abcam

Product datasheet

Anti-Histone H2A.X antibody [EPR22820-23] - ChIP Grade - BSA and Azide free ab256544

敲除 验证 重组 RabMAb

9 图**像**

Anti-Histone H2A.X 抗体 [EPR22820-23] - ChIP Grade - BSA and Azide free		
兔单克隆抗体[EPR22820-23] to Histone H2A.X - ChIP Grade – BSA and Azide free		
Rabbit		
适用于: PepArr, ChIP, IHC-P, IP, WB, ICC/IF, Flow Cyt (Intra)		
与反应: Human		
Synthetic peptide. This information is proprietary to Abcam and/or its suppliers.		
WB: Wild-type HAP1 whole, Histone H2A.X knockout HAP1 whole, Human brain, HeLa, 293T and HEK-293 lysates. IHC-P: Human breast carcinoma and Human testis tissues. ICC/IF: HeLa cells. Flow Cyt (intra): HeLa cells. IP: HeLa cells.		
ab256544 is the carrier-free version of ab229914.		
Our <u>carrier-free</u> antibodies are typically supplied in a PBS-only formulation, purified and free of BSA, sodium azide and glycerol. The carrier-free buffer and high concentration allow for increased conjugation efficiency.		
This conjugation-ready format is designed for use with fluorochromes, metal isotopes, oligonucleotides, and enzymes, which makes them ideal for antibody labelling, functional and cell-based assays, flow-based assays (e.g. mass cytometry) and Multiplex Imaging applications.		
Use our <u>conjugation kits</u> for antibody conjugates that are ready-to-use in as little as 20 minutes with <1 minute hands-on-time and 100% antibody recovery: available for fluorescent dyes, HRP, biotin and gold.		
This product is compatible with the Maxpar [®] Antibody Labeling Kit from Fluidigm, without the need for antibody preparation. Maxpar [®] is a trademark of Fluidigm Canada Inc.		
 This product is a recombinant monoclonal antibody, which offers several advantages including: High batch-to-batch consistency and reproducibility Improved sensitivity and specificity Long-term security of supply Animal-free production For more information <u>see here</u>. Our RabMAb[®] technology is a patented hybridoma-based technology for making rabbit 		

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性能		
形式	Liquid	
存 放 说明	Shipped at 4°C. Store at +4°C. Do Not Freeze.	
存储溶液	pH: 7.2 Constituent: PBS	
无载体	是一一一一一一一一一一一一一一一一一一一一一一一一一一一一一一一一一一一一一一	
纯 度	Protein A purified	
克隆	单 克隆	
克隆 编号	EPR22820-23	
同种型	lgG	

应用

The Abpromise guarantee

Abpromise™承诺保证使用ab256544于以下的经测试应用

"应用说明"部分下显示的仅为推荐的起始稀释度;实际最佳的稀释度/浓度应由使用者检定。

应用	Ab评论	说明
PepArr		Use at an assay dependent concentration.
ChIP		Use at an assay dependent concentration.
IHC-P		Use at an assay dependent concentration.
IP		Use at an assay dependent concentration.
WB		Use at an assay dependent concentration. Predicted molecular weight: 15 kDa.
ICC/IF		Use at an assay dependent concentration.
Flow Cyt (Intra)		Use at an assay dependent concentration.

靶标

功能

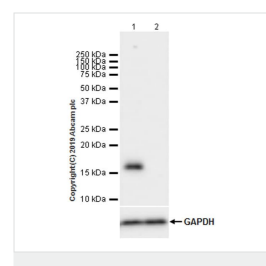
Variant histone H2A which replaces conventional H2A in a subset of nucleosomes. Nucleosomes wrap and compact DNA into chromatin, limiting DNA accessibility to the cellular machineries which require DNA as a template. Histones thereby play a central role in transcription regulation, DNA repair, DNA replication and chromosomal stability. DNA accessibility is regulated via a complex set of post-translational modifications of histones, also called histone code, and nucleosome remodeling. Required for checkpoint-mediated arrest of cell cycle progression in response to low doses of ionizing radiation and for efficient repair of DNA double strand breaks (DSBs) specifically when modified by C-terminal phosphorylation.

