Acetyl-Histone H3.1 (K4) Recombinant Antibody



RACO0025

Reactivity:

Human

Human

Product Information

Size: Protein Background:

50ul Core component of nucleosome. 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

nucleocenne remodeling

Source: nucleosome remodeling.

Isotype: HIST1H3A

Rabbit IgG Uniprot

Applications: P68431

ELISA, WB, ICC, IF Synonyms:

Recommended dilutions: Histone H3/a, Histone H3/b, Histone H3/c, Histone H3/d, Histone H3/f,

Histone H3/h, Histone H3/j, Histone H3/j, Histone H3/k, Histone H3/l, HIST1H3A, H3FA, WB:1:500-1:2000, ICC:1:50-1:500, IF:1:30
1:200

Histone H3/h, Histone H3/j, Histone H3/k, Histone H3/l, HIST1H3A, H3FA, AND, HIST1H3C, H3FL, AND, H3FL, AND, H3FL, AND, H3FL, AND, H3FL, AND, H3FL, H3FL, AND, H3FL, H3FL,

H3FD, AND, HIST1H3F, H3FI, AND, HIST1H3G, H3FH, AND, HIST1H3H, H3FK, AND,

HIST1H3I, H3FF, AND, HIST1H3J, H3FJ

Immunogen:

A synthesized peptide.

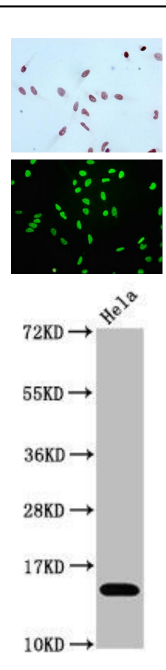
Storage:

Gene ID:

Rabbit IgG in phosphate buffered saline, pH 7.4, 150mM NaCl, 0.02% sodium azide and

50% glycerol.

Product Images



Immunocytochemistry analysis of RACO0025 diluted at 1:100 and staining in Hela cells performed on a Leica BondTM system. After dewaxing and hydration, antigen retrieval was mediated by high pressure in a citrate buffer (pH 6.0). Section was blocked with 10% normal goat serum 30min at RT. Then primary antibody (1% BSA) was incubated at 4°C overnight. The primary is detected by a biotinylated secondary antibody and visualized using an HRP conjugated SP system.

Immunofluorescence staining of Hela cells(treated by 15mM sodium butyrate for 30min) with RACO0025 at 1:68, counter-stained with DAPI. The cells were fixed in 4% formaldehyde, permeabilized using 0.2% Triton X-100 and blocked in 10% normal Goat Serum. The cells were then incubated with the antibody overnight at 4°C. The secondary antibody was Alexa Fluor 488-congugated AffiniPure Goat Anti-Rabbit IgG (H+L).

Western Blot

Positive WB detected in(Hela whole cell lysate)(treated by 15mM sodium butyrate for 30min)All lanes: Acetyl-Histone H3.1(K4)antibody at $1(1\mu q)ml$

Secondary

Goat polyclonal to rabbit IgG at 1:50000 dilution

Predicted band size: 15 KDa Observed band size: 15 KDa