 51-150) (HRP)



Go to Product page

( )	11/	IN	/ie	A .
	/ // <del> </del>	۱ ات	/   (−	' \/\/

Quantity:	100 μL	
Target:	PEX5	
Binding Specificity:	AA 51-150	
Reactivity:	Mouse	
Host:	Rabbit	
Clonality:	Polyclonal	
Conjugate:	This PEX5 antibody is conjugated to HRP	
Application:	ELISA, Immunohistochemistry (Paraffin-embedded Sections) (IHC (p)), Immunohistochemistry (Frozen Sections) (IHC (fro))	

### Product Details

Immunogen:	KLH conjugated synthetic peptide derived from human PEX5	
Isotype:	IgG	
Cross-Reactivity:	Mouse	
Predicted Reactivity:	Human,Rat,Dog,Cow,Sheep,Pig,Horse	
Purification:	Purified by Protein A.	

## **Target Details**

Target:	PEX5
Alternative Name:	PEX5 (PEX5 Products)

#### Target Details

Bac	kar	ound:

Synonyms: FLJ50634, FLJ50721, FLJ51948, Peroxin 5, Peroxin-5, Peroxisomal biogenesis factor 5, Peroxisomal C terminal targeting signal import receptor, Peroxisomal C-terminal targeting signal import receptor, Peroxisomal targeting signal 1 receptor, Peroxisome receptor 1, pex5, PEX5\_HUMAN, PTS1 BP, PTS1 receptor, PTS1-BP, PTS1R, PXR1.

Background: The product of this gene binds to the C-terminal PTS1-type tripeptide peroxisomal targeting signal (SKL-type) and plays an essential role in peroxisomal protein import. Peroxins (PEXs) are proteins that are essential for the assembly of functional peroxisomes. The peroxisome biogenesis disorders (PBDs) are a group of genetically heterogeneous autosomal recessive, lethal diseases characterized by multiple defects in peroxisome function. The peroxisomal biogenesis disorders are a heterogeneous group with at least 14 complementation groups and with more than 1 phenotype being observed in cases falling into particular complementation groups. Although the clinical features of PBD patients vary, cells from all PBD patients exhibit a defect in the import of one or more classes of peroxisomal matrix proteins into the organelle. Defects in this gene are a cause of neonatal adrenoleukodystrophy (NALD), a cause of Zellweger syndrome (ZWS) as well as may be a cause of infantile Refsum disease (IRD). Alternatively spliced transcript variants encoding different isoforms have been identified. [provided by RefSeq, Oct 2008]

Gene ID:

5830

UniProt:

P50542

Pathways:

Monocarboxylic Acid Catabolic Process

#### **Application Details**

Application Notes:

IHC-P 1:200-400

IHC-F 1:100-500

Restrictions:

For Research Use only

#### Handling

Format:

Liquid

Concentration:

 $1 \mu g/\mu L$ 

Buffer:

Aqueous buffered solution containing 0.01M TBS ( pH 7.4) with 1 % BSA, 0.03 % Proclin300 and

50 % Glycerol.

Preservative:

ProClin

## Handling

Precaution of Use:	This product contains ProClin: a POISONOUS AND HAZARDOUS SUBSTANCE, which should be handled by trained staff only.
Storage:	-20 °C
Storage Comment:	Store at -20°C. Aliquot into multiple vials to avoid repeated freeze-thaw cycles.
Expiry Date:	12 months