

Glycogen Phosphorylase a (GPa) Activity Assay Kit

Note: Take two or three different samples for prediction before test. Operation Equipment: Ultraviolet spectrophotometer/Microplate reader Catalog Number: BC3345 Size: 100T/96S

Components:

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	Reagent	Size	Storage
	Extract solution	110 mL×1	4°C
	Reagent I	20 mL×1	4°C
13	Reagent II	Powder×1	4°C
302	Reagent III	Powder×1	4°C
	Reagent IV	Powder×1	-20°C
	Reagent V	Powder×2	-20°C
	Reagent VI	Powder×2	-20°C

Solution preparation:

1. **Reagent II:** Dissolved with 0.5 mL of distilled water before use. Mix thoroughly. It can be divided into small tubules and stored for 4 weeks at -20°C. Avoid repeating freeze/thaw cycles.

2. **Reagent III:** Dissolved with 0.5 mL of distilled water before use. Mix thoroughly. It can be divided into small tubules and stored for 4 weeks at -20°C. Avoid repeating freeze/thaw cycles.

3. **Reagent IV:** Dissolved with 1.25 mL of distilled water before use. Mix thoroughly. It can be divided into small tubules and stored for 4 weeks at -20°C. Avoid repeating freeze/thaw cycles.

4. **Reagent V:** Dissolved with 1 mL of distilled water before use. Mix thoroughly. It can be divided into small tubules and stored for 4 weeks at -20°C. Avoid repeating freeze/thaw cycles.

5. **Reagent VI:** Dissolved with 1 mL of distilled water before use. Mix thoroughly. It can be divided into small tubules and stored for 4 weeks at -20°C. Avoid repeating freeze/thaw cycles.

6. Working solution: Calculate according to the amount required for the experiment before use, according to the ratio of reagent I: reagent II: reagent III: reagent IV: distilled water= 148μ L: 4μ L: 4μ L: 4μ L: 10μ L (amount of 1T), mix well before use.

Product Description:

Glycogen phosphorylase is divided into active glycogen phosphorylase a (Glycogen phosphorylase a, GPa) and inactive glycogen phosphorylase b (Glycogen phosphorylase b, GPb) two forms. The decomposition of glycogen is mainly carried out under the catalysis of glycogen phosphorylase a. When no activator is added, glycogen phosphorylase a catalyzes the production of glucose residues from glycogen and inorganic phosphorus to glycogen and glucose 1-phosphate. Under the action of phosphoglucose mutase and 6-phosphate glucose dehydrogenase, it further catalyzes the reduction of NADP to NADPH. Measuring the

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rate of increase of NADPH at 340nm can reflect the activity of glycogen phosphorylase a.

Reagents and Equipment Required but Not Provided:

Ultraviolet spectrophotometer/microplate reader, low temperature centrifuge, constant temperature incubator/water bath, adjustable pipette, mortar/homogenizer, micro quartz cuvette/96 well UV plate, ice and distilled water.

Procedure

I. Sample preparation:

1. Tissue sample: 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 8000 \times g for 10 minutes at 4°C to remove insoluble materials, and take the supernatant on ice before testing.

2. Bacteria or cells: collecting bacteria or cells into the centrifuge tube, suggested 5 million with 1 mL of Extraction reagent. Use ultrasonication to splitting bacteria and cells (placed on ice, ultrasonic power 200w, working time 3 seconds, interval 10 seconds, repeat for 30 times). Centrifuge at 8000 \times g for 10 minutes at 4°C to remove insoluble materials, and take the supernatant on ice for testing.

3. Serum (plasma) sample: detect sample directly.

II. Determination procedure:

- 1. Preheat ultraviolet spectrophotometer/microplate reader for 30 minutes, adjust wavelength to 340 nm, set zero with distilled water.
- 2. Working solution are preheated at 37°C for 5min.
- 3. GPa activity: add 10 μ L sample, 10 μ L reagent V, 10 μ L reagent VI, 170 μ L working solution in the micro quartz cuvette/96 well UV plate, mix immediately, and record the absorbance value at 10s at 340 nm A1, quickly place it in 37°C water bath or incubator (The microplate reader has a temperature control function that can adjust the temperature to 37°C) for 10 minutes, take it out and quickly dry it and measure the absorbance value A2 in 10min10s.Calculate $\Delta A=A2-A1$.Blank tube only need to be test one or two times.

III. Calculations:

(1) Micro glass cuvette

A. Protein concentration

Unit definition: One unit of enzyme activity is defined as the amount of enzyme produce 1 nmol NADPH per minute every milligram protein in 37°C.

GPa (U/mg prot) =[$\Delta A \times V_R \div (\varepsilon \times d) \times 10^9$]÷(V_S×Cpr) ÷T=321.54× $\Delta A \div$ Cpr

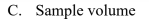
B. Sample weight

Unit definition: One unit of enzyme activity is defined as the amount of enzyme produce 1 nmol NADPH per minute every gram tissue in 37°C.

GPa (U/g weight) = $[\Delta A \times V_R \div (\varepsilon \times d) \times 10^9] \div (W \times V_S \div V_E) \div T = 321.54 \times \Delta A \div W$

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Unit definition: One unit of enzyme activity is defined as the amount of enzyme produce 1 nmol NADPH per minute every milliliter volume in 37°C.

GPa (U/mL) =[$\Delta A \times V_R \div (\epsilon \times d) \times 10^9$] $\div V_S \div T=321.54 \times \Delta A$

D. Bacteria or cells number

Unit definition: One unit of enzyme activity is defined as the amount of enzyme produce 1 nmol NADPH per minute every million bacteria or cells in 37°C.

 $GPa (U/10^{4} \text{ cells}) = [\Delta A \times V_{R} \div (\epsilon \times d) \times 10^{9}] \div (\text{cell number} \times V_{S} \div V_{E}) \div T = 321.54 \times \Delta A \div \text{cell number}$

 ε : NADPH molar extinction coefficient, 6220 L/mol/cm;

d: Cuvette light path, 1cm;

V_R: Total reaction volume, 0.2 mL;

Vs: Add sample volume,0.01mL;

V_E: extract volume, 1 mL;

W: Sample weight, g;

Cpr: Protein concentration of sample, mg/mL;

T: Reaction time, 10min

(2) 96-Well flat-bottom plates

Modify the d-1cm in the above formula to d-0.6cm (the light path of the 96-well plate) for calculation.

Note:

1. If the measured absorbance value ΔA >0.6, it is recommended to dilute the sample before measuring, and multiply the dilution factor in the calculation formula; if the measured absorbance value is low or close to the blank OD value, it is recommended to increase the sample volume before performing the measurement.

Experimental example

1. Take 0.1 g of rabbit liver tissue, add 1 mL of extract, homogenize in an ice bath, centrifuge at 8000 ×g for 10 minutes at 4°C; take the supernatant and place on ice for testing. Use micro quartz cuvette to operate according to the determination steps, $\Delta A=A2-A1=0.3472-0.2253=0.1219$, according to the formula Calculated activity: GPa(U/g weight)=[$\Delta A \times V_R \div (\epsilon \times d) \times 10^9$]÷(W ×V_S÷V_E)÷T=321.54× ΔA ÷W=391.96 U/g weight



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