

# **Product Name: POLG Rabbit Monoclonal Antibody**

Catalog #: AMRe87744

For research use only.

### **Summary**

**Description** Recombinant rabbit monoclonal antibody

HostRabbitApplicationWB

**Reactivity** Human

ConjugationUnconjugatedModificationUnmodified

**Isotype** IgG

Clonality Monoclonal
Form Liquid

Concentration

**Storage** Aliquot and store at -20°C (valid for 12 months). Avoid freeze/thaw cycles.

**Shipping** Ice bags

Supplied in 50mM Tris-Glycine(pH 7.4), 0.15M NaCl, 40% Glycerol, 0.01% sodium azide and **Buffer** 

0.05% protective protein. Stable for 12 months from date of receipt.

**Purification** Affinity Purification

## **Application**

**Dilution Ratio** WB 1:1000-1:5000

Molecular Weight Calculated MW:140 kDa; Observed MW:140 kDa

# **Antigen Information**

Gene Name POLG

Alternative Names PEO; MDP1; SCAE; MIRAS; POLG1; POLGA; SANDO; MTDPS4A; MTDPS4B

 Gene ID
 5428

 SwissProt ID
 P54098

**Immunogen** A synthetic peptide of human POLG

# **Background**

Mitochondrial DNA polymerase is heterotrimeric, consisting of a homodimer of accessory subunits plus a catalytic subunit. The protein encoded by this gene is the catalytic subunit of mitochondrial DNA polymerase. The encoded protein contains a

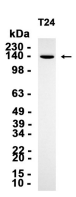
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polyglutamine tract near its N-terminus that may be polymorphic. Defects in this gene are a cause of progressive external ophthalmoplegia with mitochondrial DNA deletions 1 (PEOA1), sensory ataxic neuropathy dysarthria and ophthalmoparesis (SANDO), Alpers-Huttenlocher syndrome (AHS), and mitochondrial neurogastrointestinal encephalopathy syndrome (MNGIE). Two transcript variants encoding the same protein have been found for this gene. [provided by RefSeq, Jul 2008]

#### **Research Area**

## **Image Data**



Western blot analysis of extracts from T24 cells using POLG Rabbit Monoclonal Antibody at 1:1000.

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