



Base Catalog # P-3016

PLEASE READ THIS ENTIRE USER GUIDE BEFORE USE

The EpiQuik[™] In Situ Histone H3-K9 Methylation Assay Kit is suitable for specifically measuring histone H3-K9 methylation in situ using cultured adherent cells.











KIT CONTENTS

Components	96 assays P-3016-096	2 x 96 assays P-3016-192
GB1 (10X Wash Buffer) GB2 (Permeabilizing Buffer) GB3 (Blocking Buffer) GB4 (Antibody Buffer) GB5 (Capture Antibody, 100 μg/ml)* GB6 (Detection Antibody, 400 μg/ml)* GB7 (Developing Solution) GB8 (Stop Solution) 30% H ₂ O ₂ Solution Methylated H3-K9 Control (20 μg/ml) 8-Well Control Strips Microplates User Guide	30 ml 30 ml 20 ml 15 ml 60 μl 20 μl 12 ml 6 ml 0.5 ml 15 μl 2 1	2 x 30 ml 2 x 30 ml 2 x 20 ml 20 ml 120 µl 40 µl 24 ml 12 ml 1 ml 30 µl 4 2

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

SHIPPING & STORAGE

The kit is shipped in two parts, one part at ambient room temperature, and the second part on frozen ice packs at 4°C.

Upon receipt: (1) Store **GB6** and the **Methylated H3-K9 Control** at -20° C; (2) Store **GB8** at room temperature away from light; (3) Store **all other components** at 4° C away from light. The kit is stable for up to 6 months from the shipment date, when stored properly.

Note: Check if buffers, **GB1** and **GB4**, contain salt precipitates before using. If so, warm (at room temperature or 37°C) and shake the buffers until the salts are re-dissolved.

GENERAL PRODUCT INFORMATION

Usage Limitation: The EpiQuik[™] In Situ Histone H3-K9 Methylation Assay Kit is for research use only and is not intended for diagnostic or therapeutic application.

Quality Control: Epigentek guarantees the performance of all products in the manner described in our product instructions.

Product Updates: Epigentek reserves the right to change or modify any product to enhance its performance and design.

Intellectual Property: EpiQuik[™] is a trademark of Epigentek Group Inc.



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A BRIEF OVERVIEW

Epigenetic activation or inactivation of genes plays a critical role in many important human diseases, especially in cancer. A major mechanism for epigenetic inactivation of the genes is methylation of CpG islands in genome DNA caused by DNA methyltransferases. Histone methyltransferases (HMTs) control or regulate DNA methylation through chromatin-dependent transcription repression or activation. HMTs transfer 1-3 methyl groups from S-adenosyl-L-methionine to the lysine and arginine residues of histone proteins. ESET, G9a, SUV39-h1, SUV39-h2, SETDB1, Dim-5, and Eu-HMTase are histone methyltransferases that catalyze methylation of histone H3 at lysine 9 (H3-K9) in mammalian cells. H3-K9 methylation mediates heterochromatin formation by forming a binding site for HP1 and also participates in silencing gene expression at euchromatic sites. Increased global H3-K9 methylation is also found to be involved in some pathological processes such as cancer progress. The *EpiQuik™ In Situ* Histone H3-K9 Methylation Assay Kit provides a useful tool for measuring *in situ* histone H3-K9 methylation and 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.
- Measurement of *in situ* histone H3-K9 methylation without the need to prepare cell lysates.
- Microplate format makes the assay suitable for high throughput analysis of agents that increases or inhibit H3-K9 methylation.
- Simple, reliable, and consistent assay conditions.

PRINCIPLE & PROCEDURE

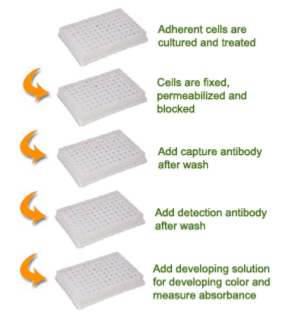
The *EpiQuik*[™] In Situ Histone H3-K9 Methylation Assay Kit is a whole cell-based detection of methylated H3-K9. In this assay, adherent cells are cultured in conventional 96-well microplates. After your experimental treatment, cells are fixed and permeabilized. The methylated H3-K9 is then detected by a high affinity H3-K9 antibody. The ratio or amount of methylated H3-K9 can be quantified through HRP conjugated secondary antibody-color development system and is proportional to the intensity of color development.











