

#### (FOR RESEARCH USE ONLY, DO NOT USE IT IN CLINICAL DIAGNOSIS!)

## SAL(Salbutamol)ELISA Kit

Catalog No:E-FS-E017 96T

This manual must be read attentively and completely before using this product.

If you have any problems, please contact our Technical Service Center for help.

Phone: 240-252-7368(USA)240-252-7376(USA)

Email: techsupport@elabscience.com

Website: www.elabscience.com

Please kindly provide us the lot number(on the outside of the box) of the kit for more efficient service.

# Test principle

This kit uses Indirect-Competitive-ELISA as the method. It can detect SAL in samples, such as urine, tissue, fodder, etc. This kit is composed of Micro ELISA Plate, HRP conjugate, antibody, standard and other supplementary reagents. The micro-plate provided in this kit has been pre-coated with SAL. During the reaction, SAL in the samples or standard competes with SAL on the solid phase supporter for sites of SAL antibody. Then Horseradish Peroxidase (HRP) conjugate is added to each micro plate well, and TMB substrate is for color development. There is a negative correlation between the OD value of samples and the concentration of SAL. You can calculate the concentration of SAL in the samples by comparing the OD of the samples to the standard curve.

#### **Technical indicator**

Sensitivity:0.1ppb(ng/mL)

**Reaction mode:** 25°C,30min~15min **Detection limit:** Tissue/urine---0.1ppb

Fodder---1ppb

Cross-reactivity: Salbutamol---100%

Dobutamine Hydrochloride---7.2%

Cimaterol ---0.1% Clenbuterol---3.6% Ractopamine ---0.1% Epinephrine ---0.1% Isoprenaline---<1%

Sample recovery rate: Urine ---90%  $\pm$  10%, Tissue---80%  $\pm$  10%, Fodder---80%  $\pm$  15%

### **Kits components**

Item	Specifications
ELISA Micro-plate	96 wells
Standard Liquid	1mL each (0ppb,0.1ppb,0.3ppb,0.9ppb,2.7ppb,8.1ppb)
High ConcentratedStandard (100ppb)	1mL
HRP Conjugate(Red cap)	5.5mL
Antibody Working Solution (Blue cap)	5.5mL
Substrate Reagent A (White cap)	6mL
Substrate Reagent B(Black cap)	6mL
Stop Solution(Yellow cap)	6mL
20×Concentrated Wash Buffer(White cap)	40mL
10×Redissolved Buffer(Yellow cap)	50mL
Product Description	1 сору

# Other supplies required

Instruments: Micro-plate reader, Printer, Homogenizer, Nitrogen Evaporators, Oscillators, Centrifuge,

Graduated pipette, Balance(sensibility 0.01g).

High-precision transferpettor: single channel(20-200μL, 100-1000μL), Multichannel(300μL).

Reagents: sodium hydroxide, Ethyl acetate, concentrated HCl, acetonitrile, isopropanol,

n-hexane, sodium sulphate anhydrous.

## **Experimental preparation**

Bring all reagents and samples to room temperature before use.

Open the micro-plate reader in advance, preheat the instrument, and set the testing parameters.

1. **Sample pretreatment Notice:** experimental apparatus should be clean, and the pipette should be disposable to avoid cross-contamination during the experiment.

### 2. Solution preparation

Solution 1: 1M HClsolution

Dissolve 8.6 mL concentrated HCl to 100mL with deionized water

Solution 2: 1M NaOH solution

Dissolve4g NaOH to 100mL with deionized water

Solution 3:Redissolved Buffer

Dilute the  $10\times$ Redissolved Buffer with deionized water at dilution of 1:9, for sample re-dissolution. It can be stored at  $4^{\circ}$ C for one month.

Solution 4: 1×Working Wash Buffer

Dilute the 20×Concentrated Wash Buffer with deionized waterat dilution of 1:19.

### 3. Sample pretreatment procedure

#### 3.1 Pretreatment ofurine:

Take 50uL of clear urine sample to detect (the turbid urine sample should be filtered or centrifuge at 4000r/minfor 5min to get clear urine sample). Temporarily used samples should be kept frozento save.

Note: Sample dilution factor: 1, minimum detection dose: 0.1ppb

## 3.2 Pretreatment1 oftissue:

Weigh  $2\pm0.05$ g of homogeneous tissue samples, add 6mL of Redissolved Buffer, oscillatethoroughlyfor 2min, centrifugeat 4000r/min at room temperature for 10 min(if the lipid content of tissue samples is high, it's suggested to incubate at  $85\,^{\circ}$ C with water bath for 10 minafter oscillating, then centrifuge). Take  $50\mu$ Lupper liquid to analyze.

