

The Resource for Virology Research

# HA(H11N2)(A/Duck/Yangzhou/906/2002) **Antigen ELISA Development Kit**

Catalog Number: IT-E3Ag-H11N2-Duck/Yangzhou/906/2002

**Description:** HA(H11N2)(A/Duck/Yangzhou/906/2002) antigen ELISA Development Kit contains the key components required for the quantitative analysis of HA(H11N2)(A/Duck/Yangzhou/906/2002) concentrations in cell culture supernatants and serum within the range of 1-1000ng/ml in a sandwich ELISA format. The components supplied in this kit are sufficient to perform the assay in five 96-well ELISA plates.

#### **REAGENTS PROVIDED**

Capture Antibody: 100µl of 1mg/ml anti-HA(H11N2) (A/Duck/Yangzhou/906/2002) monoclonal antibody.

HA(H11N2)(A/Duck/Yangzhou/906/2002) Standard: 50µl of 50µg/ml recombinant HA(H11N2)(A/Duck/ Yangzhou/906/2002).

Detection Antibody: 50µl of biotinylated monoclonal HA(H11N2)(A/Duck/Yangzhou/ antibody against 906/2002).

Streptavidin-HRP Conjugate: 50µl of HRP-conjugated streptavidin.

## **RECOMMENDED MATERIALS & SOLUTIONS\***

ELISA 96-well plates (Corning Prod # 3590 or

equivalent plate)

**Block Buffer:** 5% milk in PBS

Wash Buffer: 0.05% Tween-20 in PBS Diluent: 0.05% Tween-20, 0.5% milk in PBS **Substrate:** TMB Peroxidase Substrate Stop Solution: 2N Sulfuric Acid

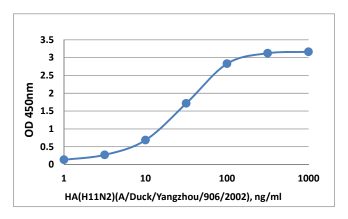
Alternatively, these could be purchased under Cat.# IT-200-002 — ELISA Plate/Buffer/Substrate Kit.

# PLATE PREPARATION

- For each 96-well plate, dilute 20µl of Capture Antibody with 10.5ml of 1xPBS to prepare a coating solution. Immediately add 100µl of the coating solution to each well. Seal the plate and incubate overnight at 4°C.
- 2. Remove the coating solution by aspirating or decanting. Invert the plate and blot it briefly against clean paper towels.
- 3. Add 300µl of Block Buffer to each well. Incubate at 37°C for 2 hours.
- 4. Aspirate to remove Block Buffer and wash the plate 4 times with 300µl of Wash Buffer per well.

### ASSAY PROCEDURE

- Standard/Sample: Dilute standard with Diluent to concentrations (1000ng/ml, 316ng/ml, 100ng/ml, 31.6ng/ml, 10ng/ml, 3.16ng/ml, 1ng/ml, Ong/ml). Immediately, add 100µl of standard and sample to each well in triplicate. Incubate at 37°C for 1 hour.
- 2. **Detection:** Aspirate and wash plate 4 times. Dilute 10µl of Detection Antibody with 10.5ml of Diluent to prepare a detection solution. Add 100µl of the detection solution into each well. Incubate at 37°C for 1 hour.
- 3. Streptavidin Peroxidase: Aspirate and wash plate 4 times. Dilute 10µl of Streptavidin-HRP conjugate with 10.5ml of Diluent. Add 100µl into each well. Incubate at 37°C for 45 minutes.
- **4.** Substrate/Stop: Aspirate and wash plate 4 times. Add 100µl of TMB Peroxidase Substrate into each well. Incubate at 37°C for 30 minutes. Then add 100µl of stop solution to each well.
- **5. Read:** Determine the optical density of each well within 30 minutes, using a microplate reader set to 450nm.
- 6. Analysis: Average the triplicate reading for each standard, control, and sample and subtract the average zero standard optical density. Create a standard curve by reducing the data using computer software capable of generating a four parameter logistic (4-PL) or other curve-fit. The HA(H11N2) (A/Duck/Yangzhou/906/2002) concentration in sample can be determined by regression analysis. If samples have been diluted, the concentration calculated from the standard curve must be multiplied by the dilution factor.



### Reference

John R. Crowther. The ELISA Guidebook (Methods in Molecular Biology), Humana Press, 2000.