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# Soil Neutral Xylanase (S-NEX) Activity Assay Kit

Note: Take two or three different samples for prediction before test.

Detection equipment: Spectrophotometer/Microplate reader

Cat No: BC5735

Size: 100T/48S

#### **Components:**

Buffer Fluid: Liquid 25 mL×1, store at 2-8°C.

Reagent I: Liquid 12 mL×1, store at 2-8°C.

**Reagent II:** Liquid 13 mL×1, store at 2-8°C.

**Standard:** Powder×1, 10mg xylose. Before use, a standard solution of 100 $\mu$ mol/mL was prepared by adding 667 $\mu$ L distilled water and stored at 2-8°C for 8 weeks.

# **Product Description:**

Soil neutral xylanase (S-NEX), also known as soil neutral hemicellulase, is mainly isolated from microorganisms with an optimal growth pH of 6-8.

In a neutral environment, S-NEX catalyzes the degradation of xylan into reducing oligosaccharides and monosaccharides. Under boiling water bath conditions, it further undergoes a color reaction with 3,5-dinitrosalicylic acid, with a characteristic absorption peak at 540nm. The color depth of the reaction solution is directly proportional to the amount of reducing sugars produced by enzymatic hydrolysis. By measuring the rate of increase in absorbance of the reaction solution at 540nm, S-NEX activity can be calculated.

# **Required but not provided:**

Spectrophotometer/Microplate reader, micro glass cuvette/96 well plate, balance, desk centrifuge, water bath, 30-50 mesh sieve, distilled water.

# **Procedure:**

I. Sample preparation(The sample size to be tested can be adjusted appropriately, and the specific proportion can be referred to in the literature)

Fresh soil samples are naturally air dried or air dried in a 37 °C oven, and sieved through a 30-50 mesh sieve.

# **II. Determination procedure**

1. Preheat spectrophotometer/microplate reader for 30 minutes, adjust wavelength to 540 nm, set the counter to zero with distilled water.

2. Dilution of standard solution: Using distilled water to dilute the Standard into 2, 1.5, 1.2, 1, 0.8, 0.4, 0.2µmol/mL of standard solution before measured.

3. Quasi-dilution table:							
Num-	Predilution concentration	Standard volume	Volume of distilled	Diluted concentration			
			Oin				

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ber	(µmol/mL)	(µL)	water (µL)	(µmol/mL)
1	100	100	900	10 💿
2	10	200	800	2
3	10	150	850	1.5
4	50 10	120	880	1.2
5	10	100	900	1
6	1 (	200	50	0.8
7	1	100	150	0.4
	1	50	200	0.2

Note:  $150\mu L$  per tube is required in the experiment.

4. Sample determination (add the following reagents in 1.5 mL EP tube in turn).

Reagent Name (µL)	Control tube (A <sub>C</sub> )	Test tube (A <sub>T</sub> )	Blank tube (A <sub>B</sub> )	Standard tube (A <sub>S</sub> )
Sample	0.05g	0.05g	- "	-
Buffer Fluid	200	200	-	-
Reagent I	Sylter	100	<u> </u>	-
Mix well and plac	e in a 50°C water bath	10 LOES		
Immediately after, heat	in a boiling water bath	Olesolen	(i)	
deactivate (be careful	not to let the lid burst o	- 74	Sig	
water from entering ar	nd altering the reaction s		- OLAL DENCE	
SOLESONE I	oom temperature.	(I)	SUFES	
Reagent I	100	) .e <sup>2</sup> -	-	
At room temperature, c	centrifuge at 12000g for			
ta	ike the supernatant		-	
Supernatant	150	150	1 St Prof	-
Standard	-		Orreson -	150
Distilled water	_	- ('C.)	150	1010E5
Reagent II	100	100	100	100

Mix well, accurately color in a boiling water bath for 5 minutes (be careful not to let the lid burst open to avoid water entering and altering the reaction system), cool to room temperature. Take 200 $\mu$ L the supernatant and measure the absorbance at 540nm in 96 well plate or micro glass cuvette. Record the absorbance values as A<sub>C</sub>, A<sub>T</sub>, A<sub>B</sub>, A<sub>S</sub>, calculate  $\Delta$ A<sub>T</sub> = A<sub>T</sub> - A<sub>C</sub>,  $\Delta$ A<sub>S</sub> = A<sub>S</sub> - A<sub>B</sub>. The blank tube and standard curve only need to be measured 1-2 times. Each test tube should have a corresponding control tube.

#### **III.** Calculation

1. Drawing of standard curve:

The standard curve is established according to the concentration of the standard tube (x, µmol/mL)



and the absorbance  $\Delta A_S$  (y,  $\Delta A_S$ ). According to the standard curve, the  $\Delta A_T$  (y,  $\Delta A_T$ ) is brought into the formula to calculate the sample concentration (x,  $\mu$ mol/mL).

## 2. Calculation of soil S-NEX activity:

Enzyme activity definition: An enzyme activity unit of a neutral xylanase is defined as the amount of enzyme required to produce 1  $\mu$ mol of reducing sugar per gram of soil per hour by decomposing xylan at 50°C and pH 7.0.

S-NEX activity (U/g soil) =  $x \times V_{RV} \div W \div T \times F = 0.15 \times x \div W \times F$ 

V<sub>RV</sub>: Total reaction volume, 0.3mL;

W: Sample mass, g;

T: Reaction time, 2h;

F: Sample dilution ratio;

## Note:

- 1. If  $\Delta A_T$  is less than 0.01, the sample size can be appropriately increased or the reaction time can be extended by 50 °C before measurement; If the  $\Delta A_T$  is greater than 1.5 or  $A_T$  is greater than 1.5, the supernatant can be diluted with distilled water before measurement, and attention should be paid to synchronously modifying the dilution factor in the calculation formula.
- 2. It is recommended to use a spiral tube to prevent the lid from bursting during the boiling water bath process and change the reaction system.

## **Experimental example:**

Take 0.05g of wild mushroom soil, perform the first reaction according to the operation table for 2 hours, then dilute the supernatant twice with distilled water and follow the measurement steps. Measure with a 96 well plate, calculate  $\Delta A_T = A_T - A_C = 0.891 - 0.200 = 0.691$ , bring in the standard curve y=0.7418x-0.2302(R<sup>2</sup>=0.9992), calculate x=1.242, and calculate S-NEX activity based on sample mass:

S-NEX activity (U/g soil) = $0.15 \times x \div W \times F = 7.452$  U/g soil.

# **Related Products:**

BC5720/BC5725Soil Acid Xylanase (S-ACX) Activity Assay KitBC5740/BC5745Soil Alkaline Xylanase (S-BAX) Activity Assay Kit

