

p40 (DeltaNp63) (PT0112R) rabbit mAb

Catalog No: YM7237

Reactivity: Human;(predicted: Mouse)

Applications: IHC;WB; ELISA

Target: p40/p63

Fields: >>MicroRNAs in cancer

Gene Name: TP63

Protein Name: p40

Human Gene Id: 8626

Human Swiss Prot

No:

Immunogen: Synthesized peptide derived from human p40 AA range:1-100

Specificity: This antibody detects endogenous levels of p40/p63

Formulation: PBS, 50% glycerol, 0.05% Proclin 300, 0.05% BSA

Source: Monoclonal, Rabbit IgG1, Kappa

Q9H3D4

Dilution: IHC 1:100-500, WB 1:500-1000, ELISA 1:5000-20000

Purification: Recombinant Expression and Affinity purified

Storage Stability: -15°C to -25°C/1 year(Do not lower than -25°C)

Molecularweight: 77kD

Background: tumor protein p63(TP63) Homo sapiens This gene encodes a member of the

p53 family of transcription factors. The functional domains of p53 family proteins include an N-terminal transactivation domain, a central DNA-binding domain and

an oligomerization domain. Alternative splicing of this gene and the use of



alternative promoters results in multiple transcript variants encoding different isoforms that vary in their functional properties. These isoforms function during skin development and maintenance, adult stem/progenitor cell regulation, heart development and premature aging. Some isoforms have been found to protect the germline by eliminating oocytes or testicular germ cells that have suffered DNA damage. Mutations in this gene are associated with ectodermal dysplasia, and cleft lip/palate syndrome 3 (EEC3); split-hand/foot malformation 4 (SHFM4); ankyloblepharon-ectodermal defects-cleft lip/palate; ADULT syndrome (acrodermato-ungual-lacrim

Function:

cofactor:Binds 1 zinc ion per subunit.,disease:Defects in TP63 are a cause of cervical, colon, head and neck, lung and ovarian cancers.,disease:Defects in TP63 are a cause of ectodermal dysplasia Rapp-Hodgkin type (EDRH) [MIM:129400]; also called Rapp-Hodgkin syndrome or anhidrotic ectodermal dysplasia with cleft lip/palate. Ectodermal dysplasia defines a heterogeneous group of disorders due to abnormal development of two or more ectodermal structures. EDRH is characterized by the combination of anhidrotic ectodermal dysplasia, cleft lip, and cleft palate. The clinical syndrome is comprised of a characteristic facies (narrow nose and small mouth), wiry, slow-growing, and uncombable hair, sparse eyelashes and eyebrows, obstructed lacrimal puncta/epiphora, bilateral stenosis of external auditory canals, microsomia, hypodontia, cone-shaped incisors, enamel hypoplasia, dystrophic nails, and

Subcellular Location:

Expression:

Nucleus.

Widely expressed, notably in heart, kidney, placenta, prostate, skeletal muscle, testis and thymus, although the precise isoform varies according to tissue type. Progenitor cell layers of skin, breast, eye and prostate express high levels of DeltaN-type isoforms. Isoform 10 is predominantly expressed in skin squamous cell carcinomas, but not in normal skin tissues.

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