

# Nitric Oxide Synthase Ultrasensitive Colorimetric Assay

Catalog Number: NSU03-K01 96 Wells For Research Use Only v. 1.1 (20JUN23)

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## **INTRODUCTION**

The traditional method for measuring nitric oxide synthase (NOS) activity is performed by radiochemical assay that measures the conversion of L-[<sup>3</sup>H]arginine to L-[<sup>3</sup>H]citrulline. This method is expensive and requires regulation of radioactive materials. The Ultrasensitive Colorimetric NOS Assay Kit is a low-cost novel assay that allows for the detection of NOS activity without the need for radioactivity. Our Ultrasensitive NOS Assay Kit employs a NADPH recycling system to allow NOS to operate linearly for hours as nitric oxide-derived nitrate and nitrite accumulate. NOS can be assayed spectrophotometrically by measuring the accumulation of its stable degradation products, nitrate and nitrite. The ratio of these two products in biological fluids, tissue culture media, etc. may vary substantially. Hence, for accurate assessment of the total nitric oxide generated, one must monitor both nitrate and nitrite. An excellent solution to this problem is the enzymatic conversion of nitrate to nitrite by the enzyme nitrate reductase (NaR), followed by quantization of nitrite using Griess Reagent. This kit allows for efficient high-throughout screening of NOS activity in resting cells or cell lysates as well as biological fluids and tissue homogenates. The kit is also ideal for in vitro NOS assays using recombinant purified NOS. All materials necessary to perform the entire assay in a 96-well microplate format are provided with the kit.

For further information about this kit, its application or the procedures in this insert, please contact the Technical Service Team at Eagle Biosciences, Inc. at <u>info@eaglebio.com</u> or at 866-411-8023.

# PRINCIPLES OF THE PROCEDURE

NADPH and L-arginine are required for the continual operation of NOS and production of nitric oxide (NO). In aqueous solution, NO rapidly degrades to nitrate and nitrite. Spectrophotometric quantization of nitrite using Griess Reagent is straightforward but does not measure nitrate. This kit employs recombinant nitrate reductase (NaR) for conversion of nitrate to nitrite prior to quantization of nitrite using Griess reagent — thus providing for accurate determination of total NOS activity.



This Nitric Oxide Synthase Ultrasensitive Colorimetric Assay kit can be used to accurately measure as little as 1 pmol/ $\mu$ L (~1 $\mu$ M) NO produced in aqueous solutions. Very little sample is required (5 to 100  $\mu$ L depending on the [NO] in the sample. The completed reaction is read at 540 nm.



#### MATERIALS PROVIDED

- 1. **Microtiter Plates –** 1, 96-well low binding, flat-bottom
- 2. Reaction Buffer 1 bottle containing 50mL of 20mM HEPES 0.5mM EDTA
- 3. NADPH Part A 1 vial containing 1.0mL of NADP+ , Glucose 6-Phosphate, L-Arginine STORE AT -20C
- 4. NADPH Part B 1 vial containing1.0mL of Glucose 6-Phosphate dehydrogenase STORE AT -20C
- 5. Nitrate Reductase 1 vial containing 1U of Nitrate Reductase STORE AT -20C
- 6. Nitrate Reductase Buffer 1.2mL of buffer
- 7. Color Reagent #1 1 bottle containing 7.0 mL of Sulfanilamide in 3N HCI
- 8. **Color Reagent #2** 1 bottle containing 7.0 mL of N-(1-Naphthyl) ethylenediamine dihydrochloride
- 9. Nitrite Standard 1 vial containing 1.5mL of 500pmol / µL NaN02

# MATERIALS NEEDED BUT NOT PROVIDED

- 1. Reagent grade water: Distilled and deionized.
- 2. Microplate reader with 540nm filter. (Note: The wavelength of the filter can be 530 to 560 nm, but 540 nm is the absorbance maximum).
- 3. Precision pipettes ranging from 5  $\mu$ L 1.0 mL and disposable tips. NOTE: If all 96 wells are to be used at one time it is suggested that a multi-channel pipettor be used.
- 4. Test tubes to dilute the standards and unknowns.
- 5. Microcentrifuge tubes or microtiter plate for incubation of cell extracts or purified NOS.
- 6. Centrifuge for microcentrifuge tubes or microtiter plate if incubating in microtiter plate.

## STORAGE CONDITIONS

The Ultrasensitive Colorimetric NOS Assay kit is shipped on wet ice and stored at 4-8C. *Upon arrival, the Nitrate Reductase, NADPH Part A and NADPH Part B should be stored at –20°C until the time of use.* After initial use of the kit, store each component according to the Materials Provided

## PROCEDURAL NOTES

- 1. Assay can be used with cell lysates or purified NOS for in vitro assays. If using purified NOS it is important the required cofactors are added to the reaction. OBR sells a NOS Cofactor Mix specifically designed for use with this kit under product number **NS70**.
- 2. If using cell lysates for the assay, it is recommended that they are suspended in phosphate buffered saline (PBS) and protein concentration determined.
- 3. It is possible for the entire assay to be run in a 96-well microplate if the researcher has a centrifuge equipped with a microplate rotor. The initial incubation should be carried out in the 96-well V-bottom plate while the Color Reaction should be done in the 96-well flat bottom plate.
- 4. For ease in setting up multiple samples, the Reaction Buffer can be combined with NADPH Part A and Part B and then aliquoted equally to each sample.
- 5. If the NO concentration in your sample is low, you can increase the sample volume in the Color Reaction to  $100 \ \mu$ L while decreasing the buffer volume.



