



VHL Polyclonal Antibody

Catalog No	YP-Ab-00589
Isotype	IgG
Reactivity	Human;Rat;Mouse;
Applications	IHC;IF;ELISA
Gene Name	VHL
Immunogen	The antiserum was produced against synthesized peptide derived from the N-terminal region of human VHL. AA range:1-50
Specificity	The antibody detects endogenous VHL
Formulation	Liquid in PBS containing 50% glycerol, 0.5% BSA and 0.02% sodium azide.
Source	Polyclonal, Rabbit,IgG
Purification	The antibody was affinity-purified from rabbit antiserum by affinity-chromatography using epitope-specific immunogen.
Dilution	IHC-p 1:50-200, ELISA 1:10000-20000. IF 1:50-200
Concentration	1 mg/ml
Purity	≥90%
Storage Stability	-20°C/1 year
Synonyms	HRCA1;Protein G7;pVHL;RCA1;VHL;VHL1;Von Hippel-Lindau disease tumor suppressor (Protein G7) (pVHL);Von Hippel-Lindau disease tumor suppressor (Protein G7;pVHL)
Observed Band	19-24kD
Calculated Molecular Weight	19-24kD
Cell Pathway	[Isoform 1]: Cytoplasm. Membrane; Peripheral membrane protein. Nucleus. Found predominantly in the cytoplasm and with less amounts nuclear or membrane-associated. Colocalizes with ADRB2 at the cell membrane.; [Isoform 3]: Cytoplasm. Nucleus. Equally distributed between the nucleus and the cytoplasm but not membrane-associated.
Tissue Specificity	Expressed in the adult and fetal brain and kidney.
Function	disease:Defects in VHL are a cause of pheochromocytoma [MIM:171300]. The pheochromocytomas are catecholamine-producing, chromaffin tumors that arise in the adrenal medulla in 90% of cases. In the remaining 10% of cases, they develop in extra-adrenal sympathetic ganglia and may be referred to as "paraganglioma." Pheochromocytoma usually presents with hypertension. Approximately 10% of pheochromocytoma is hereditary. The genetic basis for



most cases of non-syndromic familial pheochromocytoma is unknown.,
disease:Defects in VHL are a cause of renal cell carcinoma type 1 (RCC1)
[MIM:144700]; also called hypernephroma or adenocarcinoma of kidney.
Familial renal cell carcinoma syndromes form a group of diseases characterized
by a predisposition to development of renal cell carcinomas (RCCs) with various
histological subtypes.,disease:Defects in VHL are the cause of erythrocytosis
familial type

Background

von Hippel-Lindau tumor suppressor(VHL) Homo sapiens Von Hippel-Lindau syndrome (VHL) is a dominantly inherited familial cancer syndrome predisposing to a variety of malignant and benign tumors. A germline mutation of this gene is the basis of familial inheritance of VHL syndrome. The protein encoded by this gene is a component of the protein complex that includes elongin B, elongin C, and cullin-2, and possesses ubiquitin ligase E3 activity. This protein is involved in the ubiquitination and degradation of hypoxia-inducible-factor (HIF), which is a transcription factor that plays a central role in the regulation of gene expression by oxygen. RNA polymerase II subunit POLR2G/RPB7 is also reported to be a target of this protein. Alternatively spliced transcript variants encoding distinct isoforms have been observed. [provided by RefSeq, Jul 2008],

matters needing attention

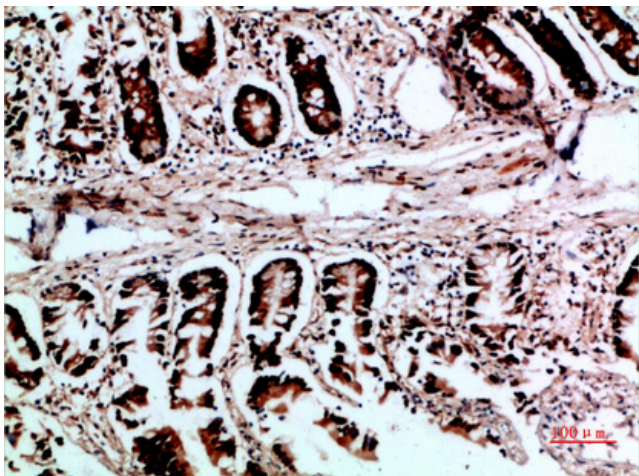
Avoid repeated freezing and thawing!

Usage suggestions

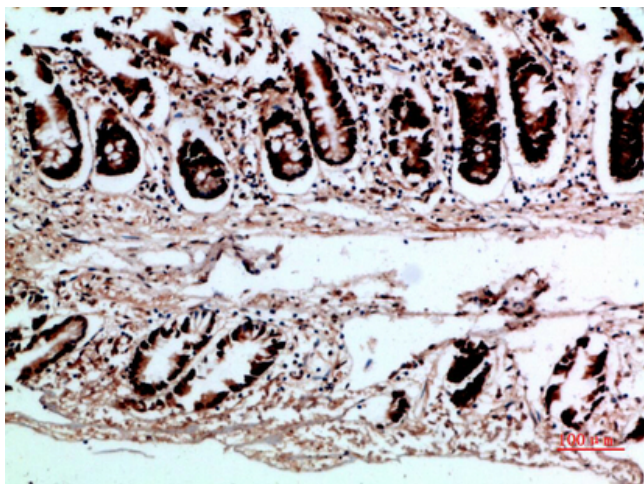
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Immunohistochemical analysis of paraffin-embedded human-colon, antibody was diluted at 1:200



Immunohistochemical analysis of paraffin-embedded human-colon, antibody was diluted at 1:200