序列相似性Belongs to the histone H2A family.发展阶段Synthesized in G1 as well as in S-phase.结构域The [ST]-Q motif constitutes a recognition sequence for kinases from the PI3/PI4-kinase family.翻译后修饰Phosphorylated on Ser-140 (to form gamma-H2AFX or H2AX139ph) in response to DNA double
strand breaks (DSBs) generated by exogenous genotoxic agents and by stalled replication forks,
and may also occur during meiotic recombination events and immunoglobulin class switching in
lymphocytes. Phosphorylation can extend up to several thousand nucleosomes from the actual
site of the DSB and may mark the surrounding chromatin for recruitment of proteins required for

DNA damage signaling and repair. Widespread phosphorylation may also serve to amplify the damage signal or aid repair of persistent lesions. Phosphorylation of Ser-140 (H2AX139ph) in response to ionizing radiation is mediated by both ATM and PRKDC while defects in DNA replication induce Ser-140 phosphorylation (H2AX139ph) subsequent to activation of ATR and PRKDC. Dephosphorylation of Ser-140 by PP2A is required for DNA DSB repair. In meiosis, Ser-140 phosphorylation (H2AX139ph) may occur at synaptonemal complexes during leptotene as an ATM-dependent response to the formation of programmed DSBs by SPO11. Ser-140 phosphorylation (H2AX139ph) may subsequently occurs at unsynapsed regions of both autosomes and the XY bivalent during zygotene, downstream of ATR and BRCA1 activation. Ser-140 phosphorylation (H2AX139ph) may also be required for transcriptional repression of unsynapsed chromatin and meiotic sex chromosome inactivation (MSCI), whereby the X and Y chromosomes condense in pachytene to form the heterochromatic XY-body. During immunoglobulin class switch recombination in lymphocytes, Ser-140 phosphorylation (H2AX139ph) may occur at sites of DNA-recombination subsequent to activation of the activation-induced cytidine deaminase AICDA. Phosphorylation at Tyr-143 (H2AXY142ph) by BAZ1B/WSTF determines the relative recruitment of either DNA repair or pro-apoptotic factors. Phosphorylation at Tyr-143 (H2AXY142ph) favors the recruitment of APBB1/FE65 and proapoptosis factors such as MAPK8/JNK1, triggering apoptosis. In contrast, dephosphorylation of Tvr-143 by EYA proteins (EYA1, EYA2, EYA3 or EYA4) favors the recruitment of MDC1containing DNA repair complexes to the tail of phosphorylated Ser-140 (H2AX139ph). Monoubiquitination of Lys-120 (H2AXK119ub) by RING1 and RNF2/RING2 complex gives a specific tag for epigenetic transcriptional repression. Following DNA double-strand breaks (DSBs), it is ubiquitinated through 'Lys-63' linkage of ubiquitin moieties by the E2 ligase UBE2N and the E3 ligases RNF8 and RNF168, leading to the recruitment of repair proteins to sites of DNA damage. Monoubiguitination and ionizing radiation-induced 'Lys-63'-linked ubiguitination are distinct events.

细胞定位

Nucleus. Chromosome.



Western blot - Anti-Histone H2A.X antibody [EPR22820-23] - ChIP Grade - BSA and Azide free (ab256544) All lanes : Anti-Histone H2A.X antibody [EPR22820-23] - ChIP Grade (<u>ab229914</u>) at 1/1000 dilution

Lane 1 : Wild-type HAP1 whole cell lysate Lane 2 : Histone H2A.X knockout HAP1 whole cell lysate

Lysates/proteins at 20 µg per lane.

Secondary

All lanes : Goat Anti-Rabbit lgG H&L (HRP) (<u>ab97051</u>) at 1/20000 dilution

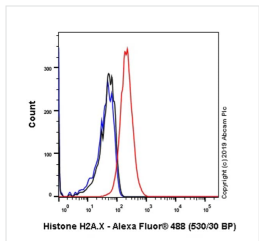
Predicted band size: 15 kDa Observed band size: 16 kDa

Ab229914 was shown to specifically react with Histone H2A.X in wild-type HAP1 cells as signal was lost in Histone H2A.X knockout cells. Wild-type and Histone H2A.X knockout samples were subjected to SDS-PAGE. **ab229914** and **ab181602** (Rabbit anti-GAPDH loading control) were incubated 1 hour at room temperature at 1/1000 dilution and 1/200,000 dilution respectively. Blots were developed with Goat Anti-Rabbit IgG, (H+L), Peroxidase conjugated (**ab97051**) secondary antibody at 1/100,000 dilution for 1 hour at room temperature before imaging. The blot was developed on a BIO-RAD® ChemiDocâ,¢ MP instrument using the ECL technique. Blocking/Diluting buffer and concentration: 5% NFDM/TBST Exposure Time: 37 seconds.

