

ab115127 – Histone H3 (phospho S10) Assay Kit (Colorimetric)



For the measurement of global histone H3 phosphorylation at ser10

This product is for research use only and is not intended for diagnostic use.

Table of Contents

INTI	RODUCTION	
1.	BACKGROUND	2
2.	ASSAY SUMMARY	3
GEN	NERAL INFORMATION	
	PRECAUTIONS	4
	STORAGE AND STABILITY	4
	MATERIALS SUPPLIED	5
	MATERIALS REQUIRED, NOT SUPPLIED	5
	LIMITATIONS	6
8.	TECHNICAL HINTS	6
ASS	SAY PREPARATION	
9.	REAGENT PREPARATION	7
10.	SAMPLE PREPARATION	8
ASS	SAY PROCEDURE	
11.	ASSAY PROCEDURE	10
DA1	TA ANALYSIS	
12.	ANALYSIS	11
RES	SOURCES	
13.	TROUBLESHOOTING	12
	NOTES	14

INTRODUCTION

1. BACKGROUND

The phosphorylation of histone H3 at serine 10 is conserved through eukaryotes, and an increase in phosphorylation has been shown to correlate with gene activation and cell growth. In vitro studies have shown that phosphorylation of histone H3 at ser10 is coupled to acetylation at the nearby Lysine-14 residue. Histone H3 phosphorylation at ser10 is also negatively impacted by histone methylation at lysine 9. It was observed that histone H3 phosphorylation at ser10 is regulated by the cell cycle and has been used as a mitotic marker. H3 phosphorylation (ser10) is critical for neoplastic cell transformation. Several protein kinases, including aurora B. PPI, and PKC, are responsible for histone H3 phosphorylation at ser10. Inhibition or activation of these protein kinases can cause the change in intracellular histone H3 phosphorylation at ser10. Detection in the change of histone H3 phosphorylation at ser10 associated with the cell cycle, apoptosis, inhibitor or activator treatment, would provide useful information for better understanding the pathological process of some diseases and for protein kinase-targeted drug development.

ab115127 provides a tool for measuring global phospho histone H3 (ser10).

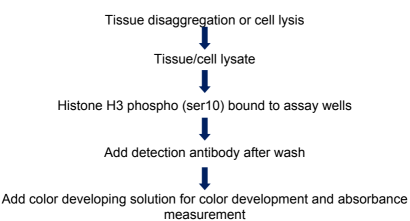
This kit has the following features:

- Quick and efficient procedure, which can be finished within 3 hours.
- Innovative colorimetric assay without the need for radioactivity, electrophoresis, or chromatography.
- Specifically captures phospho histone H3 (ser10) with the detection limit as low as 2 ng/well.
- The control is conveniently included for the quantification of phosphorylated histone H3 (ser10).
- Strip microplate format makes the assay flexible: manual or high throughput.
- Simple, reliable, and consistent assay conditions.

INTRODUCTION

The Histone H3 (phospho S10) Assay Kit (Colorimetric) is designed for measuring global histone H3 phosphorylation at ser10. In an assay with this kit, the phosphorylated histone H3 at ser10 is captured to the strip wells coated with an anti-phospho histone H3 (ser10) antibody. The captured phospho histone H3 (ser10) can then be detected with a labeled detection antibody, followed by a color development reagent. The ratio of phospho histone H3 (ser10) is proportional to the intensity of absorbance. The absolute amount of phospho histone H3 (ser10) can be quantitated by comparing to the standard control.

2. ASSAY SUMMARY



GENERAL INFORMATION

3. PRECAUTIONS

Please read these instructions carefully prior to beginning the assay.

All kit components have been formulated and quality control tested to function successfully as a kit. Modifications to the kit components or procedures may result in loss of performance.

4. STORAGE AND STABILITY

Store kit as given in the table upon receipt and away from light.

Observe the storage conditions for individual prepared components in sections 9 & 10.

For maximum recovery of the products, centrifuge the original vial prior to opening the cap.

Check if Wash and Antibody Buffers contain salt precipitates before use. If so, warm at room temperature or 37°C and shake the buffer until the salts are re-dissolved.

GENERAL INFORMATION

5. MATERIALS SUPPLIED

Item	Quantity (48 tests)	Quantity (96 tests)	Storage Condition (Before Preparation)
10X Wash Buffer	10 mL	20 mL	4°C
Antibody Buffer	6 mL	12 mL	4°C
Detection Antibody, 1 mg/mL*	5 μL	10 μL	-20°C
Color Developer	5 mL	10 mL	4°C
Stop Solution	3 mL	6 mL	RT
Standard Control (100 µg/mL)*	10 µL	20 μL	-20°C
8-Well Assay Strip (with Frame)	4	9	4°C
8-Well Standard Control Strips	2	3	4°C

^{*}Spin the solution down to the bottom after thawing and prior to use.

