Amplite™ Fluorimetric Hydrogen Peroxide Assay Kit

Red Fluorescence

Ordering Information	Storage Conditions	Instrument Platform
Product Number: 11501 (500 assays)	Keep in freezer Avoid exposure to light	Fluorescence microplate readers

Introduction

Hydrogen peroxide (H_2O_2) is a reactive oxygen metabolic by-product that serves as a key regulator for a number of oxidative stress-related states. It is involved in a number of biological events that have been linked to asthma, atherosclerosis, diabetic vasculopathy, osteoporosis, a number of neurodegenerative diseases and Down's syndrome. Perhaps the most intriguing aspect of H_2O_2 biology is the recent report that antibodies have the capacity to convert molecular oxygen into hydrogen peroxide to contribute to the normal recognition and destruction processes of the immune system. Measurement of this reactive species will help to determine how oxidative stress modulates a variety of intracellular pathways.

This AmpliteTM Fluorimetric Hydrogen Peroxide Assay Kit uses our non-fluorescent AmpliteTM Red peroxidase substrate to quantify hydrogen peroxide in solutions and cell extracts. It can also be used to detect a variety of oxidase activities through enzyme-coupled reactions. The kit is an optimized "mix and read" assay that is compatible with HTS liquid handling instruments. It provides a sensitive, one-step fluorometric assay to detect as little as 3 picomoles of H_2O_2 in a 100 μ L assay volume (30 nM, Figure 1). The assay can be performed in a convenient 96-well or 384-well microtiter-plate format and readily adapted to automation. Its signal can be easily read by either a fluorescence microplate reader at Ex/Em = ~540/590 nm or an absorbance microplate reader at ~570 nm.

Kit Key Features

Broad Application: Can be used for quantifying hydrogen peroxide in solutions, in cell extracts and in

live cells; and can also be used for detecting a variety of oxidase activities

through enzyme-coupled reactions.

Sensitive: Detect as low as 10 picomoles of H_2O_2 in solution.

Continuous: Easily adapted to automation without a separation step.

Convenient: Formulated to have minimal hands-on time. No wash is required.

Non-Radioactive: No special requirements for waste treatment.

Kit Components

Components	Amount		
Component A: Amplite™ Red Peroxidase Substrate	1 vial		
Component B: H ₂ O ₂	1 vial (3% stabilized solution, 200 μL)		
Component C: Assay Buffer	1 bottle (100 mL)		
Component D: Horseradish Peroxidase	1 vial (20 units)		
Component E: DMSO	1 vial (1 mL)		

Assay Protocol for One 96-well Plate

Brief Summary

Prepare H_2O_2 reaction mixture (50 μ L) \rightarrow Add H_2O_2 standards or test samples (50 μ L) \rightarrow Incubate at room temperature for 10-30 minutes \rightarrow Monitor fluorescence intensity at Ex/Em = 540/590 nm

Note: Thaw all the kit components at room temperature before starting the experiment.

1. Prepare stock solutions:

1.1 100X AmpliteTM Red peroxidase substrate stock solution: Add 250 μL of DMSO (Component E) into the vial of AmpliteTM Red Substrate (Component A). The stock solution should be used promptly; any remaining solution should be aliquoted and refrozen at -20 °C.

Note: Avoid repeated freeze-thaw cycles and protect from light.

1.2 <u>20 U/mL Peroxidase stock solution:</u> Add 1 mL of Assay Buffer (Component C) into the vial of Horseradish Peroxidase (Component D).

Note: The unused HRP solution should be divided into single use aliquots and stored at -20 °C.

1.3 20 mM H₂O₂ stock solution: Add 22.7 μL of 3% H₂O₂ (0.88 M, Component B) into 977 μL of Assay Buffer (Component C).

Note: The diluted H_2O_2 solution is not stable. The unused portion should be discarded.

2. Prepare H_2O_2 reaction mixture:

Prepare the H₂O₂ reaction mixture according to the following table and keep from light:

Table 1 H₂O₂ Reaction mixture for one 96-well plate (2X)

Components	Volume
Amplite™ Red Peroxidase Substrate Stock Solution (100X, from Step 1.1)	50 μL
20 U/ml Peroxidase Stock Solution (from Step 1.2)	200 μL
Assay Buffer (Component C)	4.75 mL
Total volume	5 mL

3. Prepare serial dilutuions of H_2O_2 standard (0 to 10 μM):

Warning 1: The component A is unstable in the presence of thiols such as DTT and β-ercaptoethanol. Thiols higher than 10 μM (final concentration) would significantly decrease the assay dynamic range.

Warning 2: NADH and electricians (not used forms CSH) may interfere with the assay.

Warning 2: NADH and glutathione (reduced form: GSH) may interfere with the assay.