This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and sodium azide (<u>ab229914</u>).



Immunoprecipitation - Anti-Histone H2A.X antibody [EPR22820-23] - ChIP Grade - BSA and Azide free (ab256544)



Flow Cytometry (Intracellular) - Anti-Histone H2A.X antibody [EPR22820-23] - ChIP Grade - BSA and Azide free (ab256544) Histone H2A.X was immunoprecipitated from 0.35 mg HeLa (human cervix adenocarcinoma epithelial cell) whole cell lysate with <u>ab229914</u> at 1/30 dilution (2ug in 0.35mg lysates). Western blot was performed on the immunoprecipitate using <u>ab229914</u> at 1/1000 dilution. VeriBlot for IP Detection Reagent (HRP) (ab131366) was used at 1/5000 dilution.

Lane 1: HeLa (human cervix adenocarcinoma epithelial cell) whole cell lysate

Lane 2: ab229914 IP in HeLa whole cell lysate

Lane 3: Rabbit monoclonal lgG ($\underline{ab172730}$) instead of $\underline{ab229914}$ in HeLa whole cell lysate

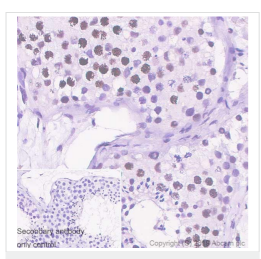
Blocking and dilution buffer and concentration/ 5% NFDM/TBST.

Exposure time/ 30 seconds.

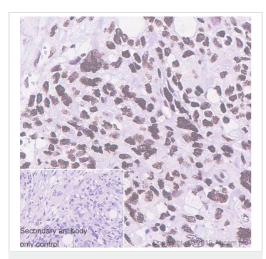
This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and sodium azide (**ab229914**).

Intracellular flow cytometric analysis of 4% paraformaldehyde fixed, 90% methanol permeabilized HeLa (human cervix adenocarcinoma epithelial cell) cells labelling Histone H2A.X with **ab229914** at 1/50 dilution (Red), compared with a Rabbit monoclonal IgG (**ab172730**) isotype control (Black)and an unlabelled control (cells without incubation with primary antibody and secondary antibody) (Blue). A Goat anti rabbit IgG (Alexa Fluor[®] 488, **ab150077**) at 1/2000 dilution was used as the secondary antibody.

This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and sodium azide (<u>ab229914</u>).



Immunohistochemistry (Formalin/PFA-fixed paraffinembedded sections) - Anti-Histone H2A.X antibody [EPR22820-23] - ChIP Grade - BSA and Azide free (ab256544)



Immunohistochemistry (Formalin/PFA-fixed paraffinembedded sections) - Anti-Histone H2A.X antibody [EPR22820-23] - ChIP Grade - BSA and Azide free (ab256544) Immunohistochemical analysis of paraffin-embedded Human testis tissue labeling Histone H2A.X with <u>ab229914</u> at 1/200 dilution (2.56 ug/ml) followed by a ready to use Rabbit specific IHC polymer detection kit HRP/DAB (<u>ab209101</u>). Nuclear staining on the human testis (PMID/ 24059746). The section was incubated with <u>ab229914</u> for 15 mins at room temperature. The immunostaining was performed on a Leica Biosystems BOND[®] RX instrument. Counterstained with Hematoxylin.

Secondary antibody only control: Secondary antibody is a ready to use Rabbit specific IHC polymer detection kit HRP/DAB (ab209101).

Heat mediated antigen retrieval with Tris-EDTA buffer (pH 9.0, epitope retrieval solution 2) for 20 mins.

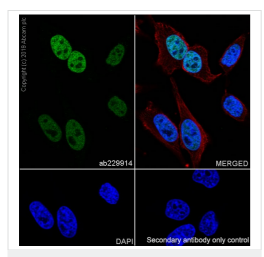
This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and sodium azide (<u>ab229914</u>).

