

Product datasheet for TP301756

PEX19 (NM_002857) Human Recombinant Protein

Product data:

Product Type:	Recombinant Proteins
Description:	Recombinant protein of human peroxisomal biogenesis factor 19 (PEX19), transcript variant 1, 20 µg
Species:	Human
Expression Host:	HEK293T
Expression cDNA Clone or AA Sequence:	>RC201756 protein sequence Red=Cloning site Green=Tags(s)

MAAAEEGCSVGAEADRELEELLESALDDDFDKAKPSPAPPSTTTAPDASGPQKRSPGDTAKDALFASQEKF
FQELFDSELASQATAEFKAMKELAE EEPHLVEQFQKLSEAGRVSMDTSQQEFTSCLKETLSGLAKNA
TDLQNSSMSEELTKAMEGLMDEGDGEGNILPIMQSIMQNLLSKDVLPSLKEITEKYPEWLQSHRESL
PPEQFEKYQEHSVMCKICEQFEAETPTDSETTQKARFEMVLDLMQQLQDLGHPPKELAGEMPPGLNFDL
DALNLSGPPGASGEQCLIM

TRTRPLEQKLISEEDLAANDILDYKDDDDKV

Tag:	C-Myc/DDK
Predicted MW:	32.6 kDa
Concentration:	>0.05 µg/µL as determined by microplate BCA method
Purity:	> 80% as determined by SDS-PAGE and Coomassie blue staining
Buffer:	25 mM Tris-HCl, 100 mM glycine, pH 7.3, 10% glycerol
Preparation:	Recombinant protein was captured through anti-DDK affinity column followed by conventional chromatography steps.
Note:	For testing in cell culture applications, please filter before use. Note that you may experience some loss of protein during the filtration process.
Storage:	Store at -80°C.
Stability:	Stable for 12 months from the date of receipt of the product under proper storage and handling conditions. Avoid repeated freeze-thaw cycles.
RefSeq:	NP_002848
Locus ID:	5824



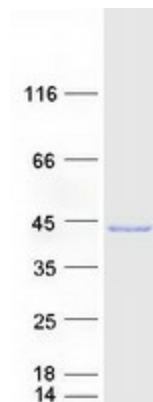
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UniProt ID:	P40855 , A0A0S2Z497
RefSeq Size:	3722
Cytogenetics:	1q23.2
RefSeq ORF:	897
Synonyms:	D1S2223E; HK33; PBD12A; PMP1; PMPI; PXF; PXMP1

Summary: This gene is necessary for early peroxisomal biogenesis. It acts both as a cytosolic chaperone and as an import receptor for peroxisomal membrane proteins (PMPs). Peroxisins (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. These disorders have at least 14 complementation groups, with more than one phenotype being observed for some 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 Zellweger syndrome (ZWS), as well as peroxisome biogenesis disorder complementation group 14 (PBD-CG14), which is also known as PBD-CGJ. Alternative splicing results in multiple transcript variants. [provided by RefSeq, Aug 2010]

Protein Families: Druggable Genome

Product images:



Coomassie blue staining of purified PEX19 protein (Cat# TP301756). The protein was produced from HEK293T cells transfected with PEX19 cDNA clone (Cat# [RC201756]) using MegaTran 2.0 (Cat# [TT210002]).