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PEROXsayTM-LIPID

A lipid compatible quantitative peroxide assay

INTRODUCTION

PEROXsayTM-*LIPID* is a colorimetric quantitative peroxide assay that measures the oxidation of ferrous (Fe²⁺) ions to ferric (Fe³⁺) ions in the presence of lipids. Basically, the peroxides oxidize ferrous ions to ferric ions. The acidic pH of the PEROXsayTM Component 2 allows the ferric (Fe³⁺) ion to complex with xylenol orange, a constituent of PEROXsayTM-*LIPID* Component 1, resulting in a change in absorbance that is proportional to the peroxide concentration.

The PEROXsay[™]-*LIPID* is suitable for the following applications; measurement of lipid peroxidation of low density lipoproteins and liposomes, quantifying level of protein damaging peroxides in detergents, and monitoring protein glycation. The PEROXsay[™] assay is designed for microtiter plates, but can be scaled up for use with 1ml cuvettes.

ITEM(S) SUPPLIED		Cat# 786-441
	PEROXsay [™] -LIPID Component 1	50ml
	PEROXsay [™] Component 2	0.5ml

STORAGE CONDITIONS

The kit is shipped at ambient temperature. Upon arrival, store the kit at 4°C, when stored properly the kit is stable for 1 year.

ITEMS NEEDED BUT NOT SUPPLIED WITH KIT

30% Hydrogen peroxide solution (8.8M)

PREPARATION BEFORE USE:

Assay Solution:

For microtiter plate assays, you require 200µl Assay Solution for each sample and for cuvettes you will require 1ml Assay Solution.

Add 1 volume of PEROXsayTM Component 2 to 100 volumes PEROXsayTM-*LIPID* Component 1 and mix. The Assay Solution must be made fresh on the day of the assay.

Hydrogen Peroxide Standards:

- 1. Add 5µl 30% Hydrogen Peroxide solution to 440ml deionized (DI) water to give a 100µM concentration.
- 2. Serially dilute the 100μM hydrogen peroxide solution four times to give hydrogen peroxide standards of 6.25, 12.5, 25 and 50μM.

NOTE: To standardize the starting 30% hydrogen peroxide solution, use the molar coefficient of 43.6 M⁻¹cm⁻¹ for hydrogen peroxide at 240nm.

PROTOCOL

NOTE:

- A. The linear range for this assay is 0-50µM. Dilute samples with higher peroxide concentrations. In addition, samples with >1mM peroxide may cause bleaching and low absorbance reading, to alleviate this issue assay a 1:100 dilution in parallel.
- B. For samples that may have chelating proteins, transition metals or strong absorbance at or near 560nm, use a blank of PEROXsay[™]-LIPID Component 1 without PEROXsay[™] Component 2. Subtract this blank from the assayed sample to control for the above interferences.



- 1. For each volume of sample, add 10 volumes of Assay Solution.

 For a microtiter plate, add 200µl Assay Solution to each well containing 20µl sample.
- 2. Mix and then incubate at room temperature for 30 minutes.
- 3. After incubation, measure the absorbance at 560nm. *Absorbances can be read at 560-600nm, for plate readers use 595nm*
- 4. Plot a standard curve using the absorbances of the hydrogen peroxide samples and calculate the concentration of peroxides in your sample.

RELATED PRODUCTS

- 1. <u>Proteomic Grade Detergents:</u> A selection of non-ionic detergents are available that have ultra low levels of protein damaging peroxides and aldehydes.
- 2. $\underline{PEROXsay}^{\text{TM}}$ (786-440): A more sensitive peroxide assay that requires extraction of lipids.

<u>NOTE</u>: For other related products, visit our web site at <u>www.GBiosciences.com</u> or contact us.