

Keratin 17 (Phospho Ser44) rabbit pAb

Catalog No: YP1377

Reactivity: Human; Rat; Mouse;

Applications: WB;ELISA;IHC

Target: Cytokeratin 17

Fields: >>Estrogen signaling pathway;>>Staphylococcus aureus infection

Gene Name: KRT17

Protein Name: Keratin 17 (Ser44)

Human Gene Id: 3872

Human Swiss Prot

Q04695

No:

Mouse Gene ld: 16667

Mouse Swiss Prot

Q9QWL7

No:

Rat Gene Id: 287702

Rat Swiss Prot No: Q6IFU8

Immunogen: Synthesized phosho peptide around human Keratin 17 (Ser44)

Specificity: This antibody detects endogenous levels of Human Keratin 17 (phospho-Ser44)

Formulation : Liquid in PBS containing 50% glycerol, 0.5% BSA and 0.02% sodium azide.

Source: Polyclonal, Rabbit, IgG

Dilution: WB 1:500-2000;IHC 1:50-300; ELISA 2000-20000

Purification: The antibody was affinity-purified from rabbit serum by affinity-chromatography

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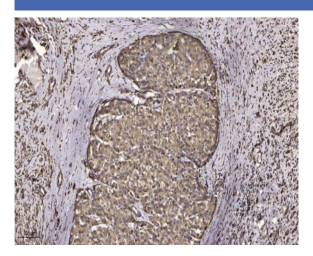
Modifications:

Phospho

using specific immunogen. **Concentration:** 1 mg/ml -15°C to -25°C/1 year(Do not lower than -25°C) **Storage Stability:** Observed Band: 48kD This gene encodes the type I intermediate filament chain keratin 17, expressed **Background:** in nail bed, hair follicle, sebaceous glands, and other epidermal appendages. Mutations in this gene lead to Jackson-Lawler type pachyonychia congenita and steatocystoma multiplex. [provided by RefSeg, Aug 2008], **Function:** disease:Defects in KRT17 are a cause of pachyonychia congenita type 2 (PC2) [MIM:167210]; also known as pachyonychia congenita Jackson-Lawler type. PC2 is an autosomal dominant ectodermal dysplasia characterized by hypertrophic nail dystrophy resulting in onchyogryposis (thickening and increase in curvature of the nail), palmoplantar keratoderma and hyperhidrosis, follicular hyperkeratosis, multiple epidermal cysts, absent/sparse eyebrow and body hair, and by the presence of natal teeth..disease:Defects in KRT17 are a cause of steatocystoma multiplex (SM) [MIM:184500]. SM is a disease characterized by round or oval cystic tumors widely distributed on the back, anterior trunk, arms, scrotum, and thighs., disease: KRT16 and KRT17 are coexpressed only in pathological situations such as metaplasias and carcinomas of the uterine cervix and in psoriasis vulgaris., function: May play a role in the Subcellular Cytoplasm. Location: Expressed in the outer root sheath and medulla region of hair follicle specifically **Expression:** from eyebrow and beard, digital pulp, nail matrix and nail bed epithelium, mucosal stratified squamous epithelia and in basal cells of oral epithelium, palmoplantar epidermis and sweat and mammary glands. Also expressed in myoepithelium of prostate, basal layer of urinary bladder, cambial cells of sebaceous gland and in exocervix (at protein level). Sort: 8884 No4: Host: Rabbit



Products Images



Immunohistochemical analysis of paraffin-embedded human liver cancer. 1, Antibody was diluted at 1:200(4° overnight). 2, Tris-EDTA,pH9.0 was used for antigen retrieval. 3,Secondary antibody was diluted at 1:200(room temperature, 45min).