

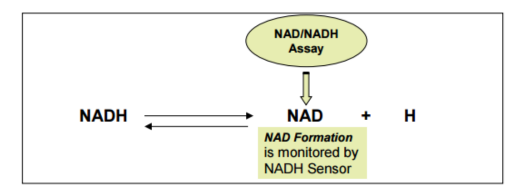
Elite[™] NAD/NADH Assay Kit (Red Fluorescence)

CATALOG NUMBER: CA-N218, 400 assays

Description

Nicotinamide adenine dinucleotide (NAD+) and nicotinamide adenine dinucleotide phosphate (NADP+) are two important cofactors found in cells. NADH is the reduced form of NAD+, and NAD+ is the oxidized form of NADH. NAD forms NADP with the addition of a phosphate group to the 2' position of the adenyl nucleotide through an ester linkage. NADP is used in anabolic biological reactions, such as fatty acid and nucleic acid synthesis, which require NADPH as a reducing agent. In chloroplasts, NADP is an oxidizing agent important in the preliminary reactions of photosynthesis. The NADPH produced by photosynthesis is used as reducing power for the biosynthetic reactions in the Calvin cycle of photosynthesis.

The traditional NAD/NADH and NADP/NADPH assays are run by monitoring the changes in NADH or NADPH absorption at 340 nm. This Elite™ Fluorimetric NAD/NADH Assay Kit provides a convenient method for sensitive detection of NAD and NADH. The enzymes in the system specifically recognize NAD/NADH in an enzyme cycling reaction that significantly increases detection sensitivity. In addition, this assay has very low background since it is run in the red visible range that considerably reduces the interference resulted from biological samples. The assay has demonstrated high sensitivity and low interference at Ex/Em = 540/590 nm.



The EliteTM Fluorimetric NAD/NADH Assay Kit provides a sensitive, one-step assay to detect as little as 10 picomoles of NAD(H) in a 100 μ l assay volume (100nM; Figure 1). The assay can be performed in a convenient 96-well or 384-well microtiter-plate format and readily adapted to automation without a separation step. Its signal can be easily read by either a fluorescence microplate reader at Ex/Em = 530-570 nm/590-600 nm (maximum Ex/Em = 540/590 nm) or an absorbance microplate reader at ~576 nm.

Features

- **Broad application**: NAD/NADH detection in solution or cell extracts.
- Sensitive: Detect as low as 10 picomoles of NAD/NADH in solution.
- Continuous: Easily adapted to automation without a separation step.
- Convenient: Formulated to have minimal hands-on time. No wash required.
- Non-radioactive: No special requirement for waste disposal.

Kit Components

Component A: NAD/NADH Recycling Enzyme Mixture,
 2 bottles (lyophilized powder)

• Component B: NADH Sensor Buffer, 1 bottle (20 ml)

Component C: NADH Standard,
 Component G: NAD/NADH Lysis Buffer,
 1 vial (142 µg)
 1 bottle (10 ml)

Please consider the environment before printing.



Storage

Keep in freezer (-20 °C) and avoid exposure to light.

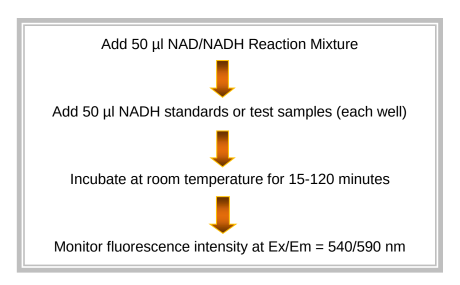
All reagents are stable for at least 6 months after receipt when stored properly at the recommended conditions.

Materials Required (but not supplied)

- 96-well black microplates with black bottom
- Fluorescence microplate reader

Assay Protocol (for One 96-Well Plate)

Summary of the Assay



1. Prepare NADH stock solution:

Add 200 µl of PBS buffer into the vial of NADH standard (**Component C**) to have 1 mM (1 nmol/ µl) NADH stock solution.

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

2. Prepare NAD/NADH reaction mixture:

Add 10 ml of NAD/NADH Sensor Buffer (**Component B**) to the bottle of NAD/NADH Recycling Enzyme Mixture (**Component A**), and mix well.

Note: This NAD/NADH reaction mixture is enough for two 96-well plates. The unused NAD/NADH reaction mixture should be divided into single use aliquots and stored at -20 °C.

3. Prepare serial dilutions of NADH standard (0 to 10 μ M):

3.1. Add 10 μ I of NADH stock solution (from Step 1) into 990 μ I PBS buffer (pH 7.4) to generate 10 μ M (10 pmols/ μ I) NADH standard solution.

Note: Diluted NADH standard solution is unstable, and should be used within 4 hours.

3.2. Take 200 μ I of 10 μ M NADH standard solution 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 NADH standard.



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3.3. Add serial dilutions of NADH standard and NAD/NADH containing test samples into a solid black 96-well microplate as described in Tables 1 and 2.

Note: Prepare cells or tissue samples as desired or as described in appendix A. NAD/NADH Lysis Buffer (**Component G**) can be used for lysing the cells.