Note: Sample dilution factor: 4, minimum detection dose: 0.4ppb

### 3.3Pretreatment 2 oftissue:

(1) Weigh  $2\pm0.05$ g of homogeneous tissue samples,, add 1mL of Redissolved Buffer, oscillate

- thoroughly, add 4mL of acetonitrile, oscillate thoroughly, add 1mLof isopropanol, oscillate thoroughly for 5min, centrifugeat 4000r/min at room temperature for 10 min.
- (2) Take 3mL of upper liquid, add  $50\mu$ L of 1M NaOH, add 7mL of Ethyl acetate, oscillate thoroughly for 5 min, centrifugeat 4000r/min at room temperature for 10 min. Take all upper liquid to blow-dry at  $56\,^{\circ}$ C nitrogen or air.
- (3) Dissolve the dried residual with 1mLof Redissolved Buffer, add 1mL of n-hexane, mix 30s; centrifuge at 4000r/min at 15℃ for 5 min.
- (4) Take 50µLlower liquid to analyze.

Note: Sample dilution factor: 1, minimum detection dose: 0.1ppb

#### 3.4 Pretreatment offodder:

- (1) Weigh1±0.05g of homogeneousfoddersample, add 10mL of methyl alcohol and 5g of sodium sulphate anhydrous, oscillate for 2min, Centrifuge at 4000r/min at room temperature for 10min.
- (2) Take 1mL of upper liquidafter centrifugation, blow-dryat 56°C nitrogen, dissolve the dried residual with 1mLof Redissolved Buffer, add 1mL of n-hexane, mix 30s; Centrifuge at 4000r/min at room temperature for 5 min.
- (3) Take 50uLlower liquid to analyze.

Note: Sample dilution factor: 10, minimum detection dose: 1ppb

## Assav procedure

Centrifuge the sample again after thawing before the assay. Bring all reagents to room temperature before use. All the reagents should be mixed thoroughly by gently swirling before pipetting. Avoid foaming. Dilute 40mL of concentrated wash buffer into 800mL wash working buffer with deionized or distilled water.

- 1. Number: number the sample and standard in order(multiplewell), and keep a record of standard wells and sample wells.
- 2. Add sample: add  $50\mu\text{L}$  of Standard, Blank, or Sample per well, then add  $50\mu\text{L}$  of HRP conjugate to each well, then add  $50\mu\text{L}$  of antibody working solution, cover the plate with sealer we provided, oscillate for 5s gently to mix thoroughly, incubate for 45min at  $25\,^{\circ}\text{C}$ .
- 3. Wash: uncover the sealer carefully, remove the liquid in each well. Immediately add 250μL of wash working buffer to each well and wash. Repeat wash procedure for 5 times, 30s intervals/time. Invert the plate and pat it against thick clean absorbent paper (If bubbles exist in the wells, clean tips can be used to prick them).
- **4. HRP conjugate:**add 100μL HRP conjugate to each well, shading light incubation for 30min at 25°C.
- **5.** Wash: repeat step 3
- 6. Color Development: add  $50\mu$ L of substrate solution A to each well, and then add  $50\mu$ L of substrate solution B. Gently oscillate for 5s to mix thoroughly. Incubate shading light for 15min at  $25^{\circ}$ C.
- 7. Stop reaction: add 50 µL of stop solution to each well, oscillate gently to mix thoroughly.
- 8. OD Measurement: determine the optical density(OD value) of each wellat 450 nm with a

microplate reader (the 450/630 nm double wavelength is recommended). This step should be finished in 10min after stop reaction.

# Result analysis

### 1. Absorbance (%)= $A/A_0 \times 100\%$

A: Average absorbance of standard or sample

A<sub>0</sub>: Average absorbance of 0 ppb Standard

### 2. Drawing and calculation of standard curve

Create a standard curve by plotting the absorbance percentage of each standard on the y-axis against the log concentration on the x-axis to draw a semi-logarithmic plot. Add average absorbance value to standard curve to get corresponding concentration. If samples have been diluted, the concentration calculated from the standard curve must be multiplied by the dilution factor.

For this kit, it is more convenient to use professional analysis software for accurate and fast analysis on a large number of samples.

#### **Notes**

- 1. Overall OD value will be lower when reagents is not brought to room temperature before use or room temperature is below25°C.
- 2. During the washing procedure, if the wells turn dry, it will lead to bad linear standard curve and poor repeatability, move on to the next step immediately after wash.
- 3. Mix thoroughly and wash the plate completely. The consistency of wash procedure can strongly affect the reproducibility of this ELISA kit.
- 4. Micro ELISA plate should be covered by plate sealer. Avoid the reagents to strong light.
- 5. Do not use expired kit and reagents of different batches.
- 6. TMB should be abandoned if it turns color. When OD value of standard(concentration: 0)<0.5 unit( $A_{450nm}<$ 0.5), it indicates reagent is deteriorated.
- 7. Stop solution is caustic, avoid contact with skin and eyes.

## Storage and valid period

**Storage**: Store at 2-8°C. Avoid freeze / thaw cycles.

**Valid Period**: 1 year, production date is on the packing box.