

Product datasheet for TL310493

OriGene Technologies, Inc.

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PEX19 Human shRNA Plasmid Kit (Locus ID 5824)

Product data:

Product Type: shRNA Plasmids

Product Name: PEX19 Human shRNA Plasmid Kit (Locus ID 5824)

Locus ID: 5824

Synonyms: D1S2223E; FLJ55296; HK33; housekeeping gene, 33kD; OTTHUMP00000031848; peroxisomal

biogenesis factor 19; peroxisomal farnesylated protein; PMP1; PMPI; PXF; PXMP1

Vector: pGFP-C-shLenti (TR30023)

E. coli Selection: Chloramphenicol (34 ug/ml)

Mammalian Cell

Selection:

Puromycin

Format: Lentiviral plasmids

Components: PEX19 - Human, 4 unique 29mer shRNA constructs in lentiviral GFP vector(Gene ID = 5824).

5µg purified plasmid DNA per construct

29-mer scrambled shRNA cassette in pGFP-C-shLenti Vector, TR30021, included for free.

RefSeq: NM 001131039, NM 001193644, NM 002857, NR 036492, NR 036493, NM 002857.1,

NM 002857.2, NM 002857.3, NM 001193644.1, NM 001131039.1, BC000496, BC000496.2,

BC064979, NM 002857.4

UniProt ID: P40855

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). 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. 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]







shRNA Design:

These shRNA constructs were designed against multiple splice variants at this gene locus. To be certain that your variant of interest is targeted, please contact techsupport@origene.com. If you need a special design or shRNA sequence, please utilize our custom shRNA service.

Performance Guaranteed: OriGene guarantees that the sequences in the shRNA expression cassettes are verified to correspond to the target gene with 100% identity. One of the four constructs at minimum are guaranteed to produce 70% or more gene expression knock-down provided a minimum transfection efficiency of 80% is achieved. Western Blot data is recommended over qPCR to evaluate the silencing effect of the shRNA constructs 72 hrs post transfection. To properly assess knockdown, the gene expression level from the included scramble control vector must be used in comparison with the target-specific shRNA transfected samples.

For non-conforming shRNA, requests for replacement product must be made within ninety (90) days from the date of delivery of the shRNA kit. To arrange for a free replacement with newly designed constructs, please contact Technical Services at techsupport@origene.com. Please provide your data indicating the transfection efficiency and measurement of gene expression knockdown compared to the scrambled shRNA control (Western Blot data preferred).