Tel: 609-228-6898

Email: tech@MedChemExpress.com



Lyso Red Kit

1 Components

Component	HY-KD1028-100T
Lyso Red	$1~\text{mM} \times 50~\mu\text{L}$

2 Introduction

Lyso Red is a fluorescent probe with red fluorescence for specific labeling of lysosomes in living mammalian cells, with strong water solubility and pH stability; it has a maximum excitation wavelength of 564 nm and a maximum emission wavelength of 590 nm. The MCE Lyso Red kit enables the specific labeling of lysosomes in living cells with good labeling and anti-bleaching ability.

3 Characteristics

Maximum excitation wavelength: 564 nm Maximum emission wavelength: 590 nm

Color: red

Subcellular structural localization: lysosome

4 Self-contained reagents

- 1. Serum-free cell culture medium
- 2. PBS buffer

5 Protocol

Preparation of Lyso Red working solution

1. The working concentration of Lyso Red is as follows:

Component	Working Concentration
Lyso Red	5-10 μM

2. Serum-free cell culture medium is required for the preparation of the working solution. For 24- and 48-well plates, the amount of Lyso Red working solution per well is 200 μ L and 100 μ L, respectively; for 15 mm and 20 mm confocal imaging dishes, the amount of Lyso Red working solution is 100 μ L and 200 μ L, respectively. The following table can be used to prepare Lyso Red working solution.

Working Solution	Lyso Green	Serum-free medium
100 μL	1 μL	99 μL
200 μL	2 μL	198 μL

Note: The working solution should cover the cells completely when staining, and the amount of working solution can be adjusted proportionally according to the specific situation.

Lyso Red Incubation Procedure

- 1. Prepare the cells.
- 2. Prepare the working solution for incubation.
- 3. Wash the cells growing in the well plate or confocal dish with appropriate amount of PBS, and then take appropriate amount of serum-free medium to rinse the cell surface once.
- 4. Add the working solution and incubate the cells with the working solution for 1 h.
- 5. Take the appropriate amount of PBS and wash 3 times, add the appropriate amount of serum-free medium and leave for 15 min.
- 6. Take appropriate amount of PBS and wash 3 times, replace with serum cell culture medium and leave for 15 min.
- 7. Imaging.



-20°C, 1 year

Keep away from light

7 Precautions

- 1. In order to ensure cell activity and labeling effect, the cell confluence should reach 70%-90% before incubation.
- 2. The optimal incubation environment for the probe is 37°C, 5% CO₂ cell culture incubator.
- 3. Fluorescent dyes are subject to fluorescence quenching, so please image as soon as possible after incubation and rinsing.
- 4. This product is limited to scientific research by professionals and should not be used for clinical diagnosis or treatment, food or medicine.
- 5. For your safety and health, please wear lab coat and disposable gloves.

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