Immunohistochemical analysis of paraffin-embedded Human breast carcinoma tissue labeling Histone H2A.X with <u>ab229914</u> at 1/200 dilution (2.56 ug/ml) followed by a ready to use Rabbit specific IHC polymer detection kit HRP/DAB (<u>ab209101</u>). Nuclear staining on the human breast carcinoma (PMID/ 27006338). The section was incubated with <u>ab229914</u> for 15 mins at room temperature. The immunostaining was performed on a Leica Biosystems BOND[®] RX instrument. Counterstained with Hematoxylin.

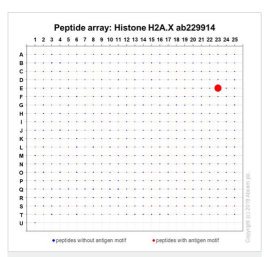
Secondary antibody only control: Secondary antibody is a ready to use Rabbit specific IHC polymer detection kit HRP/DAB (ab209101)

Heat mediated antigen retrieval with Tris-EDTA buffer (pH 9.0, epitope retrieval solution 2) for 20 mins.

This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and sodium azide (**ab229914**).



Immunocytochemistry/ Immunofluorescence - Anti-Histone H2A.X antibody [EPR22820-23] - ChIP Grade - BSA and Azide free (ab256544)



Peptide Array - Anti-Histone H2A.X antibody [EPR22820-23] - ChIP Grade - BSA and Azide free (ab256544) Immunofluorescent analysis of 4% Paraformaldehyde-fixed, 0.1% TritonX-100 permeabilized HeLa (human cervix adenocarcinoma epithelial cell) cells labelling Histone H2A.X with **ab229914** at 1/100 dilution, followed by Ab229914 anti- Histone H2A.X **ab150077** AlexaFluor[®]488 Goat anti-Rabbit secondary antibody at 1/1000 dilution (Green). Confocal image showing nuclear staining in HeLa cell line is observed. Ab195889 Anti-alpha Tubulin antibody [DM1A] - Microtubule Marker (Alexa Fluor[®] 594) was used to counterstain tubulin at 1/200 dilution (Red). The Nuclear counterstain was DAPI (Blue).

Secondary antibody only control: Secondary antibody is <u>ab150077</u> AlexaFluor[®]488 Goat anti-Rabbit secondary at 1/1000 dilution.

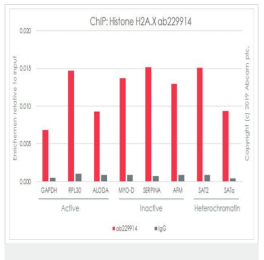
This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and sodium azide (<u>ab229914</u>).

All batches of **ab229914** are tested in Peptide Array against 501 different modified and unmodified histone peptides; each peptide is printed on the array at six concentrations (each in triplicate).

Circle area represents affinity between the antibody and a peptide: all antigen-containing peptides are displayed as red circles, all other peptides as blue circles. The affinity is calculated as area under curve when antibody binding values are plotted against the corresponding peptide concentration. Each circle area is normalized to the peptide with the strongest affinity.

The complete dataset, including full list of all peptides and information on the position of each peptide in the diagram, can be downloaded <u>here</u>.

This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and sodium azide (<u>ab229914</u>).



ChIP - Anti-Histone H2A.X antibody [EPR22820-23] -ChIP Grade - BSA and Azide free (ab256544)



Grade - BSA and Azide free (ab256544)

Please note: All products are "FOR RESEARCH USE ONLY. NOT FOR USE IN DIAGNOSTIC PROCEDURES"

Chromatin was prepared from HeLa cells according to the Abcam X-ChIP protocol*. Cells were fixed with formaldehyde for 10min.

The ChIP was performed with 25 μ g of chromatin, 5 μ g of **ab229914** (red), or 5 μ g of rabbit normal IgG **ab172730** (gray) and 20 μ I of Protein A/G sepharose beads. The immunoprecipitated DNA was quantified by real time PCR (Taqman approach for active and inactive loci, Sybr green approach for heterochromatic loci). Primers and probes are located in the first kb of the transcribed

This data was developed using the same antibody clone in a different buffer formulation containing PBS, BSA, glycerol, and sodium azide (<u>ab229914</u>).

region.

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