

# **Anthocyanin Reductase Activity Assay Kit**

**Note:** Take two or three different samples for prediction before test. **Operation Equipment:** Spectrophotometer /Microplate Reader

**Cat No:** BC4095 **Size:**100T/48S

### **Components:**

Extract solution: 50mL×1. Storage at 4°C, shake thoroughly before use.

**Reagent I:** 25mL×1. Storage at 4°C.

**Reagent II:** Powder×2. Storage at 4°C, dissolve thoroughly with 1 ml of distilled water before use.

**Reagent III:** Powder×1. Storage at -20°C, dissolve thoroughly with 1 ml of distilled water and 1 ml of alcohol before use. It can be stored at -20°C after dispensing to avoid repeated freezing and thawing.

Reagent IV: 1mL×1. Storage at 4°C.

# **Product Description:**

Anthocyanin reductase (ANR) is a key enzyme in the biosynthesis pathway of procyanidins, which converts anthocyanins into the cis-flavan-3-alcohol. It plays an important role in plants regulation.

ANR converts cyanidin chloride to flavane-3-alcohol under the action of NADPH. The activity of ANR can be reflected by measuring the reduction rate of NADPH at 340nm.

## Reagents and Equipment Required but Not Provided:

Spectrophotometer/ microplate reader, micro quartz cuvette/96 well flat-bottom UV plate, water bath, low temperature centrifuge, adjustable transferpettor, mortar/homogenizer, ice, alcohol and distilled water.

### Sample preparation:

- 1. Tissue: Add 1 ml of extract solution into 0.1g of tissue, fully grinding on ice. Centrifuge at 12000rpm and 4°C for 15min, supernatant on ice is used for test.
- 2. Cells or microbial sample: collect cells or microbial sample to centrifuge and remove supernatant. Suggested 5 million with 1mL of extract solution, split bacteria and cells with ultrasonication (power 20%, work time 3s, interval 10s, for 30 times), 12000rpm 4°C centrifuge at 12000rpm and 4°C for 15min, supernatant on ice is used for test.

#### **Procedure:**

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

2. Add the following reagents:

Reagent name	Test tube (T)	Contrast tube (C)
Reagent I(μL)	170	170



Reagent II (μL)	10	10
Reagent III (µL)	5	5
Sample (µL)	10	- ,,0,10,

20/6 cm	Mix thoroughly at 37°C for 30 min	500
Reagent IV(µL)	5	5
Sample(µL)	COligina.	10

Mix thoroughly, detect absorbance of test tube and contrast tube at 340nm, named A(T), A(C),  $\triangle A=A(C)-A(T)=A2-A1$ .

#### Calculation:

# ultra-micro quartz cuvette

### 1. Protein concentration:

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

ANR (U/mg prot) = 
$$\Delta A \div (\epsilon \times d) \times 10^9 \times Vrv \div (V_s \times Cpr) \div T = 107.18 \times \Delta A \div Cpr$$

# 2. Sample weight:

Unit definition: One unit of enzyme activity is defined as the amount of enzyme catalyzes the decreasing of 1 nmol of NADPH per min every gram tissue in the reaction system.

ANR 
$$(U/g) = \Delta A \div (\varepsilon \times d) \times 10^9 \times V_{rv} \div (W \div V_{sv} \times V_s) \div T = 107.18 \times \Delta A \div W$$

#### 3. Cells or bacteria:

Unit definition: One unit of enzyme activity is defined as the amount of enzyme catalyzes the decreasing of 1 nmol of NADPH every 10<sup>4</sup> cells or bacteria in the r eaction system per min.

ANR (U/10<sup>4</sup>cell) =
$$\Delta A \div (\epsilon \times d) \times 10^9 \times V_{rv} \div (500 \div V_{sv} \times V_s) \div T = 0.2144 \times \Delta A$$

Vrv: total reaction volume, 0.0002 mL;

ε: NADPH molar extinction coefficient, 6.22×10<sup>3</sup>L/mol/cm;

d: light path of cuvette, 1cm;

Vs: supernatant volume (mL), 0.01 mL;

Cpr: sample protein concentration (mg/mL);

T: Reaction time (min), 30 min;

W: Sample weight(g);

Vsv: Extraction volume, 1 mL;

500: 5 million cells.

 $10^9$ : unit conversion coefficient, 1 mol =  $10^9$  nmol.

### 96 well UV plate

Change d=1cm to d=0.6cm in the formula.

Note:

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- 1. Dilute react mixture with reagent 1 or decrease sample volume if  $\triangle A>0.4$  or A(C)>1( $\triangle A>0.2$  or A(C)>1 with 96 well UV plate). Increase react time (45min or 60min) and sample volume if  $\triangle A$  is too low.
- 2. After adding reagent 4, the determination should be completed within 15 minutes.
- 3. Detect sample concentrate separately.

# **Experimental Examples:**

1. Take 0.1g of apple and add 1mL extract to homogenize and grind, take the supernatant and operate according to the measurement procedure, measure by the micro quartz cuvette and calculate  $\Delta A$ =Ac-At=0.9333-0.8095=0.1238, calculate the enzyme based on the sample weight:

ANR Activity (U/g weight) = $107.18 \times \Delta A \div W = 107.18 \times 0.1238 \div 0.1 = 132.69 \text{ U/g U/g weight}$ .

### **Related Products:**

BC1360/BC1365 Uric Acid (UA) Content Assay Kit BC1340/BC1345 Plant Total Phenol Content Assay Kit BC1330/BC1335 Plant Flavonoids Content Assay Kit



