

Product datasheet for AR51020PU-S

Peroxin 26 / PEX26 (1-246, His-tag) Human Protein

Product data:

Product Type: Recombinant Proteins Description: Peroxin 26 / PEX26 (1-246, His-tag) human protein, 0.1 mg Species: Human E. coli **Expression Host:** MGSSHHHHHH SSGLVPRGSH MGSMKSDSST SAAPLRGLGG PLRSSEPVRA VPARAPAVDL Expression cDNA Clone or AA Sequence: LEEAADLLVV HLDFRAALET CERAWQSLAN HAVAEEPAGT SLEVKCSLCV VGIQALAEMD RWQEVLSWVL QYYQVPEKLP PKVLELCILL YSKMQEPGAV LDVVGAWLQD PANQNLPEYG ALAEFHVQRV LLPLGCLSEA EELVVGSAAF GEERRLDVLQ AIHTARQQQK QEHSGSEEAQ KPNLEGSVSH KFLSLPMLVR QLWDSAVSH Tag: His-tag Predicted MW: 29.3 kDa **Concentration:** lot specific **Purity:** >90% by SDS - PAGE **Buffer:** Presentation State: Purified State: Liquid purified protein Buffer System: 20 mM Tris-HCl buffer (pH 8.0) containing 0.15M NaCl, 20% glycerol, 1 mM DTT **Preparation:** Liquid purified protein Storage: Store undiluted at 2-8°C for one week or (in aliquots) at -20°C to -80°C for longer. Avoid repeated freezing and thawing. Stability: Shelf life: one year from despatch. RefSeq: NP 001121121 Locus ID: 55670 Q7Z412, A0A024R100 **UniProt ID:** Cytogenetics: 22q11.21 PBD7A; PBD7B; PEX26M1T; Pex26pM1T Synonyms:



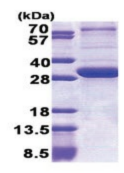
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Peroxin 26 / PEX26 (1-246, His-tag) Human Protein - AR51020PU-SSummary:This gene belongs to the peroxin-26 gene family. It is probably required for protein import
into peroxisomes. It anchors PEX1 and PEX6 to peroxisome membranes, possibly to form
heteromeric AAA ATPase complexes required for the import of proteins into peroxisomes.
Defects in this gene are the cause of peroxisome biogenesis disorder complementation group
8 (PBD-CG8). 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).
Alternatively spliced transcript variants have been identified for this gene. [provided by

Product images:



RefSeq, Dec 2010]

15% SDS-PAGE (3ug)

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