

Butyrylcholinesterase Inhibitor Activity Assay Kit

Detection Equipment: Spectrophotometer/Microplate Reader

Catalog Number: BC5985

Size: 100T/96S

Components: Please carefully check the volume of the reagent and the volume in the bottle before use. If you have any questions, please contact Solarbio staff in time.

Reagent name	Size	Preservation condition	
Extract	Liquid 120 mL×1	2-8°C storage	
Reagent I	Powder ×1 -20°C storage		
Reagent II	Liquid 15 mL×1	2-8°C storage	
Reagent III	Liquid 12 mL×1	2-8°C storage	
Reagent IV	Powder ×1	-20°C storage	
Reagent V	Liquid 1 mL×1	1 mL×1 -20°C storage	

Solution reparation:

- 1. Reagent I: Before use, take 1 Reagent I, add 1.2mL distilled water, fully dissolve, and storage the reagent for 4 weeks after subpacking at -20°C to avoid repeated freeze-thaw (The reagent is a freeze-dried reagent, there may be a large difference or even a small amount of macroscopic observation between different bottles, this phenomenon does not affect the use, the actual quality is the same).
- 2. Reagent IV: Add 6mL Reagent II before clinical use, fully dissolve, and the reagent can be storage for 4 weeks at -20°C to avoid repeated freezing and thawing.
- 3. Reagent V:10mmol/L Livansmin solution. Before clinical use, 15µL 10mmol/L Livansmin solution was taken, and 985µL distilled water was added to prepare 0.15mmol/L Livansmin solution. (This reagent was used for positive tube experiment, and it was selected.)

Description:

Butyrylcholinesterase (BchE, EC3.1.1.8), also known as plasma cholinesterase, pseudocholinesterase, is a serine hydrolase that is synthesized by the liver and enters the blood and is present in almost all animal tissues. Compared with acetylcholinesterase (AchE), BchE can effectively hydrolyze larger choline esters, such as butyrylcholine and benzoylcholine, and can remove the toxic effects of nerve agents such as organophosphorus pesticides and carbamate pesticides. Studies have shown that BchE can be used as an important target for the treatment of Alzheimer's disease, and BchE inhibitors are used to improve memory loss and cognitive dysfunction in Alzheimer's patients.

BchE catalyzes the hydrolysis of butyrylcholine to choline, and the reaction of choline with dithio-nitrobenzoic acid (DTNB) to produce 5-merhydryl-nitrobenzoic acid (TNB). BchE inhibitors reduce the hydrolysis of butyrylcholine by inhibiting BchE activity. TNB has an absorption peak at 412nm, and the BchE inhibitor activity can be calculated by measuring the change of absorbance at 412nm.



Butyrylcholine

BchE Choline

DTNB TNB (412nm)

Note: Before the experiment, it is recommended to select 2-3 sample with large expected differences for pre-experiment.

Reagents and Equipment Required but Not Provided:

Spectrophotometer/microplate reader, balance, oven, pulverizer/mortar/homogenizer, 30-50 mesh screen, ultrasonic cleaning machine, centrifuge, water bath/constant temperature incubator, micro glass cuvette/96 well flat-bottom plate, adjustable pipette, ice and distilled water.

Protocol:

- I. Sample processing (The sample size to be tested can be appropriately adjusted, and the specific proportion can be referred to the literature)
- 1. Tissue sample: The sample was dried to constant weight, crushed, after 30-50 mesh sieve, weighed about 0.1g, added 1mL of extraction solution, and extracted by ultrasonic extraction method, ultrasonic power 300W, 60°C, extraction for 30min. 12000rpm, 25°C, centrifuge for 10min, take the supernatant, and volume with the extraction solution to 1mL, to be measured.
- **2. Powder sample:** Take an appropriate amount of sample, add an appropriate amount of Extract , and prepare a solution with appropriate concentration to be measured.

Note: If the powder sample is insoluble in Extract, it can be dissolved with a suitable solvent, prepared into a solution of $100\times$ or greater concentration, and then diluted to $1\times$ concentration with the extraction solution.

II. Measurement Steps

1. Spectrophotometer/microplate reader for more than 30min, adjust the wavelength to 412nm, and zero the distilled water.

2. Operation table: (Add the following reagents in micro glass cuvette/96 well flat-bottom plate)

Reagent name (µL)	Blank tube 1	Blank tube 2	Test tube	Positive tube (optional)
Reagent I	10	- 9	10	10
Reagent II	50	60	50	50
Extract	10	10	-	_ C_O/6 2018
Sample) -	10	(C) - V
Reagent V	-191, Pur	-	-	10
	Mix well and incub	pate at 25°C for 10	min in dark	
Reagent III	100	100	100	100
Reagent IV	50	50	50	50

Thoroughly mixed, the absorbance value A at 10s was measured at 412nm, and the absorbance value A 'at 5min10s was measured accurately at 37°C for 5min(If the enzyme marker has a temperature control function, adjust the temperature to 37°C), and was recorded as A blank 1, A $_{\text{blank}}$ 2, A $_{\text{text}}$, A $_{\text{positive}}$ and A' $_{\text{blank}}$ 1, A ' $_{\text{blank}}$ 2, A' $_{\text{text}}$, A ' $_{\text{positive}}$, respectively. Calculate ΔA_{blank} = (A ' $_{\text{blank}}$ 1-A $_{\text{blank}}$ 1) - (A' $_{\text{blank}}$ 2-A $_{\text{blank}}$ 2), ΔA_{text} = (A ' $_{\text{text}}$ -A $_{\text{text}}$) - (A' $_{\text{blank}}$ 2-A $_{\text{blank}}$ 2). Blank tubes 1 and 2 only need



to be done 1-2 times.

