



Nitric Oxide Synthase Assay Kit

Catalog Number KA1634

96 assays

Version: 04

Intended for research use only

www.abnova.com

Table of Contents

Introduction	3
Intended Use	3
Principle of the Assay	3
General Information	4
Materials Supplied	4
Storage Instruction	4
Materials Required but Not Supplied	4
Precautions for Use	4
Assay Protocol	5
Reagent Preparation	5
Assay Procedure	5
Data Analysis.....	7
Calculation of Results.....	7
Resources	8
References	8
Plate Layout	9

Introduction

Intended Use

Application:

- ✓ Direct Assays: NOS activity in biological samples.
- ✓ Drug Discovery/Pharmacology: effects of drugs on NOS activity.

Features:

- ✓ Sensitive and accurate: Detection range 0.25 - 25 U/L in 96-well plate.
- ✓ Rapid and reliable: Can be completed in 40 min if reduction of NO_3^- to NO_2^- is performed at 60°C.

Principle of the Assay

Nitric oxide (NO) is a reactive radical that plays an important role in many key physiological functions. NO, an oxidation product of arginine by nitric oxide synthase (NOS), is involved in host defense and development, activation of regulatory proteins and direct covalent interaction with functional biomolecules.

Simple, direct and non-radioactive procedures for measuring NOS are becoming popular in Research and Drug Discovery. Nitric Oxide Synthase Assay Kit involves two steps: a NOS reaction step during which NO is produced followed by an NO detection step. Since the NO generated by NOS is rapidly oxidized to nitrite and nitrate, the NO production is measured following reduction of nitrate to nitrite using an improved Griess method. The procedure is reduced to as short as 40 min.

General Information

Materials Supplied

List of component

Component	Amount
Assay Buffer	10 mL
Substrate	600 µL
GDH	120 µL
Reagent A	12 mL
Reagent B	500 µL
Reagent C	12 mL
Reagent D (Dried)	2 tubes
Reagent E	1.5 mL
ZnSO ₄	1 mL
Standard	1 mL
NaOH	1 mL

Storage Instruction

- ✓ Store Assay Buffer, Substrate, Reagent D, Reagent E and GDH at -20°C. Store all other reagents at 4°C. Shelf life of six months after receipt.
- ✓ Use Reagent D within 1 week after reconstitution.

Materials Required but Not Supplied

- ✓ Pipetting devices
- ✓ eppendorf tubes
- ✓ eppendorf centrifuge
- ✓ clear, flat bottomed 96 well plates or cuvettes
- ✓ plate reader or spectrophotometer
- ✓ heat block or hot water bath (optional)

Precautions for Use

Reagents are for research use only. Please refer to Material Safety Data Sheet for detailed information.

Assay Protocol

Reagent Preparation

Reconstitute one tube of Reagent D with 300 μL dH_2O . (If assaying more than 60 wells, reconstitute both tubes of Reagent D)

Store unused reconstituted Reagent D at -20°C and use within 1 week.

Assay Procedure

Note:

Prior to assay, equilibrate all components to room temperature.

Prewarm Assay Buffer to 37°C . Keep GDH on ice.

✓ Sample treatment:

Tissue or cell samples are homogenized in 1 x PBS (pH 7.4). Centrifuge at 10,000 g or higher at 4°C . Use supernatant for NOS assay.

✓ Standard preparation:

Prepare 200 μL 500 μM Premix by mixing 100 μL 1.0 mM Standard and 100 μL distilled water. Dilute standards in 1.5-mL centrifuge tubes as described in the Table.

