

# NBN Monoklonaler Antikörper

Katalog-Nr.: CL488-66980

## Allgemeine Informationen

Katalog-Nr.:	GenBank-Zugangsnummer:	Reinigungsmethode:
CL488-66980	BC136803	Protein-A-Reinigung
Größe:	GenID (NCBI):	CloneNo.:
100ul , Konzentration: 1000 µg/ml von 4683		1E11E10
Nanodrop;	Vollständiger Name:	Empfohlene Verdünnungen:
	nibrin	IF 1:50-1:500
Wirt:	Berechnete Masse:	Anregungs-/Emissionsmaxima-
Maus	85 kDa	Wellenlängen:
Isotyp:		493 nm / 522 nm
IgG2a		
Immunogen Katalognummer:		
AG19320		

## Anwendungen

Geprüfte Anwendungen:	Positivkontrollen:
FC (Intra), IF	IF : HepG2-Zellen,
Getestete Reaktivität:	
Human	

## Hintergrundinformationen

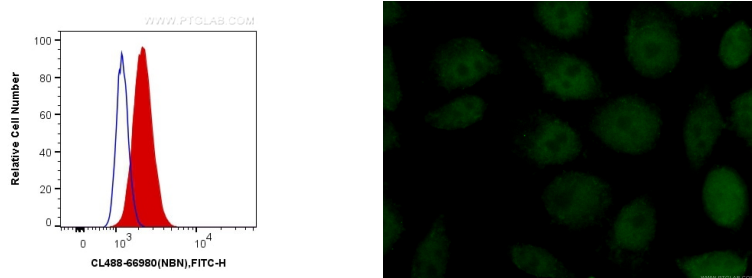
NBN, also named as NBS, NBS1 and P95, is a component of the MRE11/RAD50/NBN (MRN complex) which plays a critical role in the cellular response to DNA damage and the maintenance of chromosome integrity. The complex is involved in double-strand break (DSB) repair, DNA recombination, maintenance of telomere integrity, cell cycle checkpoint control and meiosis. The complex possesses single-strand endonuclease activity and double-strand-specific 3'-5' exonuclease activity, which are provided by MRE11A. NBN modulate the DNA damage signal sensing by recruiting PI3/PI4-kinase family members ATM, ATR, and probably DNA-PKcs to the DNA damage sites and activating their functions. NBN also functions in telomere length maintenance by generating the 3' overhang which serves as a primer for telomerase dependent telomere elongation. NBN is a major player in the control of intra-S-phase checkpoint and there is some evidence that NBN is involved in G1 and G2 checkpoints. Defects in NBN are the cause of Nijmegen breakage syndrome (NBS). Defects in NBN are a cause of genetic susceptibility to breast cancer (BC). Defects in NBN may be associated with aplastic anemia. Defects in NBN might play a role in the pathogenesis of childhood acute lymphoblastic leukemia (ALL). The antibody is specific to NBN. The full-length NBN protein, with an apparent molecular weight of 95 kDa and the two protein fragments of 26 and 70 kDa arising from the c.657\_661del5 (p.K219fsX19) mutation, and the 80 kDa protein found in patient RR with the mutation c.742\_743insGG leading to excision of exons 6 and 7 from the NBN mRNA are shown. (PMID: 26265251) The predicted molecular weight of NBN protein (p95) is 85kDa, actually detection result is about 95kDa (PMID: 23762398).

## Lagerung

Lagerungsbedingungen:  
Bei -20°C lagern. Vor Licht schützen. Nach dem Versand ein Jahr stabil.  
Lagerungspuffer:  
BS mit 50% Glyzerin, 0,05% Proclin300, 0,5% BSA, pH 7,3.  
Aliquotieren ist nicht notwendig bei -20°C Lagerung

\*\*\* 20ul-Größen enthalten 0.1% BSA

## Ausgewählte Validierungsdaten



1X10<sup>6</sup> HeLa cells were intracellularly stained with 0.4 ug Coralite® Plus 488 Anti-Human NBN (CL488-66980, Clone:1E11E10) (red), or 0.4 ug Control Antibody. Cells were fixed and permeabilized with Transcription Factor Staining Buffer Kit (PF00011).

Immunofluorescent analysis of (4% PFA) fixed HepG2 cells using CL488-66980 (NBN antibody) at dilution of 1:100.