

Product datasheet for TP301904M

OriGene Technologies, Inc.

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CLN6 (NM_017882) Human Recombinant Protein

Product data:

Product Type: Recombinant Proteins

Description: Recombinant protein of human ceroid-lipofuscinosis, neuronal 6, late infantile, variant (CLN6),

100 µg

Species: Human
Expression Host: HEK293T

Expression cDNA Clone >RC201904 protein sequence or AA Sequence: Red=Cloning site Green=Tags(s)

MEATRRQHLGATGGPGAQLGASFLQARHGSVSADEAARTAPFHLDLWFYFTLQNWVLDFGRPIAMLVF

Ρ

LEWFPLNKPSVGDYFHMAYNVITPFLLLKLIERSPRTLPRSITYVSIIIFIMGASIHLVGDSVNHRLLFS GYQHHLSVRENPIIKNLKPETLIDSFELLYYYDEYLGHCMWYIPFFLILFMYFSGCFTASKAESLIPGPA LLLVAPSGLYYWYLVTEGQIFILFIFTFFAMLALVLHQKRKRLFLDSNGLFLFSSFALTLLLVALWVAWL

WNDPVLRKKYPGVIYVPEPWAFYTLHVSSRH

TRTRPLEQKLISEEDLAANDILDYKDDDDKV

Tag: C-Myc/DDK
Predicted MW: 35.7 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 060352



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Locus ID: 54982

UniProt ID: Q9NWW5

RefSeq Size: 2258
Cytogenetics: 15q23
RefSeq ORF: 933

Synonyms: CLN4A; HsT18960; nclf

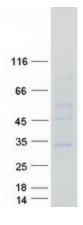
Summary: This gene is one of eight which have been associated with neuronal ceroid lipofuscinoses

(NCL). Also referred to as Batten disease, NCL comprises a class of autosomal recessive, neurodegenerative disorders affecting children. The genes responsible likely encode proteins involved in the degradation of post-translationally modified proteins in lysosomes. The primary defect in NCL disorders is thought to be associated with lysosomal storage function.

[provided by RefSeq, Oct 2008]

Protein Families: Transmembrane

Product images:



Coomassie blue staining of purified CLN6 protein (Cat# [TP301904]). The protein was produced from HEK293T cells transfected with CLN6 cDNA clone (Cat# [RC201904]) using MegaTran 2.0 (Cat# [TT210002]).