No	Premix + H_2O	Nitrite (μM)
1	50 μL + 0 μL	500
2	30 μL + 20 μL	300
3	15 μL + 35 μL	150
4	0 μL + 50 μL	0

✓ NOS Reaction:

If samples will not require deproteinization (i.e. purified NOS), add 20 μL of each sample and standard to separate labeled eppendorf tubes. Each sample requires at least two tubes: one reaction tube and one sample blank tube. Immediately prior to starting the reaction, prepare enough NOS Working Reagent (NOS WR) for all sample reaction tubes and standards by mixing per reaction tube: 65 μL Assay Buffer, 4 μL Substrate, 4 μL Reconstituted Reagent D, 10 μL Reagent E and 1 μL GDH. For the sample blanks, use 8 μL dH_2O instead of the Substrate and Reagent D. Add 80 μL of the appropriate NOS WR to each tube and incubate at 37°C for 20 min. After 20 min immediately add 200 μL of the NO Detection Reagent (NO DR) (see next section: NO Measurement) to each tube to kill the NOS reaction.

For samples requiring deproteinization which include serum, plasma, whole blood, cell culture media containing FBS, tissue or cell lysates, add 25 μL of each sample and standard to separate labeled eppendorf tubes. Each sample requires at least two tubes: one reaction tube and one sample blank tube.

Immediately prior to starting the reaction, prepare enough NOS WR for all sample reaction tubes and standards by mixing per reaction tube: 80 μL Assay Buffer, 5 μL Substrate, 5 μL Reconstituted Reagent D, 13 μL Reagent E and 1 μL GDH. For the sample blanks, use 10 μL dH_2O instead of the Substrate and Reagent D. Add 100 μL of the appropriate NOS WR to each tube and incubate at 37°C for 20 min. After 20 min immediately proceed to the deproteinization step.

Deproteinization:

Add 7 μL ZnSO_4 to each sample and standard tube. Vortex and then add 7 μL NaOH. Vortex again and centrifuge 10 min at 14,000 rpm. Transfer 100 μL of the clear supernatant to a clean tube and proceed to the NO Measurement step.

✓ NO Measurement:

Immediately prior to starting the reaction, prepare enough NO Detection Reagent (NO DR) for all samples and standards by mixing per reaction tube: 100 μL Reagent A, 4 μL Reagent B and 100 μL Reagent C. Add 200 μL of the NO DR to each sample and standard tube and incubate for 5 min at 60°C. (Alternatively, the reaction can be run at 37°C for 60 min or RT for 150 min.)

Briefly centrifuge the reaction tubes to pellet any condensation and transfer 250 μL of each reaction to separate wells in a 96 well plate. Read OD at 500-570 nm (peak 540 nm).

Data Analysis

Calculation of Results

1. Subtract blank OD (Std 4) from the standard OD values and plot the OD against standard concentrations. Determine the slope using linear regression fitting. The NOS activity of the Sample is then calculated as:

$$\text{NOS Activity} = \frac{\text{OD}_{\text{Sample}} - \text{OD}_{\text{Blank}}}{\text{Slope}} \times \frac{1}{t} \text{ (U/L)}$$

2. $\text{OD}_{\text{Sample}}$ and OD_{Blank} are optical density values of the sample and sample blank, respectively. t is the reaction time (20 min).
3. Unit definition: one unit of NOS catalyzes the production of 1 μmole of nitric oxide per minute under the assay conditions (pH 7.5 and 37°C).
4. General Considerations:
Antioxidants and nucleophiles (e.g. β -mercaptoethanol, glutathione, dithiothreitol and cysteine) may interfere with this assay. Avoid using these compounds during sample preparation. However, if β -mercaptoethanol or dithiothreitol must be used, an equal concentration needs to be added to the standards.

Resources

References

1. Ghigo, D. (2006). Cycling of NADPH by glucose 6-phosphate dehydrogenase optimizes the spectrophotometric assay of nitric oxide synthase activity in cell lysates. *Nitric Oxide* 15: 148-53.
2. Knowles, R. G. and Moncada, S. (1994). Nitric oxide synthases in mammals. *Biochem. J.* 298: 249-58.
3. Förstermann, U. et al. (1991). Isoforms of nitric oxide synthase. Characterization and purification from different cell types. *Biochem Pharmacol.* 42:1849-57.

Plate Layout

12								
11								
10								
9								
8								
7								
6								
5								
4								
3								
2								
1								
	A	B	C	D	E	F	G	H