

**(FOR RESEARCH USE ONLY. DO NOT USE IT IN CLINICAL DIAGNOSIS !)**

**Human Japanese Encephalitis Virus (JEV) IgG ELISA Kit**

Catalog No: GEH039

96T

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

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

Tel: 400-967-3365

Email: [techsupport@uni-science.com](mailto:techsupport@uni-science.com)

Website: [www.elabscience.cn](http://www.elabscience.cn)

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

## Test principle

This ELISA kit uses Indirect-ELISA as the principle to detect the Japanese Encephalitis Virus (JEV) IgG in human serum. The ELISA Microtiter plate provided in this kit has been pre-coated with purified Japanese Encephalitis Virus antigen. Samples are added to the ELISA Microtiter plate wells and the JEV-IgG in which will combine with the pre-coated antigen to form antigen-antibody compound. Free components are washed away. The HRP conjugated Mouse-anti-human IgG antibody is added to each well and react with the compound to form “antigen- antibody-HRP antibody” compound. The TMB substrate is added to initiate the color developing reaction. The presence of JEV-IgG can be determined according to the OD value after colorimetric assay with the Micro-plate Reader.

## Kit components

| Item                            | Specifications |
|---------------------------------|----------------|
| ELISA Microtiter plate          | 96 wells       |
| Positive Control                | 1 mL           |
| Negative Control                | 1 mL           |
| HRP Conjugated Working Solution | 12 mL          |
| Sample Diluent                  | 12 mL          |
| 20×Concentrated Wash Buffer     | 50 mL          |
| Substrate Reagent A             | 6 mL           |
| Substrate Reagent B             | 6 mL           |
| Stop Solution                   | 6 mL           |
| Plate Sealer                    | 3 pieces       |
| Sealed Bag                      | 1 piece        |
| Manual                          | 1 copy         |

## Other materials required but not supplied

Micro-plate Reader with 450 nm wavelength filter or dual-wavelength (450/630 nm)

High-precision transferpettor, EP tubes and disposable pipette tips

37°C Incubator or water bath

Deionized water

Absorbent paper

Loading slot for Wash Buffer

**Notes**

1. Wear gloves and work clothes during experiment, and the disinfection and isolation system should be strictly executed. All the waste should be handled as contaminant.
2. The Stop Solution is corrosive, it should be avoided to contact with skin and clothing. Wash immediately with plenty of water if contact it carelessly.
3. The ELISA Microtiter plate obtained from cold storage conditions should be adjusted to room temperature before use. The unused plate should be kept in a sealed bag with desiccant.
4. 20×Concentrated Wash Buffer at low temperature condition is easy to crystallize, it should be adjusted to room temperature to dissolve completely before use.
5. Each well must be filled with liquid when washing to prevent residual free enzyme.
6. The tested sample should be kept fresh.
7. The results shall depend on the readings of the Micro-plate Reader.
8. Do not use components from different batches of kit.

**Storage and expiry date**

Store unopened at 2-8°C. Do not freeze.

Please store the opened kit at 2-8°C, protect from light and moisture. The shelf life of the opened kit is up to 1 months.

**Expiry date:** expiration date is on the box.

**Sample preparation**

1. **Serum:** Human serum can be used as detected sample. Fresh collected serum samples should be fully centrifuged, then take clear liquid for test. Suspended fibrous protein may cause a false positive.
2. Anticoagulant (such as EDTA, sodium citrate and heparin, etc.) in samples do not affect the results.
3. Avoid of samples with suspended fibrous protein, aggregation or severe hemolysis.  
(hemoglobin > 10 g/L), hyperlipemia (triglyceride >20 g/L), high bilirubin (bilirubin > 0.2 g/L). Obviously contaminated samples can't be detected.
4. Samples can be stored at 2-8°C for one week. If samples not tested in a week, store them below -15°C no more than a month, and avoid freeze-thaw cycles.
5. **Wash Buffer:** The 20×Concentrated Wash Buffer should be adjusted to room temperature to make the sediment dissolved fully before use, then dilute it with deionized water at 1:19.

---

**Assay procedure**

Restore all reagents and samples to room temperature (25°C) before use. All the reagents should be mixed thoroughly by gently swirling before pipetting. Avoid foaming. The unused ELISA Microtiter plate should be sealed as soon as possible and stored at 2-8°C.

1. **Number:** number the sample and control in order (multiple well), and keep a record of control wells and sample wells. Set 1 well for blank control, 3 wells for negative control and 2 wells for positive control. **Samples need test in duplicate.** (Blank well is not necessary for dual-wavelength detection)
2. **Add sample:**
  - a) Add 100 µL of **Negative/Positive Control** respectively to 3 negative control wells, 2 positive control wells, keep the blank control well empty.
  - b) Dilute the tested **Serum** with **Sample Diluent** at 1:10 into sample well (add 100 µL of sample diluent and add 10 µL of serum), mix fully.
3. **Incubate:** gently tap the plate to mix thoroughly. Cover the ELISA Microtiter plate with sealer. Incubate for 30 min at 37°C in shading light.
4. **Wash:** remove the plate sealer and aspirate the liquid of each well. Repeat the washing procedure for 5 times with **Wash Buffer** and immerse for 30-60 sec each 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).
5. **HRP conjugate:** add 100 µL of **HRP Conjugated Working Solution** to each well except the blank control well, mix fully.
6. **Incubate:** gently tap the plate to mix thoroughly, cover the ELISA Microtiter plate with sealer. Incubate for 30 minutes at 37°C in shading light.
7. **Wash:** repeat step 4.
8. **Add substrate:** add 50 µL of **Substrate Reagent A** and 50 µL of **Substrate Reagent B** to each well. Gently tap the plate to mix thoroughly. Cover with a new plate sealer. Incubate for 15 min at 37°C in shading light.
9. **Stop reaction:** add 50 µL of **Stop Solution** to each well, gently tap the plate to mix thoroughly.
10. **OD Measurement:** set the Microplate Reader wavelength at 450 nm (it is recommended to set the dual wavelength at 450 nm/630 nm) to detect A value of each well. Blank well is not essential when using dual wavelength 450 nm/630 nm for detection.

**Reference value**

Normally, Blank well (just chromogenic agent and Stop Solution) absorbance  $\leq 0.08$ .

Positive control (PC)  $A_{450} > 0.80$  and Negative control (NC)  $A_{450} < 0.10$ .

**Interpretation of test results**

Calculate the Cut Off: Cut Off (C.O) =  $0.10 + \text{negative control (NC) average A value}$  (when NC average  $A_{450} < 0.05$ , calculate at 0.05; while NC average  $A_{450} \geq 0.05$ , calculate at the actual value).

1. Positive result: Sample absorbance  $\geq$  Cut Off.
2. Negative result: Sample absorbance  $<$  Cut Off.
3. Negative result indicates no JEV-IgG antibody detected in samples, while positive result is just the opposite. JEV-IgG is the important indicator of JEV infection.

**Limitations of test method**

1. This test is only used as the qualitative detection of JEV-IgG antibodies in serum of human.
2. The detection results of this kit are only for reference. For confirmation of the result, please combine the symptoms and other methods of detection, this detection cannot be used as the only criteria for result.