6. For best results, complete the reading of the plate within 20 minutes.

#### PREPARATION

- 1. Nitrate Reductase: Reconstitute with 1ml of Nitrate Reductase Buffer. Vortex briefly every five minutes for fifteen minutes total.
- 2. Nitrate Standard: Nitrite Standard must first be diluted from 500  $\mu$ M to 100  $\mu$ M by adding 240  $\mu$ L of the Nitrite Standard to 960  $\mu$ L of Reaction Buffer. This should be labeled Diluted Standard and stored on ice until used. The standard curve is then created by further diluting the Diluted Standard according to Table 2 below.

Standard	Final Concentration (µM)	ddH2O (µL)	Diluted Standard (µL)
B <sub>0</sub>	0	1000	0
S <sub>1</sub>	0.5	995	5
S <sub>2</sub>	1	990	10
<b>S</b> 3	5	950	50
<b>S</b> 4	10	900	100
<b>S</b> 5	25	750	250
<b>S</b> 6	50	500	500
<b>S</b> <sub>7</sub>	100	0	250

#### Table 2: Preparation of the Standard Curve.

**NOTE:** Standard solutions may be stored at 4°C for later use.

## PROCEDURE

#### In V-well microplate or microcentrifuge tube

- 1. Add 40–500 μg of protein from lysates or 0.2 1.0 Unit of recombinant or purified NOS in a volume of 30 μl to a tube or well.
- 2. Add 200 µL Reaction Buffer
- 3. Add 10 µL of NADPH Part A
- 4. Add 10 µL of NADPH Part B
- 5. If using purified NOS, add 10µL of the NS70 NOS Cofactor Mix
- 6. Mix and incubate for 1 6 hours at 37°C.
- 7. Chill on ice for 5 minutes.
- 8. Add 10 µL of the reconstituted Nitrate Reductase to each sample, vortex tube or tap plate to mix, and incubate for 20 minutes at room temperature.
- 9. Centrifuge at 12,500 rpm for 5 minutes at 4°C.

## In flat-bottom microtiter plate

- 10. Add 100 µL of Standards or Samples to the microplate in duplicate. Depending on [NO], the samples may need to be diluted in Reaction Buffer. See Scheme 1 for a sample plate layout.
- 11. Add 50 µL Color Reagent #1 and shake briefly.



Add 50 µL Color Reagent #2. Shake for 5 minutes at room temperature.
 Read absorbance values at 540 nm in Microtiter plate reader.

	1	2	3	4	5	6	7	8	9	10	11	12
А	BO	BO	U1	U1	U9	U9	U17	U17	U25	U25	U33	U33
В	S <sub>1</sub>	S <sub>1</sub>	U2	U2	U <sub>10</sub>	U <sub>10</sub>	U <sub>18</sub>	U <sub>18</sub>	U <sub>26</sub>	U <sub>26</sub>	U34	U34
С	S2	S2	U3	U3	U11	U11	U19	U19	U27	U27	U35	U35
D	S3	S <sub>3</sub>	U4	U4	U <sub>12</sub>	U <sub>12</sub>	U <sub>20</sub>	U <sub>20</sub>	U <sub>28</sub>	U <sub>28</sub>	U36	U36
E	S4	S4	U5	U5	U <sub>13</sub>	U <sub>13</sub>	U <sub>21</sub>	U <sub>21</sub>	U29	U29	U37	U37
F	S5	S5	U6	U6	U14	U14	U22	U22	U30	U30	U38	U38
G	S6	S6	U7	U7	U15	U15	U23	U23	U31	U31	U39	U39
Н	S7	S7	U8	U8	U16	U16	U24	U24	U32	U32	U40	U40

# Table 3: Sample microplate layout

 $S_0$  = Blanks: Add buffer in place of standard or sample.

 $U_{1-40}$  = Unknown sample to be assayed.

# CALCULATIONS

- 1. Subtract the average O.D. value of the blank wells (S<sub>0</sub>) from all other pairs of wells.
- 2. Average the O.D. values for each pair of duplicate wells.
- 3. Plot a standard curve using the average O.D. value for each standard value versus the concentration of standard. If samples are less than 10  $\mu$ M, plot standards B0, S1, S2, S3, and S4. For samples over 10  $\mu$ M, plot standards S4, S5, S6, and S7.
- 4. Determine the concentration of each unknown by interpolation from the standard curve.
- 5. Samples can be compared by determining  $\mu moles$  of NO produced /  $\mu g$  protein / unit time.



Figure 1: Typical Standard Curve

Nitric Oxide Synthase Ultrasensitive Colorimetric Assay Catalog Number: NSU03-K01





#### WARNINGS AND PRECAUTIONS

- 1. **DO NOT** use components beyond the specified expiration date.
- 2. Universal precautions should be employed with handling of the components provided in this kit.
- 3. Use aseptic techniques when opening and removing reagents from vials and bottles.

#### REFERENCES

Ghigo, D., et al. (2006). *Nitric Oxide* **15**:148-153 Schmidt, H.H., et al. (1995). *Biochemica* **2**:22-23

#### WARRANTY INFORMATION

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For further information about this kit, its application, or the procedures in this kit, please contact the Technical Service Team at Eagle Biosciences, Inc. at info@eaglebio.com or at 866-411-8023.