

## Phospho-RB(S612) Antibody Blocking peptide

Synthetic peptide Catalog # BP3236a

# **Specification**

## Phospho-RB(S612) Antibody Blocking peptide - Product Information

**Primary Accession** 

P06400

# Phospho-RB(S612) Antibody Blocking peptide - Additional Information

**Gene ID 5925** 

#### **Other Names**

Retinoblastoma-associated protein, p105-Rb, pRb, Rb, pp110, RB1

# **Target/Specificity**

The synthetic peptide sequence used to generate the antibody <a href=/product/products/AP3236a>AP3236a</a> was selected from the 605-619 <CR>region of human Phospho-RB-S612. A 10 to 100 fold molar excess to antibody is recommended. Precise conditions should be optimized for a particular assay.

### **Format**

Peptides are lyophilized in a solid powder format. Peptides can be reconstituted in solution using the appropriate buffer as needed.

#### Storage

Maintain refrigerated at 2-8°C for up to 6 months. For long term storage store at -20°C.

#### **Precautions**

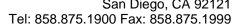
This product is for research use only. Not for use in diagnostic or therapeutic procedures.

## Phospho-RB(S612) Antibody Blocking peptide - Protein Information

## Name RB1

#### **Function**

Tumor suppressor that is a key regulator of the G1/S transition of the cell cycle (PubMed:<a href="http://www.uniprot.org/citations/10499802" target="\_blank">10499802</a>). The hypophosphorylated form binds transcription regulators of the E2F family, preventing transcription of E2F-responsive genes (PubMed:<a href="http://www.uniprot.org/citations/10499802" target="\_blank">10499802</a>). Both physically blocks E2Fs transactivating domain and recruits chromatin- modifying enzymes that actively repress transcription (PubMed:<a href="http://www.uniprot.org/citations/10499802" target="\_blank">10499802</a>). Cyclin and CDK-dependent phosphorylation of RB1 induces its dissociation from E2Fs, thereby activating transcription of E2F responsive genes and triggering entry into S phase (PubMed:<a href="http://www.uniprot.org/citations/10499802" target="\_blank">10499802</a>). RB1 also promotes the G0-G1 transition upon phosphorylation and activation by CDK3/cyclin-C (PubMed:<a href="http://www.uniprot.org/citations/15084261" target="\_blank">15084261</a>). Directly





involved in heterochromatin formation by maintaining overall chromatin structure and, in particular, that of constitutive heterochromatin by stabilizing histone methylation. Recruits and targets histone methyltransferases SUV39H1, KMT5B and KMT5C, leading to epigenetic transcriptional repression. Controls histone H4 'Lys-20' trimethylation. Inhibits the intrinsic kinase activity of TAF1. Mediates transcriptional repression by SMARCA4/BRG1 by recruiting a histone deacetylase (HDAC) complex to the c-FOS promoter. In resting neurons, transcription of the c-FOS promoter is inhibited by BRG1- dependent recruitment of a phospho-RB1-HDAC1 repressor complex. Upon calcium influx, RB1 is dephosphorylated by calcineurin, which leads to release of the repressor complex (By similarity).

#### **Cellular Location**

Nucleus. Note=During keratinocyte differentiation, acetylation by KAT2B/PCAF is required for nuclear localization.

#### **Tissue Location**

Expressed in the retina. Expressed in foreskin keratinocytes (at protein level) (PubMed:20940255)

### Phospho-RB(S612) Antibody Blocking peptide - Protocols

Provided below are standard protocols that you may find useful for product applications.

#### • Blocking Peptides

Phospho-RB(S612) Antibody Blocking peptide - Images

# Phospho-RB(S612) Antibody Blocking peptide - Background

Retinoblastoma (RB) is an embryonic malignant neoplasm of retinal origin. It almost always presents in early childhood and is often bilateral. Spontaneous regression ('cure') occurs in some cases.[supplied by OMIM].

### Phospho-RB(S612) Antibody Blocking peptide - References

Dasgupta, P., et al., Mol. Cell. Biol. 24(21):9527-9541 (2004).Cui, X., et al., Hum. Pathol. 35(10):1189-1195 (2004).Borah, S., et al., J. Virol. 78(19):10336-10347 (2004).Dasgupta, P., et al., J. Biol. Chem. 279(37):38762-38769 (2004).Lohmann, D.R., et al., J. Biol. Chem. 129(1):23-28 (2004).