

Anti-Aurora B Antibody

Mouse Monoclonal Antibody Catalog # AH13604

Product Information

Application	WB, IF, FC
Primary Accession	<u>Q96GD4</u>
Other Accession	<u>442658</u>
Reactivity	Human
Host	Mouse
Clonality	Monoclonal
Isotype	Mouse / IgG2b
Clone Names	AURKB/1521
Calculated MW	39311

Additional Information

Gene ID	9212
Other Names	AIK2; AIM-1; ARK-2; AurB; AURKB; Aurora-1; Aurora and Ipl1 like midbody associated protein 1; Aurora kinase B; Aurora-B; Aurora-related kinase 2; Aurora/IPL1-related kinase 2; IPL1; Protein phosphatase 1 regulatory subunit 48 (PPP1R48); Serine/threonine-protein kinase 12; Serine/threonine-protein kinase aurora-B; STK1; STK12; STK5
Application Note	Flow Cytometry (0.5-1ug/million cells); ,Immunofluorescence (1-2ug/ml); ,Western Blotting (0.5-1ug/ml),Optimal dilution for a specific application should be determined.
Format	200ug/ml of Ab purified from Bioreactor Concentrate by Protein A/G. Prepared in 10mM PBS with 0.05% BSA & 0.05% azide. Also available WITHOUT BSA & azide at 1.0mg/ml.
Storage	Store at 2 to 8°C.Antibody is stable for 24 months.
Precautions	Anti-Aurora B Antibody is for research use only and not for use in diagnostic or therapeutic procedures.

Protein Information

Name	AURKB
Function	Serine/threonine-protein kinase component of the chromosomal passenger complex (CPC), a complex that acts as a key regulator of mitosis (PubMed: <u>11516652</u> , PubMed: <u>12925766</u> , PubMed: <u>14610074</u> , PubMed: <u>14722118</u> , PubMed: <u>29449677</u>). The CPC complex has essential

functions at the centromere in ensuring correct chromosome alignment and segregation and is required for chromatin-induced microtubule stabilization and spindle assembly (PubMed: <u>11516652</u> , PubMed: <u>12925766</u> , PubMed: <u>14610074</u> , PubMed: <u>14722118</u> , PubMed: <u>26829474</u>). Involved in the bipolar attachment of spindle microtubules to kinetochores and is a key regulator for the onset of cytokinesis during mitosis (PubMed: <u>15249581</u>). Required for central/midzone spindle assembly and cleavage furrow formation (PubMed: <u>12458200</u> , PubMed: <u>12686604</u>). Key component of the cytokinesis checkpoint, a process required to delay abscission to prevent both premature resolution of intercellular chromosome bridges and accumulation of DNA damage: phosphorylates CHMP4C, leading to retain abscission-competent VPS4 (VPS4A and/or VPS4B) at the midbody ring until
abscission checkpoint signaling is terminated at late cytokinesis (PubMed: <u>22422861</u> , PubMed: <u>24814515</u>). AURKB phosphorylates the CPC complex subunits BIRC5/survivin, CDCA8/borealin and INCENP (PubMed: <u>11516652</u> , PubMed: <u>12925766</u> , PubMed: <u>14610074</u>). Phosphorylation
of INCENP leads to increased AURKB activity (PubMed: <u>11516652</u> , PubMed: <u>12925766</u> , PubMed: <u>14610074</u>). Other known AURKB substrates involved in centromeric functions and mitosis are CENPA, DES/desmin, GPAF, KIF2C, NSUN2, RACGAP1, SEPTIN1, VIM/vimentin, HASPIN, and histone H3 (PubMed: <u>11756469</u> , PubMed: <u>11784863</u> , PubMed: <u>11856369</u> ,
PubMed: <u>12689593</u> , PubMed: <u>14602875</u> , PubMed: <u>16103226</u> , PubMed: <u>21658950</u>). A positive feedback loop involving HASPIN and AURKB contributes to localization of CPC to centromeres (PubMed: <u>21658950</u>). Phosphorylation of VIM controls vimentin filament segregation in cytokinetic
process, whereas histone H3 is phosphorylated at 'Ser-10' and 'Ser-28' during mitosis (H3S10ph and H3S28ph, respectively) (PubMed: <u>11784863</u> , PubMed: <u>11856369</u>). AURKB is also required for kinetochore localization of BUB1 and SGO1 (PubMed: <u>15020684</u> , PubMed: <u>17617734</u>). Phosphorylation of p53/TP53 negatively regulates its transcriptional activity (PubMed: <u>20959462</u>). Key regulator of active promoters in resting B- and T-lymphocytes: acts by mediating phosphorylation of H3S28ph at active promoters in resting B-cells, inhibiting RNF2/RING1B-mediated ubiquitination of histone H2A and
enhancing binding and activity of the USP16 deubiquitinase at transcribed genes (By similarity). Acts as an inhibitor of CGAS during mitosis: catalyzes phosphorylation of the N-terminus of CGAS during the G2-M transition, blocking CGAS liquid phase separation and activation, and thereby preventing CGAS-induced autoimmunity (PubMed: <u>33542149</u>). Phosphorylates KRT5 during anaphase and telophase (By similarity). Phosphorylates ATXN10 which promotes phosphorylation of ATXN10 by PLK1 and may play a role in the regulation of cytokinesis and stimulating the proteasomal degradation of ATXN10 (PubMed: <u>25666058</u>).
Nucleus. Chromosome. Chromosome, centromere. Chromosome, centromere, kinetochore. Cytoplasm, cytoskeleton, spindle. Midbody. Note=Localizes on chromosome arms and inner centromeres from prophase through metaphase and then transferring to the spindle midzone and midbody from anaphase through cytokinesis (PubMed:20929775). Colocalized with gamma tubulin in the midbody (PubMed:17726514). Proper localization of the active, Thr-232- phosphorylated form during metaphase may be

Tissue Location

Cellular Location

High level expression seen in the thymus. It is also expressed in the spleen, lung, testis, colon, placenta and fetal liver. Expressed during S and G2/M phase and expression is up-regulated in cancer cells during M phase.

(and probably targeting of the CPC) to the inner centromere occurs predominantly in regions with overlapping mitosis-specific histone phosphorylations H3pT3 and H2ApT12 (PubMed:20929775).

dependent upon interaction with SPDYC (PubMed:20605920). Colocalized with SIRT2 during cytokinesis with the midbody (PubMed:17726514). Localization

Background

Recognizes a protein of 39kDa, which is identified as Aurora B. The serine/threonine protein kinase aurora B (Aurora B) is a chromosomal passenger protein critical for accurate chromosome segregation, cytokinesis, protein localization to the centromere and kinetochore, correct microtubule-kinetochore attachment, and regulation of the mitotic checkpoint. Aurora B forms a tight complex with inner centrosome protein and survivin. Inactivation of any of these proteins causes similar defects in chromosome segregation. A significant overexpression of Aurora B has been found in a variety of human tumors including non-small cell lung carcinoma, astrocytoma, seminoma and carcinomas of the colon, prostate, endometrium and thyroid. The expression level of Aurora B is associated with cell proliferation and prognosis in these tumors.

Images



Western Blot Analysis (A) Recombinant Protein (B) Human Liver Lysate Using Aurora B Monoclonal Antibody (AURKB/1521).

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