

Xite™ Red beta-D-galactopyranoside

Catalog number: 14035 Unit size: 1 mg

Component	Storage	Amount
Xite™ Red beta-D-galactopyranoside	Freeze (< -15 °C), Minimize light exposure	1 mg

OVERVIEW

XiteTM Red beta-D-galactopyranoside provides a simple and sensitive tool to detect beta-galactosidase (β-gal) activity. Compared to the existing red beta-galactosidase substrates (e.g., the commonly used resorufin beta-D-galactopyranoside), it has much better cell permeability. XiteTM Red beta-D-galactopyranoside provides a simple and sensitive tool to detect beta-galactosidase activity. XiteTM Red beta-D-galactopyranoside might be used as a simple tool for measuring cellular senescence in cells since β-gal has been identified as a reliable marker for cellular senescence. XiteTM Red beta-D-galactopyranoside enters readily cells where it gets cleaved by β-gal, producing XiteTM Red, a strongly fluorescent product. The strongly fluorescent XiteTM Red is well retained in cells, making it easy to be detected with a flow cytometer and fluorescence microscope. In addition, XiteTM Red beta-D-galactopyranoside is fixable. The red fluorescence generated by XiteTM Red beta-D-galactopyranoside can be readily combined with other color fluorescent probes such as DAPI or GFP for multicolor fluorescence analysis.

AT A GLANCE

Protocol summary

- 1. Treat samples as desired.
- Prepare and add Xite™ Red beta-D-galactopyranoside working solution to samples
- 3. Incubate samples at 37 °C for 15 to 45 minutes
- Monitor the fluorescence intensity using flow cytometer with 575/26 nm filter (PE channel) or using fluorescence microscopy with Cy3/TRITC filter set

KEY PARAMETERS

Flow cytometer

Excitation 488 nm laser
Emission 575/26 nm filter
Instrument specification(s) PE channel

Fluorescence microscope

Excitation Cy3/TRITC filter set
Emission Cy3/TRITC filter set
Recommended plate Black wall/clear bottom

PREPARATION OF STOCK SOLUTIONS

Unless otherwise noted, all unused stock solutions should be divided into single-use aliquots and stored at -20 °C after preparation. Avoid repeated freeze-thaw cycles.

Xite™ Red beta-D-galactopyranoside stock solution

Add appropriate amount of DMSO into Xite™ Red beta-D-galactopyranoside to make 2-5 mM Xite™ Red beta-D-galactopyranoside stock solution.

Note Store the unused Xite $^{\text{TM}}$ Red beta-D-galactopyranoside stock solution at -20 °C in single use aliquots.

PREPARATION OF WORKING SOLUTION

Xite™ Red beta-D-galactopyranoside working solution

Prepare 1-20 μM of XiteTM Red beta-D-galactopyranoside working solution in buffer of your choice.

Note Xite™ Red beta-D-galactopyranoside working solution should be used promptly.

Note The concentration of the Xite™ Red beta-D-galactopyranoside should be optimized for different cell types and conditions.

SAMPLE EXPERIMENTAL PROTOCOL

The following protocol can be used as a guideline and should be optimized according to the needs.

- 1. Treat your samples as desired.
- Remove the treatment and wash cells with buffer of your choice such as DPBS.
- Add Xite[™] Red beta-D-galactopyranoside working solution for 15-45 minutes and incubate the samples at 37 °C incubator.

Note Optimal time for incubation needs to be determined experimentally.

- Remove the working solution and wash cells with buffer of your choice
- Resuspend the cells in buffer of your choice and monitor the fluorescence intensity with flow cytometer using 575/26 nm filter (PE channel) or fluorescence microscope with Cy3/TRITC filter set.

EXAMPLE DATA ANALYSIS AND FIGURES

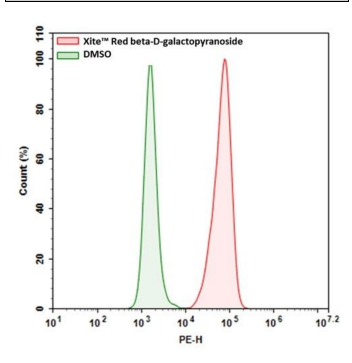


Figure 1. Expression of β-gal was measured with XiteTM Red beta-D-galactopyranoside. 9L-LacZ cells (cells that overexpressed β-gal) were incubated with XiteTM Red beta-D-galactopyranoside for 30 mins at 37 °C. The signal was acquired with PE channel using a NovoCyte Flow Cytometer (ACEA Biosciences).

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