

Human VHL antibody

Catalog Number: ATGA0492

PRODUCT INFORMATION

Catalog number

ATGA0492

Clone No.

AT82B10

Product type

Monoclonal Antibody

UnitProt No.

P40337

NCBI Accession No.

NP_000542

Alternative Names

Von Hippel-Lindau tumor suppressor isoform 1, Von Hippel-Lindau tumor suppressor isoform 1, HRCA1, RCA1, VHL1, Von Hippel-Lindau tumor suppressor isoform 1 pVHL, G7 protein, Elongin binding protein, HRCA 1, RCA 1, VHL, VHL 1, VHLH, Von Hippel Lindau disease tumor suppressor, von Hippel Lindau syndrome, von Hippel Lindau tumor suppressor, AVH0616

Additional Information

AVH0616 has been replaced with a catalog number ATGA0492.

PRODUCT SPECIFICATION

Antibody Host

Mouse

Reacts With

Human

Concentration

1mg/ml (determined by BCA assay)

Formulation

Liquid in. Phosphate-Buffered Saline (pH 7.4) with 0.02% Sodium Azide, 10% glycerol

Immunogen

Recombinant human VHL (1-154aa) purified from E. coli

Isotype

IgG2b kappa

Purification Note

By protein-A affinity chromatography

Application

ELISA, WB, ICC/IF

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Usage

The antibody has been tested by ELISA, Western blot analysis and ICC/IF to assure specificity and reactivity. Since application varies, however, each investigation should be titrated by the reagent to obtain optimal results.

Storage

Can be stored at +2C to +8C for 1 week. For long term storage, aliquot and store at -20C to -80C. Avoid repeated freezing and thawing cycles.

BACKGROUND

Description

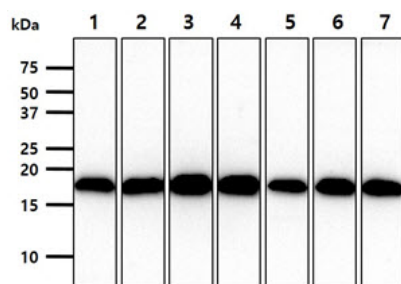
Von Hippel-Lindau disease (VHL) is a dominant inherited syndrome characterized by the predisposition to develop various kinds of benign and malignant tumors, including clear cell renal carcinomas, pheochromocytomas and hemangioblastomas of the central nervous system and retina. VHL syndrome is caused by germline mutation in the VHL tumor suppressor, and VHL tumors are associated with loss or mutation of the remaining wild-type allele. VHL has two domains: a roughly 100-residue NH₂-terminal domain rich in beta sheet (beta-domain) and a smaller alpha-helical domain (alpha-domain), held together by two linkers and a polar interface. VHL protein is also involved in the degradation of hypoxia-inducible factor (HIF).

General References

Latif F., et al. (1993) Science. 260(5112): 1317-20.
Duan DR., et al. (1995) PNAS. 92(14): 6459-63.
Maxwell PH., et al. (1999) Nature. 399(6733): 271-5.

DATA

Western blot analysis (WB)



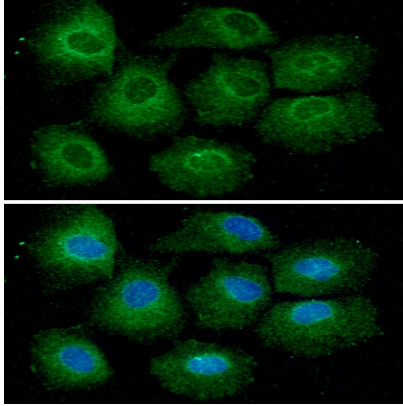
The cell lysates (40ug) were resolved by SDS-PAGE, transferred to PVDF membrane and probed with anti-human VHL antibody (1:1000). Proteins were visualized using a goat anti-mouse secondary antibody conjugated to HRP and an ECL detection system.

Lane 1.: HepG2 cell lysate
Lane 2.: HeLa cell lysate
Lane 3.: Raji cell lysate
Lane 4.: Jurkat cell lysate
Lane 5.: A549 cell lysate
Lane 6.: MCF7 cell lysate
Lane 7.: PC3 cell lysate

Immunocytochemistry/Immunofluorescence (ICC/IF)

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ICC/IF analysis of VHL in Hep3B. The cell was stained with ATGA0492 (1:100). The secondary antibody (green) was used Alexa Fluor 488. DAPI was stained the cell nucleus (blue).