Technical support: support@abbkine.com

Website: https://www.abbkine.com

CheKine™ Tannase (TAN) Activity Colorimetric Assay Kit

Cat #: KTB1542 Size: 48 T/96 T

FQ	Tannase (TAN) Activity Colorimetric Assay Kit		
REF	Cat #: KTB1542	LOT	Lot #: Refer to product label
	Detection range: 0.3125-20 µmoL/mL (The		Sensitivity: 0.156 µmoL/mL (The Sensitivity corres
	Detection range corresponds to the standard)		-ponds to the standard)
	Applicable samples: Plant Tissues, Fungi, Bacteria		
Ĵ	Storage: Stored at 4°C for 6 months		

Assay Principle

Tannase, full name is Tannin Acyl Hydrolase (Tannase, EC 3.1.1.20). Tannase exists in tannin-rich plants and also widely exists in microorganisms. Tannase hydrolyses the ester bonds and depside bonds in gallic acid tannins to release gallic acid and glucose. The enzyme can be produced by molds such as Aspergillus Niger, Aspergillus oryzae. It can be used to treat tannin and protein in beer to make it clear and transparent. It can also be used to remove the astringency of persimmon and other products. And it can also be used to make instant tea to prevent turbid fermented tea. CheKine™ Tannase (TAN) Activity Colorimetric Assay Kit provides a convenient tool for detection of Tannase activity. The principle is to use propyl gallate (PG) as the substrate for the enzymatic reaction of tanninase, which has a characteristic absorption peak at 270 nm. The Tannase activity measure the change of the absorbance at 270 nm before and after the reaction, and calculate the Tannase activity of the sample can be calculated by measuring the absorbance at 270 nm.

Materials Supplied and Storage Conditions

Wit a summaria	Si	ze	O	
Kit components	48 T	96 T	Storage conditions	
Extraction Buffer	100 mL	100 mL×2	4°C	
Substrate	5 mL	10 mL	4°C, protected from light	
Standard	1	2	4°C, protected from light	

Materials Required but Not Supplied

- · Microplate Reader capable of measuring absorbance at 270 nm
- · Water Bath, Ice Maker, Refrigerated Centrifuge
- 96-well UV microplate, Precision Pipettes, Disposable Pipette Tips
- Anhydrous Ethanol
- · Homogenizer (for Tissue Samples)



Version 20220421

Reagent Preparation

Extraction Buffer: Ready to use as supplied. Equilibrate to room temperature before use. Store at 4°C.

Substrate: Ready to use as supplied. Equilibrate to room temperature before use. Store at 4°C, protected from light.

Standard: Add 1.178 mL of Anhydrous Ethanol to dissolve before use. The concentration is 20 μmol/mL. This solution can be stored at 4°C for one week or stored at -20°C for long time.

Standard curve setting: Dilute 20 µmol/mL standard with Extraction Buffer as shown in the table below.

	Volume of Standard	Volume of Extraction Buffer (μL)	The concentration of Standard (μmol/mL)
Std.1	200 μL of 20 μmol/mL	0	20
Std.2	100 μL of Std.1(20 μmol/mL)	100	10
Std.3	100 μL of Std.2 (10 μmol/mL)	100	5
Std.4	100 μL of Std.3 (5 μmol/mL)	100	2.5
Std.5	100 μL of Std.4 (2.5 μmol/mL)	100	1.25
Std.6	100 μL of Std.5 (1.25 μmol/mL)	100	0.625
Std.7	100 μL of Std.6 (0.625 μmol/mL)	100	0.313

Sample Preparation

Note: Fresh samples are recommended, If not assayed immediately, samples can be stored at -80°C for one month. Processed samples must be assayed immediately. If the protein concentration of the sample is need to determined, it is recommended to use Abbkine Cat #: KTD3001 Protein Quantification Kit (BCA Assay) to measure the protein concentration of the sample.

- 1. Plant Tissues: Weigh 0.1 g tissue, add 1 mL Extraction Buffer and homogenize on ice. Centrifuge at 10,000 g for 10 min at 4°C. Use supernatant for assay, and place it on ice to be tested.
- 2. Fungi or Bacteria: Collect 5x10⁶ Fungi or Bacteria into the centrifuge tube, wash Fungi or Bacteria with cold PBS, discard the supernatant after centrifugation; add 1 mL Extraction Buffer to ultrasonically disrupt the Fungi or Bacteria in ice bath 5 min (power 20% or 200 W, ultrasonic 3 s, interval 7 s, repeat 30 times). Centrifuge at 10,000 g for 10 min at 4°C. Use supernatant for assay, and place it on ice to be tested.

