

Produktinformation



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Zellkultur & Verbrauchsmaterial



Diagnostik & molekulare Diagnostik



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Product datasheet MON240257



Rabbit anti-ATRX antibody, clone SQab22251 (monoclonal)

Clone no. SQab22251 MONOSAN

Product name Rabbit anti-ATRX antibody, clone SQab22251 (monoclonal)

Host Rabbit

Applications IHC-P

Species reactivity Human

Conjugate -

Immunogen Recombinant protein fragment within Human ATRX.

Isotype -

Clonality Monoclonal

Clone number SQab22251

Size 100 ul

Concentration n/a

Format Purification with Protein A.

Storage buffer PBS, 0.01% Sodium azide, 40% Glycerol and 0.05%BSA.

Storage until expiry date -20°C

FOR RESEARCH USE ONLY. NOT FOR USE IN DIAGNOSTIC PROCEDURES

Product datasheet

MON240257



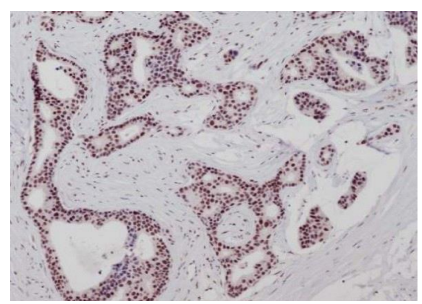
Rabbit anti-ATRX antibody, clone SQab22251 (monoclonal)

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Additional info

Application note: IHC-P: Antigen Retrieval: Heat mediated was performed using Tris/EDTA buffer (pH 9.0). Incubate the samples at RT (18-25°C) for 30 min. * The dilutions indicate recommended starting dilutions and the optimal dilutions or concentrations should be determined by the scientist. Storage instruction: For continuous use, store undiluted antibody at 2-8°C for up to a week. For long-term storage, aliquot and store at -20°C or below. Storage in frost free freezers is not recommended. Avoid repeated freeze/thaw cycles. Suggest spin the vial prior to opening. The antibody solution should be gently mixed before use. Background: The protein encoded by this gene contains an ATPase/helicase domain, and thus it belongs to the SWI/SNF family of chromatin remodeling proteins. This protein is found to undergo cell cycledependent phosphorylation, which regulates its nuclear matrix and chromatin association, and suggests its involvement in the gene regulation at interphase and chromosomal segregation in mitosis. Mutations in this gene are associated with an X-linked mental retardation (XLMR) syndrome most often accompanied by alpha-thalassemia (ATRX) syndrome. These mutations have been shown to cause diverse changes in the pattern of DNA methylation, which may provide a link between chromatin remodeling, DNA methylation, and gene expression in developmental processes. Multiple alternatively spliced transcript variants encoding distinct isoforms have been reported. [provided by RefSeq, Aug 2013]

Images



Immunohistochemistry: Formalin/PFA-fixed and paraffin-embedded sections of Human breast carcinoma tissue stained with anti-ATRX antibody [SQab22251]. Antigen Retrieval: Heat tissue section in Tris-EDTA buffer (pH 9.0).

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References

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