

Nitrite Reductase (NiR) Activity Assay Kit

Note: The reagents have been changed, so please be aware of and follow this instruction strictly.

Operation Equipment: Spectrophotometer

Catalog Number: BC1540

Size:50T/24S

Components:

Extract solution: Liquid 30 mL×1. Store at 2-8°C.

Reagent I: Liquid 8 mL×1. Store at 2-8°C.

Reagent II: Powder×2. Store at 2-8°C. Dissolve with 6mL of distilled water before use. It could be stored at 2-8°C for two weeks.

Reagent III: Powder×1. Store at 2-8°C. Dissolve with 15mL of distilled water at 70-80°C before use. It could be stored at 2-8°C for three months.

Reagent IV: Liquid 25 mL×1. Store at 2-8°C.

Reagent V: Liquid 25 mL×1. Store at 2-8°C. If there is precipitation in the reagent, it could be dissolved at 70-80°C.

Standard: Liquid 1 mL×1. Store at 2-8°C. 10 μmol/mL of NaNO₂ standard solution.

Working solution: Reagent IV and Reagent V are mixed by the ratio of 1:1 to make working solution. Prepare when the solution will be used.

Product Description:

Nitrite reductase (NiR) is a key enzyme in the reduction of nitrite nitrogen, and plays an important role in the biogeochemical nitrogen cycle, which is widely found in microbes and plants. It catalyze the reduction of nitrite nitrogen, reduce the accumulation of nitrite nitrogen and its toxic effect on the growth and development of organisms.

Nitrite reductase can reduce NO₂⁻ to NO, and reduce the NO₂⁻ in the sample to participate in the diazotization reaction to produce a purple-red compound, that is, the change in absorbance at 540nm can reflect the activity of nitrite reductase in soil.

Reagents and Equipment Required but Not Provided:

Spectrophotometer, centrifuge, adjustable transferpettor, balance, mortar/homogenizer, 1mL glass cuvette, ice and distilled water.

Procedure:

I. Sample preparation

1. Tissue: according to the proportion of tissue weight (g): extraction solution volume (mL) of 1:5-10 to extract. It is suggested that 0.1 g of tissue with 1 mL of Extraction reagent and fully homogenized on ice bath. Centrifuge at $10000 \times g$ for 10 minutes at $4^{\circ}C$ to remove insoluble materials, and take the supernatant

on ice before testing.



2. Bacteria or cells: Collect bacteria or cells into the centrifuge tube, after centrifugation discard supernatant. It is suggested that add 1 mL of Extraction reagent to 5 million of bacteria or cells. Use ultrasonication to split bacteria or cells (place on ice, ultrasonic power 200W, working time 3 seconds, interval 7 seconds, repeat for 3 minutes). Centrifuge at 10000 ×g for 10 minutes at 4°C to remove insoluble materials and take the supernatant on ice for testing.

II. Determination

- 1. Preheat spectrophotometer for 30 min, adjust the wavelength to 540 nm and set counter to zero with distilled water.
- 2. Standard working solution: dilute 10μ mol/mL standard solution to 0.08, 0.06, 0.05, 0.025, 0.0125, 0.00625, 0.003125, 0.0015625 μ mol/mL with distilled water.
- 3. Add reagent to a 1.5 mL EP tube:

Reagent (µL)	Matrix tube (Am)	Control tube (Ac)	Test tube (At)	Standard tube (As)	Blank tube (Ab)
Sample	- ~0	100	100	-	-
Distilled water	100	200	7.00	-	-
Reagent I	200	-	200	-	-
Reagent II	200	200	200	-	- 0
Viol.	After mixing, react at 25°C for 1 h.			-	101069
Reagent III	200	200	200	- C	0/000
Co Super	Fully shake for 30s, stand for 5min and take supernatant.			8	-
Supernatant	350	350	350	-	-
Standard	- 5%	<u>-</u>	0	350	-
Distilled water	-5	-	CALDINE'S	-	350
Working solution	700	700	700	700	700

Mix well and stand for 5min. Measure the absorbance value at the wavelength of 540nm, and record them as Am, Ac, At, As and Ab, and calculate Δ At = Am-(At-Ac), Δ As = As-Ab. Each test tube should be provided with one contrast tube. Standard curve, Matrix tube (Am) and Blank tube (Ab) only be measured once or twice.

III. Calculation:

1. Standard curve

Taking the concentration of each standard solution as the x-axis and its corresponding ΔA_S as the y-axis, draw a standard curve to get the standard equation y = kx + b, and bring ΔA_S into the equation to get x (μ mol/mL).

2. Calculation

1) Protein concentration:

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Unit definition: One unit of enzyme activity is defined as the amount of enzyme catalyzes the reduction of 1µmol NO₂- per hour every milligram protein in the reaction system.

NiR (U/mg prot) =
$$x \times V1 \div V2 \div Cpr \div T = x \times 7 \div Cpr$$

2) Sample weight

Unit definition: One unit of enzyme activity is defined as the amount of enzyme catalyzes the reduction of 1µmol NO₂- per hour every gram tissue sample in the reaction system.

NiR (U/g weight) =
$$x \times V1 \div V2 \times V_E \div W \div T = x \times 7 \div W$$

3) Bacteria or cells

Unit definition: One unit of enzyme activity is defined as the amount of enzyme catalyzes the reduction of 1µmol NO₂- per hour every 10⁴ bacteria or cells in the reaction system.

NiR (U/10⁴ cell) =
$$x \times V1 \div V2 \times V_E \div N \div T = x \times 7 \div N$$

V1: Total volume before taking supernatant, 0.7mL;

V2: Sample volume, 0.1 mL;

V_E: Extraction volume, 1.0 mL;

T: Reaction time, 1 hour;

W: Sample weight, g;

Cpr: Sample protein concentration, mg/mL;

N: Bacteria or cells number, 10⁴.

Experimental examples:

1. Take 0.1003g Iris tectorum leaf for sample processing and follow the measurement procedure. After determination with micro glass cuvette, calculate $\Delta At = Am$ -(At-Ac)=0.849- (0.713-0.008) =0.144.Bring the result into the standard curve y=13.214x-0.0311, R²=0.9915, and calculate x=0.01325. The result is calculated according to the sample mass.

NiR (U/g weight) = $x \times 7 \div W = 0.03125 \times 7 \div 0.1003 = 0.9247$ U/g weight.

Related Products:

BC0080/BC0085 Nitrate Reductase (NR) Activity Assay Kit

BC1480/BC1485 Nitrite Assay Kit (Water And Soil)

BC1490/BC1495 Food Nitrite Content Assay Kit

BC1500/BC1505 Plant Nitrate Nitrogen Assay Kit

BC1520/BC1525 Plant Ammoniacal Nitrogen Assay Kit

BC4960/BC4965 Nitrate Reductase (NR) Activity Assay Kit (Griess-Colorimetric Method)