Vinculin ELISA Kit

Support: service@enogene.com



Assay range: 2 ng/ml -85 ng/ml

96 Tests

Introduction

This kit allows for the determination of Vinculin concentrations in Human serum, plasma and other biological fluids

Principle of the assay

The kit assay Human Vinculin level in the sample, use Purified Human Vinculin antibody to coat microtiter plate wells, make solid-phase antibody, then add Vinculin to wells, Combined Vinculin antibody which With HRP labeled, become antibody - antigen - enzyme-antibody complex, after washing Completely, Add TMB substrate solution, TMB substrate becomes blue color At HRP enzyme-catalyzed, reaction is terminated by the addition of a sulphuric acid solution and the color change is measured spectrophotometrically at a wavelength of 450 nm. The concentration of Human Vinculin in the samples is then determined by comparing the O.D. of the samples to the standard curve.

Components

1	Wash solution	20ml×1bottle	7	Stop Solution	6ml×1 bottle
2	HRP-Conjugate reagent	6ml×1 bottle	8	Standard (160ng/ml)	0.5ml×1 bottle
3	Microelisa strip plate	12 well×8 strips	9	Standard diluent	1.5ml×1 bottle
4	Sample diluent	6ml×1 bottle	10	Instruction	1
5	Chromogen Solution A	6ml×1 bottle	11	Closure plate membrane	2
6	Chromogen Solution B	6ml×1 bottle	12	Sealed bags	1

Specimen requirements

- 1. Extract as soon as possible after Specimen collection, and according to the relevant literature, and should be experiment as soon as possible after the extraction. If it can't, specimen can be kept in -20 °C to preserve, Avoid repeated freeze-thaw cycles.
- 2. Can't detect the sample which contain NaN3, because NaN3 inhibits HRP active.

For Research Use Only

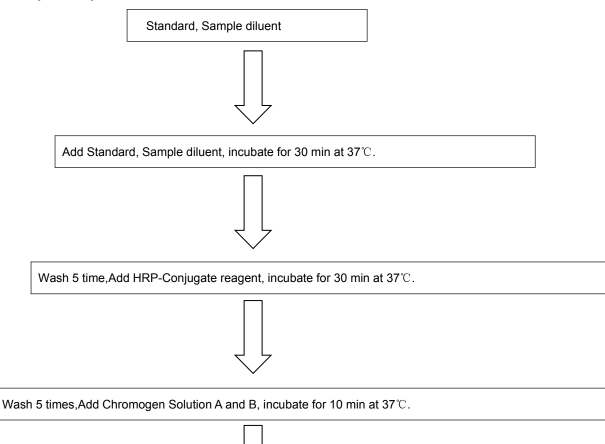
Assay procedure

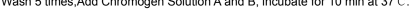
1. Dilute and add sample: Dilute Original density Standard as follow table:

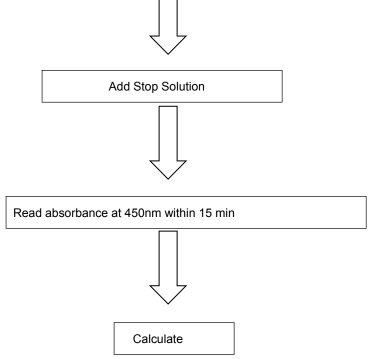
80ng/ml	5 Standard	150µl Original density Standard+150µl Standard diluent
40ng/ml	4 Standard	150µl 5 Standard+150µl Standard diluent
20ng/ml	3 Standard	150μl 4 Standard+150μl Standard diluent
10ng/ml	2 Standard	150µl 3 Standard +150µl Standard diluent
5ng/ml	1 Standard	150µl 2 Standard +150µl Standard diluent

- 2.Add sample: Set blank wells separately (blank comparison wells don't add sample and HRP-Conjugate reagent, other each step operation is same). testing sample well. Add 50µl of standard to Microelisa strip plate, add Sample dilution 40µl to testing sample well, then add testing sample 10µl (sample final dilution is 5-fold), add sample to wells, don't touch the well wall as far as possible, and Gently mix.
- 3.Incubate: After closing plate with Closure plate membrane ,incubate for 30 min at 37° C.
- 4.Prepare liquid: 30-fold (or 20-fold) wash solution diluted 30-fold (or 20-fold) with distilled water and reserve.
- 5. Washing: Uncover Closure plate membrane, discard Liquid, dry by swing, add washing buffer to every well, still for 30s then drain, repeat 5 times, dry by pat.
- 6.Add enzyme: Add HRP-Conjugate reagent 50µl to each well, except blank well.
- 7.incubate: Operation with 3.
- 8. Washing: Operation with 5.
- 9.Color reaction: Add Chromogen Solution A 50ul and Chromogen Solution B 50ul to each well, evade the light preservation for 10 min at 37°C
- 10.Stop the reaction: Add Stop Solution50µl to each well, Stop the reaction(the blue color change to yellow color).
- Detection: take blank well as zero , Read absorbance at 450nm after Adding Stop Solution and within
 15min.

Steps description







Calculate

Take the standard density as the horizontal, the OD value for the vertical ,draw the standard curve on graph paper, Find out the corresponding density according to the sample OD value by the Sample curve,

For Research Use Only

multiplied by the dilution multiple, or calculate the straight line regression equation of the standard curve

with the standard density and the OD value ,with the sample OD value in the equation, calculate the

sample density, multiplied by the dilution factor, the result is the sample actual density.

Notes

1. The kit takes out from the refrigeration environment should be balanced 1 hour in the room

temperature, ELISA plates coated if has not use up after opened, the plate should be stored in

Sealed bag.

2. Washing buffer will Crystallization separation, it can be heated the water helps dissolve when dilute .

Washing does not affect the result.

3. Add Sample with sampler Each step, And proofread its accuracy frequently, avoids the experimental

error. add sample within 5 min, if the number of sample is much, recommend to use Volley.

4. If the testing material content is excessively higher (The sample OD is bigger than the first standard

well), please dilute Sample (n-fold), Please diluente and multiplied by the dilution factor. (×n×5) .

Closure plate membrane only limits the disposable use, to avoid cross-contamination.

The substrate evade the light preservation.

7. Please according to the instruction strictly. The test result determination must take the microtiter

plate reader as a standard.

8. All samples, washing buffer and each kind of reject should according to infective material process.

Do not mix reagents with those from other lots.

Storage and validity

1. Storage: 2-8°C.

2. Validity: Six months

For Research Use Only