

MaxPab®

SNRPN purified MaxPab mouse polyclonal antibody (B01P)

Catalog # H00006638-B01P

Size 50 ug

Applications



Western Blot (Transfected lysate)

Western Blot analysis of SNRPN expression in transfected 293T cell line (<u>H00006638-T01</u>) by SNRPN MaxPab polyclonal antibody.

Lane 1: SNRPN transfected lysate(26.51 KDa). Lane 2: Non-transfected lysate.

Specification	
Product Description	Mouse polyclonal antibody raised against a full-length human SNRPN protein.
Immunogen	SNRPN (AAH03180, 1 a.a. ~ 240 a.a) full-length human protein.
Sequence	MTVGKSSKMLQHIDYRMRCILQDGRIFIGTFKAFDKHMNLILCDCDEFRKIKPKNAKQPEREEKRVL GLVLLRGENLVSMTVEGPPPKDTGIARVPLAGAARGPGVGRAAGRGVPAGVPIPQAPAGLAGPV RGVGGPSQQVMTPQGRGTVAAAAVAATASIAGAPTQYPPGRGTPPPPVGRATPPPGIMAPPPG MRPPMGPPIGLPPARGTPIGMPPPGMRPPPPGIRGPPPPGMRPPRP
Host	Mouse
Reactivity	Human
Interspecies Antigen Sequence	Mouse (99); Rat (99)
Quality Control Testing	Antibody reactive against mammalian transfected lysate.
Storage Buffer	In 1x PBS, pH 7.4
Storage Instruction	Store at -20°C or lower. Aliquot to avoid repeated freezing and thawing.

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Protocol Download

Gene Info — SNRPN	
Entrez GenelD	<u>6638</u>
GeneBank Accession#	<u>BC003180</u>
Protein Accession#	<u>AAH03180</u>
Gene Name	SNRPN
Gene Alias	DKFZp686C0927, DKFZp686M12165, DKFZp761I1912, DKFZp762N022, FLJ33569, FLJ3699 6, FLJ39265, HCERN3, MGC29886, PWCR, RT-LI, SM-D, SMN, SNRNP-N, SNURF-SNRPN
Gene Description	small nuclear ribonucleoprotein polypeptide N
Omim ID	<u>176270 182279</u>
Gene Ontology	Hyperlink
Gene Ontology Gene Summary	Hyperlink The protein encoded by this gene is one polypeptide of a small nuclear ribonucleoprotein comple x and belongs to the snRNP SMB/SMN family. The protein plays a role in pre-mRNA processing, possibly tissue-specific alternative splicing events. Although individual snRNPs are believed to re cognize specific nucleic acid sequences through RNA-RNA base pairing, the specific role of this f amily member is unknown. The protein arises from a bicistronic transcript that also encodes a pro tein identified as the SNRPN upstream reading frame (SNURF). Multiple transcription initiation sit es have been identified and extensive alternative splicing occurs in the 5' untranslated region. Ad ditional splice variants have been described but sequences for the complete transcripts have not been determined. The 5' UTR of this gene has been identified as an imprinting center. Alternative splicing or deletion caused by a translocation event in this paternally-expressed region is respons ible for Angelman syndrome or Prader-Willi syndrome due to parental imprint switch failure. [provi ded by RefSeq



Disease

- Autistic Disorder
- Genetic Predisposition to Disease