6. MATERIALS REQUIRED, NOT SUPPLIED

These materials are not included in the kit, but will be required to successfully utilize this assay:

- Orbital shaker
- Pipettes and pipette tips
- Reagent reservoir
- Microplate reader

GENERAL INFORMATION

7. LIMITATIONS

- Assay kit intended for research use only. Not for use in diagnostic procedures
- Do not use kit or components if it has exceeded the expiration date on the kit labels
- Do not mix or substitute reagents or materials from other kit lots or vendors. Kits are QC tested as a set of components and performance cannot be guaranteed if utilized separately or substituted
- Any variation in operator, pipetting technique, washing technique, incubation time or temperature, and kit age can cause variation in binding

8. TECHNICAL HINTS

- Avoid foaming or bubbles when mixing or reconstituting components.
- Avoid cross contamination of samples or reagents by changing tips between sample, standard and reagent additions.
- Ensure plates are properly sealed or covered during incubation steps.
- Complete removal of all solutions and buffers during wash steps.
- This kit is sold based on number of tests. A 'test' simply refers to a single assay well. The number of wells that contain sample, control or standard will vary by product. Review the protocol completely to confirm this kit meets your requirements. Please contact our Technical Support staff with any questions.

ASSAY PREPARATION

9. REAGENT PREPARATION

Prepare fresh reagents immediately prior to use.

9.1 1X Wash Buffer

Dilute 10X Wash Buffer with distilled water (pH 7.2-7.5) at a 1:10 ratio (1 mL of 10X Wash Buffer + 9 mL of water) to make 1X Wash Buffer.

9.2 Diluted Detection Antibody

Dilute Detection Antibody (at a 1:1000 ratio) to 1 $\mu g/mL$ with Antibody Buffer.

Suggested Plate Configuration

- Strip 1-3 (for 96 assays) or strip 1-2 (for 48 assays) standard wells (labeled as SC); the standard curve can be generated with 5-8 concentration points (includes blank)
- Example amount of standard control/well A1: 100 ng; B1: 50 ng;
 C1: 25 ng; D1: 12 ng; E1: 6 ng; F1: 3 ng; G1: 1.5 ng; H1: 0 ng
- Strip 4-12 (for 96 assays) or strip 3-6 (for 48 assays) sample wells (No label)
- Each sample or standard point can be assayed in duplicates or triplicates

ASSAY PREPARATION

10. SAMPLE PREPARATION

You may prepare histone extracts from cells/tissues treated or untreated by using your own successful method (acid extraction or high salt extraction). For your convenience and the best results, Abcam offers the Histone Extraction Kit (ab113476) optimized for use in the modified histone quantification series or alternatively, use the following protocol.

- 10.1 For tissues (treated and untreated): weigh the sample and cut the sample into small pieces (1-2 mm³) with a scalpel or scissors.
- 10.2 Transfer tissue pieces to a Dounce homogenizer.
- 10.3 Add TEB buffer (PBS containing 0.5% Triton X 100, 2 mM PMSF and 0.02% NaN₃) at 200 mg/mL, and disaggregate tissue pieces by 50-60 strokes.
- 10.4 Transfer homogenized mixture to a 15 mL conical tube and centrifuge at 3000 rpm for 5 minutes at 4°C. If total mixture volume is less than 2 mL, transfer mixture to a 2 mL vial and centrifuge at 10,000 rpm for 1 minute at 4°C.
- 10.5 Remove supernatant.
- 10.6 **For cells (treated and untreated):** harvest cells and pellet the cells by centrifugation at 1000 rpm for 5 minutes at 4°C.
- 10.7 Resuspend cells in TEB buffer at 10⁷ cells/mL and lyse cells on ice for 10 minutes with gentle stirring.
- 10.8 Centrifuge at 3000 rpm for 5 minutes at 4°C. If total volume is less than 2 mL, transfer cell lysates to a 2 mL vial and centrifuge at 10,000 rpm for 1 minute at 4°C.
- 10.9 Remove supernatant.
- 10.10 For both tissue and cells: Resuspend cell/tissue pellet in 3 volumes (approx. $200 \,\mu\text{L}/10^7$ cells or $200 \,\text{mg}$ tissues) of

ASSAY PREPARATION

- extraction buffer (0.5N HCl + 10% glycerol) and incubate on ice for 30 minutes.
- 10.11 Centrifuge at 12,000 rpm for 5 minutes at 4°C and remove the supernatant fraction to a new vial.
- 10.12 Add 8 volumes (approx. 0.6 mL/10⁷ cells or 200 mg tissues) of acetone and leave at –20°C overnight.
- 10.13 Centrifuge at 12,000 rpm for 5 minutes and air-dry the pellet. Dissolve the pellet in distilled water (30-50 μ L/10⁷ cells or 200 mg tissues).
- 10.14 Quantify the protein concentration. Aliquot the extract and store the extract at -20°C or -80°C.

