

Aldehyde Dehydrogenase(ALDH) Activity Assay Kit

Note: It is necessary to predict 2-3 large difference samples before the formal determination.

Operation Equipment: Spectrophotometer

Cat No: BC0750

Size: 50T/48S

Components:

Extract solution: Liquid 60 mL×1, store at 4°C;

Reagent I: Liquid 20 mL×1, store at 4°C;

Reagent II: Powder×1, store at -20°C and protect from light. Add 6 mL of distilled water when the solution will be used. The rest of reagent store at -20°C; It can also be prepared in proportion when the solution will be used;

Reagent III: Liquid 1 mL×1, store at 4°C and protect from light;

Reagent IV: Liquid 2 mL×1, store at 4°C;

Reagent V: Liquid 3 mL×1, store at 4°C.

Product Description:

Aldehyde dehydrogenase (EC 1.2.1.10) is a kind of aldehyde dehydrogenase. It widely exists in various animals, plants and microorganisms. In the presence of coenzyme I, it can catalyze the dehydrogenation of some primary or secondary alcohols, aldehydes or ketones, including ethanol. In humans and many animals, mitochondrial acetaldehyde dehydrogenase can transform harmful alcohols. So in the study of cell detoxification, glyoxal dehydrogenase is highly concerned; Aldehyde dehydrogenase is widely used in molecular biology and detection of related diseases.

Acetaldehyde dehydrogenase catalyzes the conversion of acetaldehyde and NAD⁺ to acetic acid and NADH. The activity of aldehyde dehydrogenase can be calculated by the change of absorbance value of NADH at 340 nm.

Required but Not Provided:

Ultraviolet spectrophotometer, desk centrifuge, water-bath, adjustable pipette, 1 mL quartz cuvette, mortar /homogenizer, ice and distilled water.

Protocol

I. Preparation:

1. Tissue:

According to the tissue weight (g): the volume of the Extract solution (mL) is 1:5~10 to prepare (it is recommended that add 1 mL of Extract solution to 0.1 g of tissue). Homogenate on ice. Centrifuge at 10000 g and 4°C for 20 minutes. Take the supernatant on ice for test.

2. Cells or bacterial

According to the number of bacteria or cells (10^4): the volume of extraction solution (mL) is 500-1000:1 to prepare (it is recommended that add 1 mL of extraction solution to 500 million of cells). Bacteria/cells is split by ultrasonication (power 300W, ultrasonic 3s, interval 7s, total time 3 min). Centrifuge at 10000 g and 4°C for 20 minutes. Take the supernatant on ice for test.

3. Liquid: detect directly.

II. Determination procedure:

1. Preheat ultraviolet spectrophotometer/microplate reader for 30 min, adjust wavelength to 340 nm, set the counter to zero with distilled water.

2. Preheat Reagent I in 37°C (mammal) or 25°C (other species) for 15 min.

3. Operation table:

Reagent Name (μL)	Blank tube (A_B)	Test tube (A_T)
Sample		200
Distilled water	500	300
Reagent I	300	300
Reagent II	100	100
Reagent III	20	20
Reagent IV	30	30
Reagent V	50	50

The above reagents are added into the 1 mL quartz cuvette in sequence. Mix thoroughly. Measure the absorbance A_1 at 340 nm for 30s. Put it in a water bath or incubator at 37°C(mammal) or 25°C (other species) for 1 min. Take it out and dry it quickly, and then measure the absorption value A_2 at 90s. $\Delta A_T = A_{2T} - A_{1T}$. $\Delta A_B = A_{2B} - A_{1B}$. $\Delta A = \Delta A_T - \Delta A_B$. Blank tube just need to test once or twice.

