For Research Use Only

ATRX-Specific Polyclonal antibody

Catalog Number: 19788-1-AP



Purification Method:

Antigen affinity purification

Basic Information

Catalog Number:

Bradford method using BSA as the

19788-1-AP

standard;

Source:

Rabbit

Isotype:

GeneID (NCBI): 150ul , Concentration: 133 $\mu g/ml$ by

UNIPROT ID:

NM 000489

P46100 Full Name:

alpha thalassemia/mental retardation syndrome X-linked (RAD54 homolog, S. cerevisiae)

GenBank Accession Number:

Calculated MW: 283 kDa

Applications

Tested Applications:

FIISA

Species Specificity:

human

Background Information

 $A TRX, also \ named \ as \ RAD54L \ and \ XH2, belongs \ to \ the \ SNF2/RAD54 \ helicase \ family. \ ATR \ could \ be \ a \ global$ transcriptional regulator. ATRX modifies gene expression by affecting chromatin. It may be involved in brain development and facial morphogenesis. Defects in ATRX are the cause of X-linked alpha-thalassemia/mental retardation syndrome (ATR-X) which is an X-linked disorder comprising severe psychomotor retardation, facial dysmorphism, urogenital abnormalities, and alpha-thalassemia. Defects in ATRX are the cause of mental retardation syndromic X-linked with hypotonic facies syndrome type 1 (MRXSHF1) which also called Carpenter-Waziri syndrome (CWS), Juberg-Marsidi syndrome (JMS), Smith-Fineman-Myers syndrome type 1 (SFM1). This antibody is specific to ATRX.

Storage

Store at -20°C. Stable for one year after shipment.

PBS with 0.02% sodium azide and 50% glycerol pH 7.3.

Aliquoting is unnecessary for -20°C storage

*** 20ul sizes contain 0.1% BSA

in USA), or 1(312) 455-8498 (outside USA)

W: ptglab.com

Selected Validation Data