Table 1. Layout of NADH standards and test samples in a solid black 96-well microplate

BL	BL	TS	TS	 			
NS1	NS1						
NS2	NS2						
NS3	NS3						
NS4	NS4						
NS5	NS5						
NS6	NS6						
NS7	NS7						

Note: NS= NADH Standards; BL=Blank Control; TS=Test Samples.

Table 2. Reagent composition for each well

NADH Standard	Blank Control	Test Sample			
Serial Dilutions: 50 µl	PBS: 50 μl	50 μl			

Note: Add the serially diluted NADH standards from 0.01 μ M to 10 μ M into wells from NS1 to NS7 in duplicates. High concentration of NADH (e.g. >100 μ M final concentration) may cause reduced fluorescence signal due to the over oxidation of NADH sensor (to a non-fluorescent product).

4. Run NAD/NADH assay in supernatants reaction:

- 4.1. Add 50 μ l of NADH reaction mixture (from Step 2) into each well of NADH standard, blank control, and test samples (from Step 3.3) to make the total NADH assay volume of 100 μ l/well.

 Note: For a 384-well plate, add 25 μ L of sample and 25 μ L of NADH reaction mixture into each well
- 4.2. Incubate the reaction at room temperature for 15 minutes to 2 hours, protected from light.
- 4.3. Monitor the fluorescence increase with a fluorescence plate reader at Ex/Em = 540/590 nm.

Note1: 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.

Note2: For measuring the NAD/NADH ratio, kit CA-N226 is recommended.

Data Analysis

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

Note: The fluorescence background increases with time, thus it is important to subtract the fluorescence intensity value of the blank wells for each data point.



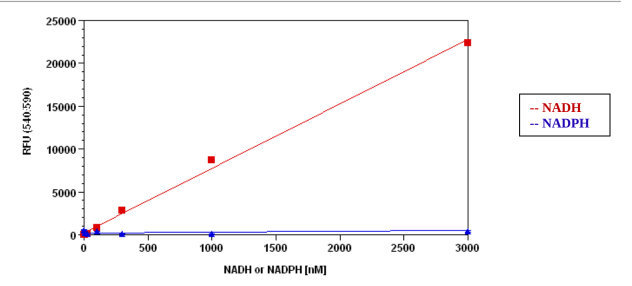


Figure 1. NADH dose response was measured with Elite™ NAD/NADH Assay Kit in a solid black 96-well plate using a NOVOStar microplate reader (BMG Labtech). As low as 100 nM (10 pmol/well) of NADH can be detected with 1 hour incubation (n=3) while there is no response from NADPH.

References

- 1. Ziegenhorn J, Senn M, Bucher T. (1976) Molar absorptivities of beta-NADH and beta-NADPH. Clin Chem, 22, 151.
- 2. Ikegami T, Kameyama E, Yamamoto SY, Minami Y, Yubisui T. (2007) Structure and Properties of the Recombinant NADH-Cytochrome b(5) Reductase of Physarum polycephalum. Biosci Biotechnol Biochem.
- 3. Kimura N, Fukuwatari T, Sasaki R, Shibata K. (2006) Comparison of metabolic fates of nicotinamide, NAD+ and NADH administered orally and intraperitoneally; characterization of oral NADH. J Nutr Sci Vitaminol (Tokyo), 52, 142.
- 4. O'Donnell JM, Kudej RK, LaNoue KF, Vatner SF, Lewandowski ED. (2004) Limited transfer of cytosolic NADH into mitochondria at high cardiac workload. Am J Physiol Heart Circ Physiol, 286, H2237.



Appendix A

Sample cell lysis preparation for the assay kits

This protocol serves as a general guide for preparing samples for the assays. For the best results, users need to choose their own method for cell lysis preparation. A method of freeze-thaw (or other lysis methods using physical disruption in PBS) is strongly recommended for the preparation of cell lysate, especially when the background noise becomes an issue for this assay.

1. Lysis of plant cells

Homogenize the leaves with the Cell/Tissue Lysis Buffer (**Component G**) at 200 mg/ml, and centrifuge at 2500 rpm for 5-10 minutes. Use the supernatant for the assay.

2. Lysis of bacterial cells

Collect bacterial cells by centrifugation (i.e. 10,000 x g, 0°C, 15 min). Add Cell/Tissue Lysis Buffer (**Component G**) to the pellet (1 mL per 100 to 10,000,000 cells), and leave at room temperature for 15 minutes. Centrifuge at 2500 rpm for 5 minutes, and use the supernatant for the assay.

3. Lysis of mammalian cells

Remove the medium from the culture plate (wells). Use about 100 μ L Cell/Tissue Lysis Buffer (**Component G**) per 1-5 million cells (or add lysis buffer 100 μ L lysis buffer per well in a 96-well cell culture plate), and leave at room temperature for 15 minutes. For the assay, use the lysate directly or centrifuge at 1500 rpm for 5min then use the supernatant.

4. Lysis of tissues

Weigh \sim 20 mg tissue, and then wash with cold PBS buffer. Homogenize the tissue with 400 μ l of Cell/Tissue Lysis Buffer (**Component G**) in a micro-centrifuge tube, and centrifuge at 2500 rpm for 5-10 minutes. Use the supernatant for the assay.

Note: use Cat# CA-N015g for ordering the Cell/Tissue Lysis Buffer separately.