

Nibrin (phospho Ser343) Polyclonal Antibody

Catalog No: YP0194

Reactivity: Human;Rat

Applications: WB;ELISA

Target: Nibrin

Fields: >>Homologous recombination;>>Cellular senescence

O60934

Q9R207

Gene Name: NBN

Protein Name: Nibrin

Human Gene Id: 4683

Human Swiss Prot

No:

Mouse Swiss Prot

No:

Rat Gene ld: 85482

Rat Swiss Prot No: Q9JIL9

Immunogen: The antiserum was produced against synthesized peptide derived from human

p95/NBS1 around the phosphorylation site of Ser343. AA range:310-359

Specificity: Phospho-Nibrin (S343) Polyclonal Antibody detects endogenous levels of Nibrin

protein only when phosphorylated at S343.

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

Source: Polyclonal, Rabbit, IgG

Dilution: WB 1:500 - 1:2000. ELISA: 1:5000. Not yet tested in other applications.

Purification: The antibody was affinity-purified from rabbit antiserum by affinity-

chromatography using epitope-specific immunogen.



Concentration: 1 mg/ml

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

Observed Band: 95kD

Cell Pathway: Homologous recombination;

Background: Mutations in this gene are associated with Nijmegen breakage syndrome, an

autosomal recessive chromosomal instability syndrome characterized by microcephaly, growth retardation, immunodeficiency, and cancer predisposition. The encoded protein is a member of the MRE11/RAD50 double-strand break repair complex which consists of 5 proteins. This gene product is thought to be involved in DNA double-strand break repair and DNA damage-induced

checkpoint activation. [provided by RefSeq, Jul 2008],

Function: disease:Defects in NBN are a cause of genetic susceptibility to breast cancer

(BC) [MIM:114480]. BC is an extremely common malignancy, affecting one in eight women during their lifetime. A positive family history has been identified as major contributor to risk of development of the disease, and this link is striking for early-onset breast cancer., disease:Defects in NBN are the cause of Nijmegen breakage syndrome (NBS) [MIM:251260]. NBS is an autosomal recessive syndrome characterized by chromosomal instability, radiation sensitivity, microcephaly, growth retardation, immunodeficiency and predisposition to cancer, particularly to lymphoid malignancies., disease:Defects in NBN may be associated with aplastic anemia [MIM:609135]. Aplastic anemia is a disease of bone-marrow failure characterized by peripheral pancytopenia and marrow

hypoplasia. Most of the cases of aplastic anemia are idiopa

Subcellular

Nucleus . Nucleus, PML body . Chromosome, telomere . Chromosome .

Location :

Localizes to discrete nuclear foci after treatment with genotoxic agents

(PubMed:26438602, PubMed:10783165, PubMed:26215093). Acetylation of 'Lys-5' of histone H2AX (H2AXK5ac) promotes NBN/NBS1 assembly at the sites

of DNA damage (PubMed:26438602). .

Expression: Ubiquitous (PubMed:9590180). Expressed at high levels in testis

(PubMed:9590180).

Sort: 10844

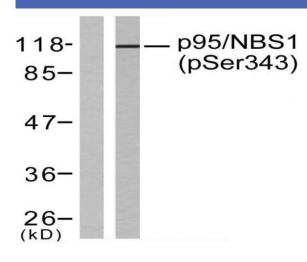
No4:

Host: Rabbit

Modifications : Phospho



Products Images



Western blot analysis of lysates from Jurkat cells, using p95/NBS1 (Phospho-Ser343) Antibody. The lane on the left is blocked with the phospho peptide.