

NADP Phosphatase (NADPase) Activity Assay Kit

Note: Take two or three different samples for prediction before test.

Operation Equipment: Spectrophotometer/Microplate reader

Cat No: BC1115

Size: 100T/48S

Product Composition: Before use, please carefully check whether the volume of the reagent is consistent with the volume in the bottle. If you have any questions, please contact Solarbio staff in time.

Reagent name	Size	Preservation Condition
Extract solution	Liquid 50 mL×1	2-8°C
Reagent I	Liquid 15 mL×1	2-8°C
Reagent II	Powder×2	2-8°C
Reagent III	Powder×1	2-8°C
Reagent IV	Powder×1	2-8°C
Reagent V	Liquid 8 mL×1	RT
Standard	Liquid 1 mL×1	2-8°C

Solution Preparation:

1. Reagent II: Add 1.1mL of Reagent I to fully dissolve for later use and prepare it fresh.
2. Reagent III: Add 8 mL of distilled water for use, dissolves and save at 2-8 °C for a week.
3. Reagent IV: Add 8 mL of distilled water for use, dissolves and storage at 2-8 °C for a week.
4. Standard: 10 mmol/L standard phosphorus stock solution. Storage at 2-8°C. Preparation of 1 μmol/mL standard phosphorus application solution: dilute the standard for 10 times, namely take 1 mL standard solution add 9mL distilled water, fully mix.
5. Preparation of phosphorus detect reagent: it is prepared that the ratio of distilled water : Reagent III: Reagent IV: ReagentV=2:1:1:1. The prepared phosphorus detect reagent should be light yellow. If it is colorless, the reagent is invalid. If it is blue, it is polluted by phosphorus. Phosphorus detect reagent is prepared when the solution will be used.

Note: It is better to use the new beaker, glass rod and glass pipette to prepare reagent, or to use disposable plastic utensils to avoid phosphorus pollution.

Product description:

NADPase mainly exists in plant tissues and is the only enzyme in organisms that catalyzes the degradation of NADP⁺ to NAD⁺. NADPase and NADK regulate the balance between NAD and NADP.

NADPase can catalyze the hydrolysis of NADP⁺ to NAD⁺ and inorganic phosphorus, the activity of NADPase is determined by determining the content of inorganic phosphorus.

Reagents and Equipment Required but Not Provided

Refrigerated centrifuge, water-bath, adjustable pipette, spectrophotometer/microplate reader, micro glasscuvette/96 well plate, homogenizer/mortar and distilled water.

Procedure

I. Extraction of crude enzyme solution:

Tissue samples: Take about 0.1 g of sample, and add 1.0 mL of Extract solution for full grinding. Centrifuge at 8000 ×g for 10 minutes at 4°C, take the supernatant and place it on ice under test.

II. Test procedure

a. Preheat the spectrophotometer/microplate reader, for more than 30 minutes, adjust the wavelength to 660 nm and adjust zero with distilled water.

b. Operate table:

Enzymatic reaction:

Reagent Name (μL)	Test Tube (A _T)	Contrast Tube (A _C)
Reagent I	120	120
Reagent II	40	-
Distilled water	-	40
37°C (mammals) or 25°C preheat for 5 minutes (other species)		
Sample	40	40

37°C (mammals) or 25°C (other species) accurately react for 20 minutes, boiling water bath for 5 minutes (cover tightly, in order to prevent moisture loss). After cooling, centrifuge at 10000 ×g for 10 minutes at room temperature, take the supernatant.

Detect phosphorus content:

Reagent Name (μL)	Standard Tube (A _S)	Blank Tube (A _B)	Test Tube (A _T)	Contrast Tube (A _C)
1 μmol/mL standard phosphorus application solution	20	-	-	-
Distilled water	-	20	-	-
Supernatant	-	-	20	20
Fixed phosphorus reagent	200	200	200	200

Mix thoroughly, 37°C (mammals) water bath 30 minutes, cool to room temperature. Then draw 200 μL to micro glass cuvette or 96 well plate, and record the absorbance value at 660 nm, note as A_S, A_B, A_T, A_C, and calculate the $\Delta A_S = A_S - A_B$, $\Delta A_T = A_T - A_C$.

