

# Product datasheet for TP302713M

## NDUFS4 (NM\_002495) Human Recombinant Protein

### **Product data:**

#### **Product Type: Recombinant Proteins Description:** Recombinant protein of human NADH dehydrogenase (ubiquinone) Fe-S protein 4, 18kDa (NADH-coenzyme Q reductase) (NDUFS4), nuclear gene encoding mitochondrial protein, 100 µg Species: Human **Expression Host:** HEK293T Expression cDNA Clone >RC202713 protein sequence or AA Sequence: Red=Cloning site Green=Tags(s) MAAVSMSVVLRQTLWRRRAVAVAALSVSRVPTRSLRTSSWRLAQDQTQDTQLITVDEKLDITTLTGVPEE HIKTRKVRIFVPARNNMQSGVNNTKKWKMEFDTRERWENPLMGWASTADPLSNMVLTFSTKEDAVSFAEK NGWSYDIEERKVPKPKSKSYGANFSWNKRTRVSTK **TRTRPLEQKLISEEDLAANDILDYKDDDDKV** Tag: C-Myc/DDK Predicted MW: 15.3 kDa **Concentration:** >0.05 µg/µL as determined by microplate BCA method > 80% as determined by SDS-PAGE and Coomassie blue staining Purity: **Buffer:** 25 mM Tris-HCl, 100 mM glycine, pH 7.3, 10% glycerol Recombinant protein was captured through anti-DDK affinity column followed by conventional **Preparation:** 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. Store at -80°C. Storage: Stable for 12 months from the date of receipt of the product under proper storage and Stability: handling conditions. Avoid repeated freeze-thaw cycles. **RefSeq:** NP 002486 4724 Locus ID: UniProt ID: O43181, A0A0S2Z433



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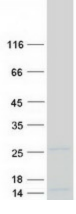
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### OriGene Technologies, Inc.

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	NDUFS4 (NM_002495) Human Recombinant Protein – TP302713M
RefSeq Size:	676
Cytogenetics:	5q11.2
RefSeq ORF:	525
Synonyms:	AQDQ; CI-18; CI-18 kDa; CI-AQDQ; MC1DN1
Summary:	This gene encodes an nuclear-encoded accessory subunit of the mitochondrial membrane respiratory chain NADH dehydrogenase (complex I, or NADH:ubiquinone oxidoreductase). Complex I removes electrons from NADH and passes them to the electron acceptor ubiquinone. Mutations in this gene can cause mitochondrial complex I deficiencies such as Leigh syndrome. Alternative splicing results in multiple transcript variants. [provided by RefSeq, Dec 2015]
Protein Families	Druggable Genome
Protein Pathway	<b>s:</b> Alzheimer's disease, Huntington's disease, Metabolic pathways, Oxidative phosphorylation, Parkinson's disease

## **Product images:**



Coomassie blue staining of purified NDUFS4 protein (Cat# [TP302713]). The protein was produced from HEK293T cells transfected with NDUFS4 cDNA clone (Cat# [RC202713]) using MegaTran 2.0 (Cat# [TT210002]).

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