Histone extracts can be used immediately or stored at-80°C for future use.

ASSAY PROCEDURE

11. ASSAY PROCEDURE

- 11.1 Determine the number of strip wells required. Leave these strips in the plate frame (remaining unused strips can be placed back in the bag. Seal the bag tightly and store at 4°C).
- 11.2 Add 50 μL of Antibody Buffer into each well. For the sample, add 1-2 μg of the histone extract into the sample wells. For the standard curve, dilute Standard Control with Antibody Buffer to 1-100 ng/μL at 5-7 points (e.g., 1.5, 3, 6, 12, 25, 50, and 100 ng/μL). Add 1 μL of Standard Control at the different concentrations into the Standard Control Wells. For the blank, do not add any nuclear extracts or standard control protein. Mix and cover the strip wells with Parafilm M and incubate at room temperature for 1-2 hours.
- 11.3 Aspirate and wash the wells with 150 μ L of 1X Wash Buffer three times.
- 11.4 Add 50 µL of diluted Detection Antibody to each well and incubate at room temperature for 60 minutes on an orbital shaker (100 rpm).
- 11.5 Aspirate and wash the wells with 150 μL of 1X Wash Buffer six times.
- 11.6 Add 100 µL of Color Developer into the wells and incubate at room temperature for 2-10 minutes away from light. Monitor the color development in the sample and standard wells (blue).
- 11.7 Add 50 µL of Stop Solution to each well to stop enzyme reaction when the color in the standard wells containing the higher concentrations of standard control turns medium blue. The color should change to yellow and absorbance can be read on a microplate reader at 450 nm within 1-15 minutes.

DATA ANALYSIS

12. ANALYSIS

Calculate % histone H3 phospho (ser10):

Phospho (ser10)
$$\frac{\% = \text{Treated (tested) sample OD} - \text{blank OD}}{\text{Untreated (control) sample OD} \times 100\%}$$

For the amount quantification plot OD versus amount of Standard Control and determine the slope as delta OD/ng.

Calculate the amount of phospho (ser10) using the following formula:

Amount (ng/mg protein) =
$$\frac{\text{Sample OD - blank OD}}{\text{Protein } (\mu g)^* \times \text{Slope}} \times 1000$$

*Histone extract amount added into the sample well at step 11.2.

RESOURCES

13. TROUBLESHOOTING

Problem	Cause	Solution
No Signal for Both the Standard Control and the Samples	Reagents are added incorrectly.	Check if reagents are added in order and if some steps of the procedure are omitted by mistake.
	Incubation time and temperature is incorrect.	Ensure the incubation time and temperature described in the protocol is followed correctly.
No Signal or Very Weak Signal for Only the Standard Control	The amount of Standard control is not added into the "standard control wells or is added insufficiently.	Ensure a sufficient amount of control is added to the well.
No Signal for Only the Sample	The protein sample is not properly extracted.	Ensure the procedure and reagents are correct for the nuclear protein extraction.
	The protein amount is added into well insufficiently.	Ensure extract contains a sufficient amount of protein.
	Protein extracts are incorrectly stored.	Ensure the protein extracts are stored at –20°C or –80°C.

RESOURCES

Problem	Cause	Solution
High Background Present for the Blank	The well is not washed sufficiently.	Check if wash at each step is performed according to the protocol.
	Contaminated by the Standard control	Ensure the well is not contaminated from adding the control protein or by using control protein contaminated tips.
	Overdevelopment	Decrease development time in step 11.6.

RESOURCES

14. <u>NOTES</u>



UK, EU and ROW

Email: technical@abcam.com | Tel: +44-(0)1223-696000

Austria

Email: wissenschaftlicherdienst@abcam.com | Tel: 019-288-259

France

Email: supportscientifique@abcam.com | Tel: 01-46-94-62-96

Germany

Email: wissenschaftlicherdienst@abcam.com | Tel: 030-896-779-154

Spain

Email: soportecientifico@abcam.com | Tel: 911-146-554

Switzerland

Email: technical@abcam.com

Tel (Deutsch): 0435-016-424 | Tel (Français): 0615-000-530

US and Latin America

Email: us.technical@abcam.com | Tel: 888-77-ABCAM (22226)

Canada

Email: ca.technical@abcam.com | Tel: 877-749-8807

China and Asia Pacific

Email: hk.technical@abcam.com | Tel: 400 921 0189 / +86 21 2070 0500

Japan

Email: technical@abcam.co.jp | Tel: +81-(0)3-6231-0940

www.abcam.com | www.abcam.cn | www.abcam.co.jp

Copyright © 2014 Abcam, All Rights Reserved. The Abcam logo is a registered trademark. All information / detail is correct at time of going to print.