III. Calculations

1. Calculation of inhibition rate

Butyryl cholinesterase inhibitors Inhibition ratio (%) = $(\Delta A_{blank} - \Delta A_{text}) \div \Delta A_{blank} \times 100\%$.

2.IC₅₀ calculation

 IC_{50} , the semi-inhibitory concentration of the inhibitor. For the samples that are determined to have inhibition on BchE, appropriate concentration gradients can be formulated, and the sample concentration is taken as the horizontal coordinate and the inhibition rate as the vertical coordinate as the inhibition curve, so as to calculate the sample concentration when the inhibition rate is 50%, that is, IC_{50} .

Note:

- 1. In order to ensure the accuracy and stability of the experimental results, please strictly control the reaction time and operation time.
- 2. When the absorbance of the sample is greater than 1.6 or the ΔA determination is close to the ΔA blank, it is recommended to increase the proportion of tissue samples in the sample processing step or prepare the powder sample into a higher concentration solution before the determination. When the ΔA determination is less than 0.004, the sample can be diluted with the extraction solution before the determination.
- 3. If it is used to compare the degree of inhibition of BchE by different reagents, extracts, drugs or tissues, the reagent, extract, drug or tissue homogenate must be prepared to the same concentration for comparison.

Experimental example:

- Take 0.1005g pomelo peel sample (dried), add 1mL Extract solution for ultrasonic extraction, centrifuge and take supernatant, according to the measurement procedure, and use 96 well flat-bottom plate to measure and calculate: ΔA _{blank} = (A '_{blank 1}-A _{blank 1}) (A' _{blank 2}-A _{blank 2}) = (1.129-0.133)-(0.141-0.120)=0.975, ΔA _{text} = (A '_{text} -A _{text}) (A' _{blank 2}-A _{blank 2}) = (0.878-0.337) (0.141-0.120) = 0.520, calculated inhibition rate:
 - BchE inhibitor inhibition rate (%) = $(0.975-0.520) \div 0.975 \times 100\% = 46.667\%$.
- 2. Take 0.1014g persimmon peel sample (dried), add 1mL Extract solution for ultrasonic extraction, centrifuge and obtain supernant, dilute 16 times according to the measurement procedure, and use 96 well flat-bottom plate to measure and calculate: ΔA _{blank} = (A' _{blank 1}-A _{blank 1}) (A' _{blank 2}-A _{blank 2}) = (1.129-0.133)-(0.141-0.120)=0.975, ΔA _{text} = (A' _{text} -A _{text}) (A' _{blank 2}-A _{blank 2}) = (0.581-0.127) (0.141-0.120) = 0.433, the inhibition rate is calculated as follows:
 - BchE inhibitor inhibition rate (%) = $(0.975-0.433) \div 0.975 \times 100\% = 55.590\%$.
- 3. Take $10\mu L$ 0.15mmol/L Livansmin solution, follow the measurement steps, and use 96 well flat-bottom plate to measure and calculate: $\Delta A_{blank} = (A'_{blank} _1 A_{blank} _1) (A'_{blank} _2 A_{blank} _2) = (1.129-0.133) (0.141-0.120) = 0.975$, $\Delta A_{text} = (A'_{text} A_{text}) (A'_{blank} _2 A_{blank} _2) = (0.631-0.126)$
 - (0.141-0.120) =0.484, the inhibition rate is calculated as follows:
 - BchE inhibitor inhibition rate (%) = $(0.975-0.484) \div 0.975 \times 100\% = 50.359\%$.



References:

- [1] Ellman GL, Courtney KD, Andres V Jr. et al. A new and rapid colorimetric determination of acetylcholinesterase activity [J]. Biochemical Pharmacology, 1961, 7(2): 88-95.
- [2] Noor Atatreh, Sara Al Rawashdah, Shaikha S Al Neyadi. et al. Discovery of new butyrylcholinesterase inhibitors via structure-based virtual screening [J]. Journal of Enzyme Inhibition and Medicinal Chemistry, 2019, 34(1): 1373-1379.
- [3] Li Q, Chen Y, Xing S. et al. Highly Potent and Selective Butyrylcholinesterase Inhibitors for Cognitive Improvement and Neuroprotection [J]. Journal of Medicinal Chemistry, 2021, 64(10): 6856-6876.

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