



# **PEX6 Rabbit pAb**

CatalogNo: YN1038

## Key Features

Host Species

Rabbit

Reactivity

Human,Rat,Mouse

ApplicationsWB,ELISA

MW • 107kD (Observed) Isotype • IgG

## **Recommended Dilution Ratios**

WB 1:500-2000 ELISA 1:5000-20000

#### **Storage**

Storage*	-15°C to -25°C/1 year(Do not lower than -25°C)
Formulation	Liquid in PBS containing 50% glycerol, and 0.02% sodium azide.

#### **Basic Information**

Clonality Polyclonal

#### Immunogen Information

Immunogen Synthesized peptide derived from human protein . at AA range: 480-560

**Specificity** PEX6 Polyclonal Antibody detects endogenous levels of protein.

## Target Information

Gene name PEX6 PXAAA1

Protein Name	Peroxisome assembly factor 2 (PAF-2) (Peroxin-6) (Peroxisomal biogenesis factor 6) (Peroxisomal-type ATPase 1)		
	Organism	Gene ID	UniProt ID
	Human	<u>5190;</u>	<u>Q13608;</u>
	Mouse		<u>Q99LC9;</u>
	Rat		<u>P54777;</u>
Cellular Localization	Cytoplasm. Peroxisome membrane . Cell projection, cilium, photoreceptor outer segment . Associated with peroxisomal membranes. Localized at the base of the outer segment of photoreceptor cells (PubMed:26593283)		
Tissue specificity	Expressed in the retina, at higher le outer and inner segments.	vels in the photoreceptor lay	yer at the joint between the
Function	Disease:Defects in PEX6 are a cause of Zellweger syndrome (ZWS) [MIM:214100]. ZWS is a fatal peroxisome biogenesis disorder characterized by dysmorphic facial features, hepatomegaly, ocular abnormalities, renal cysts, hearing impairment, profound psychomotor retardation, severe hypotonia and neonatal seizures. Death occurs within the first year of life.,Disease:Defects in PEX6 are the cause of peroxisome biogenesis disorder complementation group 4 (PBD-CG4) [MIM:601498]; also known as PBD-CGC. PBD refers to a group of peroxisomal disorders arising from a failure of protein import into the peroxisomal membrane or matrix. The PBD group is comprised of four disorders: Zellweger syndrome (ZWS), neonatal adrenoleukodystrophy (NALD), infantile Refsum disease (IRD), and classical rhizomelic chondrodysplasia punctata (RCDP). ZWS, NALD and IRD are distinct from RCDP and constitute a clinical continuum of overlapping phenotypes known as the Zellweger spectrum. The PBD group is genetically heterogeneous with at least 14 distinct genetic groups as concluded from complementation studies.,Function:Involved in peroxisome biosynthesis. Required for stability of the PTS1 receptor. Anchored by PEX26 to peroxisome membranes, possibly to form heteromeric AAA ATPase complexes required for the import of proteins into peroxisomes.,similarity:Belongs to the AAA ATPase family.,subcellular location:Associated with peroxisomal membranes, postibut. Mediates the indirect interaction between PEX1 and PEX26.,		

## Validation Data

## **Contact information**

Orders:	order@immunoway.com
Support:	tech@immunoway.com
Telephone:	877-594-3616 (Toll Free), 408-747-0185
Website:	http://www.immunoway.com
Address:	2200 Ringwood Ave San Jose, CA 95131 USA



Please scan the QR code to access additional product information: **PEX6 Rabbit pAb**  For Research Use Only. Not for Use in Diagnostic Procedures.

Immunoway - 3 / 3