III. ALDH Calculation:

1) Protein concentration:

Unit definition: One unit of enzyme activity is defined as the amount of enzyme catalyzes the production of 1 nmol NADH per minute in the reaction system every milligram tissue protein.

$$\text{ALDH (U/mg prot)} = \Delta A \div (\epsilon \times d) \times V_{RT} \div (C_{pr} \times V_{SA}) \div T \times 10^9 = 804 \times \Delta A \div C_{pr}$$

2) Sample weight:

Unit definition: One unit of enzyme activity is defined as the amount of enzyme catalyzes the production of 1 nmol NADH per minute in the reaction system every gram tissue weight.

$$\text{ALDH (U/g weight)} = \Delta A \div (\epsilon \times d) \times V_{RT} \div (W \div V_E \times V_{SA}) \div T \times 10^9 = 804 \times \Delta A \div W$$

3) Cells or germ

Unit definition: One unit of enzyme activity is defined as the amount of enzyme catalyzes the production of 1 nmol NADH per minute in the reaction system every 10^4 cells or germ.

$$\text{ALDH (U/}10^4 \text{ cell)} = \Delta A \div (\epsilon \times d) \times V_{RT} \div (\text{cells (million)} \times V_{SA} \div V_E) \div T \times 10^9 = 804 \times \Delta A \div \text{cells (million)}$$

4) Liquid volume

Unit definition: One unit of enzyme activity is defined as the amount of enzyme catalyzes the production of 1 nmol NADH per minute in the reaction system every milliliter liquid.

$$\text{ALDH (U/mL)} = \Delta A \div (\epsilon \times d) \times V_{RT} \div V_{SA} \div T \times 10^9 = 804 \times \Delta A$$

ϵ : NADH molar extinction coefficient, 6.22×10^3 L/mol/cm;

d: Light path of cuvette, 1 cm;

V_{RT} : Total reaction volume, 0.001 L;

V_{SA} : Sample volume, 0.2 mL;

V_E : Extract solution volume, 1 mL;

T: Reaction time, 1 min;

C_{pr}: Protein concentration, mg/mL;

W: Sample weight, g.

109: unit conversion factor, 1mol=10⁹ nmol.

Note:

1. The blank tube is the test hole for testing the quality of each reagent component. Under normal circumstances, the OD value should not exceed 0.3, and the change should not exceed 0.01.

2. When the ΔA is greater than 1.0, it is recommended to measure after dilution. When ΔA is less than 0.01, the reaction time can be prolonged to 5 min or 10 min for determination.

Experimental example:

1. Take 0.1g mouse kidney and add 1ml extract for homogenate grinding. Take the supernatant and dilute it 10 times. Operate according to the determination steps. Calculate $\Delta A_T = A_{2T} - A_{1T} = 0.258 - 0.203 = 0.055$, $\Delta A_B = A_{2B} - A_{1B} = 0$, $\Delta A = \Delta A_T - \Delta A_B = 0.055 - 0 = 0.055$

ALDH activity (U/g mass) = $804 \times \Delta A \div W \times 10$ (dilution) = $804 \times 0.055 \div 0.1 \times 10$ (dilution) = 4422 U/g mass

Related Products:

[1] Tongmeng Jiang, Jinmin Zhao, Shan Yu, et al. Untangling the response of bone tumor cells and bone forming cells to matrix stiffness and adhesion ligand density by means of hydrogels. *Biomaterials*. January 2019;188:130-143.(IF5.452)

[2] Chong Li, Shi Gao, Xiaotong Li, et al. Efficient metabolic evolution of engineered *Yarrowia lipolytica* for succinic acid production using a glucose-based medium in an in situ fibrous bioreactor under low-pH condition. *Biotechnology for Biofuels*. August 2018;(IF5.452)

[3] Yufei He, Xiaoyan Ci, Ying Xi, et al. Untangling the response of bone tumor cells and bone forming cells to matrix stiffness and adhesion ligand density by means of hydrogels. *Biomaterials*. September 2018;(2019)188:130-143.(IF8.806)

Related Products:

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BC2340/BC2345 Lipase(LPS) Activity Assay Kit

BC1080/BC1085 Alcohol Dehydrogenase(ADH) Activity Assay Kit