III. Calculation of Enzyme activity of NADPase:

1. Calculate by protein concentration:

Definition: One unit of enzyme activity is defined as the amount of NADPase catalyzes the decompose NADP to produce 1 μmol of inorganic phosphorus per minute every milligram of tissue protein.

$$\begin{aligned} \text{NADPase (U/mg prot)} &= \Delta A_T \div \Delta A_S \div C_S \times V_{SU} \div (C_{Pr} \times V_S \times V_{SU} \div V_{EN}) \div T \\ &= 0.25 \times \Delta A_T \div \Delta A_S \div C_{Pr} \end{aligned}$$

2. Calculate by sample fresh weight:

Definition: One unit of enzyme activity is defined as the amount of NADPase catalyzes the decompose NADP to produce 1 μmol of inorganic phosphorus per minute every gram tissue.

$$\begin{aligned} \text{NADPase (U/g fresh weight)} &= \Delta A_T \div \Delta A_S \div C_S \times V_{SU} \div (W \times V_S \div V_{EX} \times V_{SU} \div V_{EN}) \div T \\ &= 0.25 \times \Delta A_T \div \Delta A_S \div W \end{aligned}$$

C_S: 1 μmol/mL phosphorus standard application solution;

V_{SU}: The supernatant liquid volume in constant phosphorus test, 0.02 mL;

C_{pr}: Sample protein concentration, mg/mL;

V_S: The sample volume in the enzymatic reaction, 0.04 mL;

V_{EN}: Total volume of enzymatic reaction, 0.2 mL;

T: Reaction time, 20 minutes;

V_{EX}: Extract solution volume, 1 mL;

W: Sample fresh weight, g.

Note:

1. This method has the characteristics of trace, sensitive and rapid. So the test tube has strict requirements that must have not phosphorus, if the test tube pass phosphate or phosphate buffer, must be washed very clean, first boiled with detergent and water, then with tap water, finally rinse with distilled water. Using disposable plastic or new glass pipe is the best, avoiding phosphorus pollution is the key to detect success or failure.
2. The blank tube and the standard tube only need to be done once.

Experimental instances:

1. Take 0.1g of liver, add 1mL of extract solution, homogenate and grind. Take the supernatant and detect according to the measured steps. Calculate $\Delta A_T = A_T - A_C = 0.265 - 0.254 = 0.011$, $\Delta A_S = A_S - A_B = 0.533 - 0.047 = 0.486$, calculate the enzyme activity according to sample weight:

$$\text{NADPase (U/g weight)} = 0.25 \times \Delta A_T \div \Delta A_S \div W = 0.25 \times 0.011 \div 0.486 \div 0.1 = 0.057 \text{ U/g weight.}$$

2. Take 0.1g Setaria (root crop), add 1mL of extract solution, homogenate and grind. Take the supernatant and detect according to the measured steps. Calculate $\Delta A_T = A_T - A_S = 0.336 - 0.107 = 0.229$, $\Delta A_S = A_S - A_B = 0.533 - 0.047 = 0.486$, calculate the enzyme activity according to sample weight:

$$\text{NADPase (U/g weight)} = 0.25 \times \Delta A_T \div \Delta A_S \div W = 0.25 \times 0.229 \div 0.486 \div 0.1 = 1.18 \text{ U/g weight.}$$

References:

[1] Kawai S, Mori S, Mukai T, et al. Cytosolic NADP phosphatases I and II from *Arthro bacter* sp. strain KM: implication in regulation of NAD⁺/NADP⁺ balance[J]. *Journal of Basic Microbiology: An International Journal on Biochemistry, Physiology, Genetics, Morphology, and Ecology of Microorganisms*, 2004, 44(3): 185-196.

Related products:

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BC0400/BC0405 Isocitrate Dehydrogenase Cytoplasmic(ICDHc) Assay Kit
BC2100/BC21056-phosphogluconate Dehydrogenase(6-PGDH)Activity Assay Kit