

## **Product Description**

Pioneering GTPase and Oncogene Product Development since 2010

## **DII-LDL**

产品名称: Dil-LDL

货

号: 10459

货号: 0.5 mg

纯化: 98% (Co-migrates with reference on agarose gel

electrophoresis)

Concentration: Minimum 1.6 mg/ml 蛋白

描述: Human Dil-Labeled Low Density Lipo蛋白

Purified LDL is labeled with the fluorescent probe, Dil, and reisolated by ultracentrifugation (1.019-1.063). The resultant

product is exhaustively dialyzed against phosphate

背景: buffered saline, (pH 7.4), sterilized by membrane filtration

and then aseptically packaged in a solution containing phosphate-buffered saline at pH 7.4 and 0.2 mM EDTA. Each

lot is evaluated on a murine macrophage cell line for

fluorescence uptake.

The labelled LDL is stable for 6 weeks after receipt when handled aseptically and stored at 2-8°C (**Don't Freeze**).

储存和运输: Note: After prolonged storage, some precipitate may be

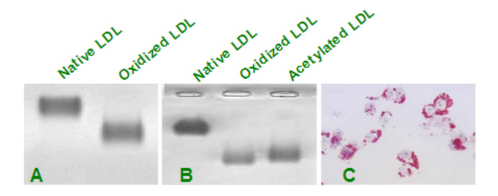
observed. This is normal for the product. Spin in

centrifugation at 5000×g for 10 minutes before using.

Packaging:

The labelled LDL requires one week lead time. Please plan

your experiments in advance and use the fresh material.



Native-LDL(n-LDL), Oxidized-LDL (ox-LDL) and -LDL(Ac-LDL) were loaded on agarose gel and electrophoresed for 60 mins. The lipoproteins were stained with Sudan Black (A and B). Oil red O staining was used to determine the formation of



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## Typical Lipoprotein Labeling Protocol

- 1. Dilute Dil-LDL to 10-40 ug/ml in growth media.
- 2. Add to cells and incubate for 2-6 hours at 37°C.
- 3. Remove media containing Dil-LDL from your culture.
- 4. Wash 3 times with probe-free media.

**A. Fluorescence Microscopy:**Visualize using standard rhodamine excitation: emission filters (or suggested wavelengths excitation:emission at 554nm:571nm or near). If fixation is desired use 3% formaldehyde in PBS. (Never use methanol or acetone fixation - Dil is soluble in organic solvents). Note: A positive culture must be stained for comparison purposes.

**A. Cell Sorting:**Label as in steps 1–5. Trypsinize or treat cultures with EDTA to produce a single cell suspension. Use labeled pure cultures of positive and negative cell types to set gates on the cell sorter. Suggested Wavelengths for Cell Sorting:Excitation: 554nmEmission: 571nm