Assay Procedure

- 1. Preheat the microplate reader for more than 30 min, and adjust the wavelength to 270 nm.
- 2. Add 50 µL of sample into an EP tube as a control tube, bath in boiling water for 5 min, and cool to room temperature.
- 3. Add the following reagents respectively into each EP tube :

	Blank tube (µL)	Standard tube (µL)	Test tube (µL)	Control tube (μL)
Extraction Buffer	200	150	100	100
Stds.	0	50	0	0
sample	0	0	50	50 (deactivated)
Substrate	o	0	50	50

Mix well, incubate at 40°C water bath for 10 min, and then immediately take a boiling water bath for 5 min. After cooling, centrifuge at 10,000 g at room temperature (25°C) for 10 min, and take the supernatant.

Add the following reagents to a 96-well UV microplate:



Version 20220421

supernatant	10	10	10	10
Extraction Buffer	190	190	190	190

^{4.} Mix well, read the values at 270 nm. Recorded as A_{Blank} , $A_{Standard}$, A_{Test} and $A_{Control}$, respectively. Finally, calculate $\Delta A_{Test} = A_{Control} \cdot A_{Test}$, $\Delta A_{Standard} = A_{Standard} - A_{Blank}$. (Only one blank well needs to be detected, a control is required for each sample).

Note: In order to guarantee the accuracy of experimental results, need to do a pre-experiment with 2-3 samples. If ΔA_{Test} is greater than 1.0, the sample can be appropriately diluted with Extraction Buffer, the calculated result multiplied by the dilution factor. If ΔA_{Test} is less than 0.001, increase the sample quantity appropriately.

Data Analysis

Note: We provide you with calculation formulae, including the derivation process and final formula. The two are exactly equal. It is suggested that the concise calculation formula in bold is final formula.

1. Drawing of standard curve

With the concentration of the standard Solution as the y-axis and the $\Delta A_{Standard}$ as the x-axis, draw the standard curve. Substitute the ΔA_{Test} into the equation to obtain the y value (μ mol/mL).

- 2. Calculate the activity of Tannase (TAN)
- (1) By sample weight

Unit Definition: 1 nmol PG reduced per min in 1 g tissue reaction system at 40°C is defined as a unit of enzyme activity.

TAN $(U/g)=y\times1,000\times F\times V_{Reaction\ Total}\div (W\times V_{Sample}\div V_{Sample\ Total})\div T\times n=8,000\times y\div W\times n$

(2) By cells number of Fungi or Bacteria

Unit Definition: 1 nmol PG reduced per min in 10⁴ cells number of Fungi or Bacteria reaction system at 40°C is defined as a unit of enzyme activity.

TAN (U/10⁴ cells)=y×1,000×F×V_{Reaction Total}÷(V_{Sample}×500÷V_{Sample Total})÷T×n=16×y×n

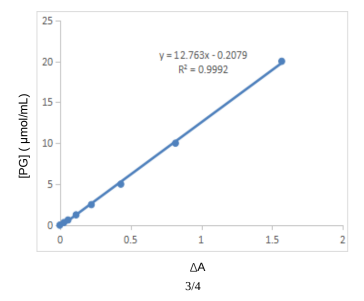
(3) By protein concentration

Unit Definition: 1 nmol PG reduced per min in 1 mg tissue protein reaction system at 40° C is defined as a unit of enzyme activity. TAN(U/mg prot)=yx1,000xFxV_{Reaction Total}÷(Cpr xV_{Sample})÷T×n=8,000xy÷Cprxn

Where: 1,000: 1 μ mol=1,000 nmol; F: Dilution factor of supernatant, F=200 μ L÷10 μ L=20; V_{Reaction Total}: total reaction volume, 0.2 mL; W: sample weight, g; V_{sample}: sample volume added, 0.05 mL; V_{sample Total}: Extraction Buffer volume added, 1 mL; T: reaction time, 10 min; n: dilution multiple of sample; 500: Total cells number of Fungi or Bacteria, 5×10⁶; Cpr: sample protein concentration, mg/mL.

Typical Data

Typical standard curve





Version 20220421

Figure 1. Standard Curve of PG in 96-well plate assay-data provided for demonstration purposes only. A new standard Curve must be generated for each assay.

Recommended Products

Catalog No.	Product Name
KTB1540	CheKine™ Plant Total Phenols (TP) Colorimetric Assay Kit
KTB1530	CheKine™ Plant Flavonoids Colorimetric Assay Kit
KTB1541	CheKine™ Tannin Colorimetric Assay Kit

Disclaimer

The reagent is only used in the field of scientific research, not suitable for clinical diagnosis or other purposes.