Schematic Procedure for Using the EpiQuik™ In Situ Histone H3-K9 Methylation Assay Kit

PROTOCOL

Before starting, perform the following:

- (A) Prepare the following required solution (not included): 37% Formaldehyde.
- (B) Ensure that all buffer solutions are clear in appearance. Shake or vortex if these buffers have precipitates.
- 1. Grow adherent cells in the 96-well microplate to 50-60% confluency. Treat cells with the appropriate amount of reagents that may increase or reduce H3-K9 methylation for the appropriate time, based on your experiment design.
- 2. Prepare the **Fixing Solution** by adding 2.16 ml of 37% formaldehyde to 18 ml of PBS. Remove culture media from the wells with a wrist-flick.
- 3. Immediately add 150 μ l of the **Fixing Solution** slowly to the wells and incubate at room temperature for 15 minutes. Remove the **Fixing Solution** from the wells with a wrist-flick; while still inverted, tap the plate gently onto absorbent paper to remove any excess fixing reagent still within the wells.
- 4. Dilute **GB1** with distilled water (pH 7.2-7.5) at a 1:10 ratio (e.g., 1 ml of **GB1** + 9 ml of distilled water). Wash the wells once (2 minutes) with $150 \,\mu$ l of the **diluted GB1**.
- 5. Remove the wash buffer with a wrist flick; while still inverted, tap the plate onto absorbent paper to remove any excess solution. Add 150 μ l of **GB2** to each well and incubate at room temperature for 5 minutes. Meanwhile, prepare 1% H₂O₂ Solution by adding 330 μ l of 30% H₂O₂ Solution into 10 ml of **GB2**.
- 6. Remove **GB2** from wells with a wrist flick and add 100 μ l of the **1%** H₂O₂ Solution into each well and incubate at room temperature for 10 minutes to remove endogenous peroxidase.



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- 7. Remove the $1\% H_2O_2$ Solution from the wells with a wrist flick and wash the wells twice with 150 μ l of the diluted GB1.
- 8. Remove the wash buffer with a wrist flick; while still inverted, tap the plate onto absorbent paper to remove any excess solution. Add 150 μ l of **GB3** to the wells and incubate at 37°C for 45 minutes. Meanwhile, add 50 μ l of **diluted GB1** to the desired number of control strip wells, followed by adding 1 μ l of *methylated H3-K9 control protein* at the different amounts (ex: 0.5-20 ng, diluted with distilled water) and incubate at room temperature for 30-45 minutes. For the blank wells, do not add any *methylated H3-K9 control protein*.
- 9. Remove GB3 with a wrist flick; while still inverted, tap the plate onto absorbent paper to remove any excess solution. Wash the wells twice with 150 μl of diluted GB1. For each wash, remove the diluted GB1 with a wrist flick; while still inverted, tap the plate onto absorbent paper. Meanwhile, aspirate the solution from the control strip wells and wash the wells with 150 μl diluted GB1 three times.
- 10. Dilute GB5 (at a 1:100 ratio) to 1 μg/ml with GB4. Add 50 μl of the diluted GB5 to the sample wells and the Methylated H3-K9 Control Strip Wells. Incubate at room temperature for 60 minutes on an orbital shaker (50-100 rpm).
- 11. Remove solution from the wells with a wrist flick and wash the wells four times with 150 µl of diluted GB1. For each wash, remove the diluted GB1 with a wrist flick; while still inverted, tap the plate onto absorbent paper.
- 12. Dilute the **GB6** (at a 1:1000 ratio) to 0.4 μ g/ml with **GB4**. Add 50 μ l of the **diluted GB6** to the wells and incubate at room temperature for 30 minutes.
- Remove solution from the wells with a wrist flick and wash the wells four times with 150 μl of diluted GB1. For each wash, remove the diluted GB1 with a wrist flick; while still inverted, tap the plate onto absorbent paper.
- 14. Add 100 μ l of **GB7** to the wells and incubate at room temperature for 2-10 minutes away from light. Monitor the color development in the sample and control wells until you observe a medium blue color.
- 15. Add 50 μ l of **GB8** to the wells and read absorbance on a microplate reader at 450 nm.
- 16. Calculate % H3-K9 methylation using the following formula:

Methylation % = $\frac{\text{O.D. (treated sample - blank)}}{\text{O.D. (untreated control - blank)}} \times 100\%$

TROUBLESHOOTING

No Signal for Both the Positive Control and the Samples

Reagents are added incorrectly.

Check if reagents are added in proper order and if any steps of the procedure may have been omitted by mistake.

Ensure the incubation time and temperature described in the protocol are followed correctly.



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Incubation time and temperature are incorrect.



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No Signal for Only the Sample

Cells are not fixed and	Ensure fixation solution and permeabilizing
permeabilized sufficiently.	solution are sufficiently added into the cells and that
	the incubation time is sufficient.

High Background Present for the Blank

The wells are not washed sufficiently.

Check if wash at each step is performed according to the protocol.

Overdevelopment.

Decrease development time in Step 14.

RELATED PRODUCTS

P-3001	EpiQuik™ DNA Methyltransferase Activity/Inhibition Assay Kit
P-3002	EpiQuik [™] Histone Methyltransferase Activity/Inhibition Assay Kit (H3-K4)
P-3003	EpiQuik [™] Histone Methyltransferase Activity/Inhibition Assay Kit (H3-K9)
P-3015	EpiQuik™ In Situ Histone H3-K4 Methylation Assay Kit
P-3017	EpiQuik™ Global Histone H3-K4 Methylation Assay Kit
P-3018	EpiQuik™ Global Histone H3-K9 Methylation Assay Kit
P-3019	EpiQuik™ DNA Demethylase Activity/Inhibition Assay Kit
P-3020	EpiQuik™ Global Histone H3-K27 Methylation Assay Kit