- 3.1 Add 1 μ L of 20 mM H₂O₂ solution (from Step 1.3) into 1999 μ L of Assay Buffer (Component C) to get a 10 μ M H₂O₂ standard.
- 3.2 Take 200 μ L of 10 μ M H₂O₂ standard to perform 1:3 serial dilutions to get 3, 1, 0.3, 0.1, 0.03, 0.01 and 0 μ M serial dilutions of H₂O₂ standard.
- 3.3 Add serial dilutions of H₂O₂ standard and H₂O₂-containing test samples into a solid black 96-well microplate as described in Tables 2 and 3.

		•					
BL	BL	TS	TS	 			
HS1	HS1			 			
HS2	HS2						
HS3	HS3						
HS4	HS4						
HS5	HS5						
HS6	HS6						·
HS7	HS7						

Table 2 Layout of H₂O₂ standards and test samples in a solid black 96-well microplate

Note: HS= H₂O₂ Standards: BL=Blank Control: TS=Test Samples

Table 3 Reagent composition for each well

H ₂ O ₂ Standard	Blank Control	Test Sample		
Serial dilutions*: 50 μL	Assay Buffer (Component C): 50 μL	50 μL		

*Note: Add the serially diluted H_2O_2 standards from 0.01 μ M to 10 μ M into wells from HS1 to HS7 in duplicate. High concentration of H_2O_2 (e.g., > 100 μ M, final concentration) may cause reduced fluorescence signal due to the overoxidation of AmpliteTM Red (to a non-fluorescent product).

4. Run H₂O₂ assay in supernatants reaction:

- 4.1 Add 50 μL of H₂O₂ reaction mixture (from Step 2) into each well of H₂O₂ standard, blank control, and test samples (see Step 3.3) to make the total H_2O_2 assay volume of 100 μ L/well. Note: For a 384-well plate, add 25 μ L of sample and 25 μ L of H_2O_2 reaction mixture in each well.
- 4.2 Incubate the reaction at room temperature for 15 to 30 minutes, protected from light.
- 4.3 Monitor the fluorescence increase with a fluorescence plate reader at Ex/Em = $540 \pm 10 / 590 \pm 10$ nm (optimal Ex/Em = 540/590 nm). Note: The contents of the plate can also be transferred to a white clear bottom plate and read by an absorbance microplate reader at the wavelength of 576 ± 5 nm. The absorption detection has lower sensitivity compared to fluorescence reading.

5. Run H₂O₂ assay for cells:

The AmpliteTM Fluorimetric Hydrogen Peroxide Assay Kit can be used to measure the release of H₂O₂ from cells. The following is a suggested protocol that can be modified to meet the specific research needs.

- 5.1 The H₂O₂ reaction mixture should be prepared as Step 2 except that the Assay Buffer (Component C) should be replaced with the media that is used in your cell culture system. Suggested media including (a) Krebs Ringers Phosphate Buffer (KRPB); (b). Hanks Balanced Salt Solution (HBSS); or (c) Serum-free media.
- 5.2 Prepare cells in a 96-well plate (50 100 μL/well), and activate the cells as desired. Note: The negative controls (media alone and non-activated cells) are included for measuring background fluorescence.
- 5.3 Add 50 μL of H₂O₂ reaction mixture (from Step 5.1) into each well of cells and H₂O₂ standards (from Step
 - Note: For a 384-well plate, add 25 μ L of cells and 25 μ L of H_2O_2 reaction mixture into each well.
- 5.4 Incubate the reaction at room temperature for 15 to 30 minutes, protected from light.
- 5.5 Monitor the fluorescence increase with a fluorescence plate reader at Ex/Em= $540 \pm 10/590 \pm 10$ nm (optimal Ex/Em = 540/590 nm).

Data Analysis

The fluorescence in blank wells (with the assay buffer only) is used as a control, and is subtracted from the values for those wells with the H_2O_2 reactions. A H_2O_2 standard curve is shown in Figure 1.

Note: The fluorescence background increases with time, thus it is important to subtract the fluorescence intensity

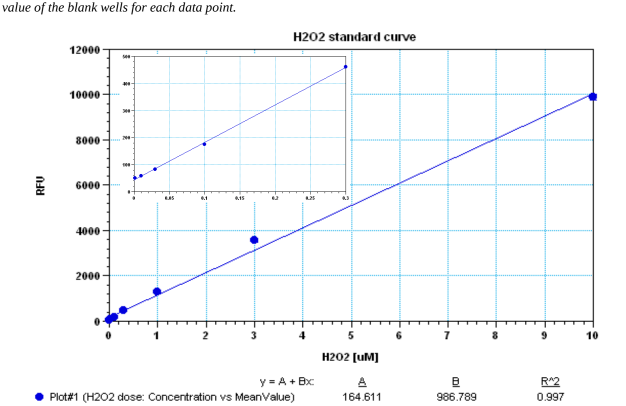


Figure 1. H_2O_2 dose response was measured in a 384-well black plate with the AmpliteTM Florimetric Hydrogen Peroxide Assay Kit using a Gemini fluorescence microplate reader (Molecular Devices). As low as 0.03 μ M H_2O_2 can be detected with 30 minutes incubation (n=3). The insert shows the low levels of H_2O_2 detection.

References

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