# PRODUCT INFORMATION SUMMARY

# Human sTRAIL ELISA Construction Kit

Product Number RHF760CKC
With Developing Reagents:

Capture Antibody 100.0 ug
Biotin-Labeled tracer 25.0 ug
Antigen Standard 1.0 ug or 10.0ug

ELISA Coating Stabilizer 50 mL Streptavidin-HRP 0.5mL TMB Substrate (50 mL x 2) WASH Buffer (20X) 100 mL

RHF760CKP: w/ Developing Reagents & 10 ELISA plates

### DESCRIPTION:

This ELISA CONSTRUCTION Kit provides antigen affinity purified capture and tracer antibodies, and antigen standard sufficient for **approximately** ten microplates.

Working concentrations must be optimized by customer.

Note: Reconstitute components only when ready to run assay.

# CAPTURE ANTIBODY:

Provided as lyophilized, 100 ug, additive-free. Reconstitute in 0.50~mL sterile or distilled water (200.0 ug/mL).

#### TRACER ANTIBODY:

Provided as 25 ug (lyophilized or as 0.5 mL liquid- see vial) of Biotin labeled, antigen-affinity purified antibody, additive-free. Reconstitute lyophilized vial in 500 uL sterile water containing 0.1% BSA.

For biotin tracer provided as 0.5 mL liquid- STORE refrigerated.

**STANDARD:** Provided as 10.0 ug of recombinant Human TRAIL. **Quick-spin** and reconstitute in 100 uL of sterile water. Further dilutions can be made in 0.1% BSA in PBS.

HANDLING/ STORAGE: Reconstitute reagents when ready to build ELISA assay. Antibodies (Capture and Tracer) can be stored for approximately one month at 4 Degrees C. Or store frozen at -20 Degrees C. for up to 6 months. Standard (rec. Human TRAIL) can be stored in liquid state (@ 4 Deg. C.) For up to one week, or store frozen at -20 Deg. C. for up to 2 months. AVOID repeat freeze-thaw.

# MATERIALS RECOMMENDED:

ELISA Microplates: Nunc Maxisorp, Prod. # 4420404

Tween -20

BSA

Streptavidin-HRP: ANTIGENIX Cat no. **S100180** or similar

TMB Substrate (ANTIGENIX cat # ES200)

Dubelco's PBS (10X)

ANTIGENIX ELISA Coating Stabilizer (cat no: EA150)

#### RECOMMENDED SOLUTIONS:

PBS: Dilute to 1XPBS in sterile water WASH BUFFER: 0.05% Tween-20 in PBS.

BLOCK BUFFER: use ANTIGENIX AMERICA coating stabilizer (EA150)

or 1% BSA in PBS

Substrate Solution: TMB Substrate Solution (ANTIGENIX # ES200)

Diluent: 0.05% Tween-20, 0.1% BSA in PBS

# PLATE PREPARATION:

1. **Dilute portion** of capture antibody with 0.05M Carbonate buffer (or PBS) to concentration **1.0 ug/mL**. Immediately **add** 100 uL to each ELISA well. Seal the plate and incubate overnight at room temperature.

- 2. Aspirate wells to remove all liquid and wash 4 times using 300 uL of wash buffer per well. After last wash, add 200 uL ANTIGENIX AMERICA ELISA coating stabilizer (cat # EA150)-recommended! and incubate for 60 minutes at room temperature. (With coating stabilizer, DO NOT let plate dry prior to use of coating stabilizer. This will stabilize and Block in one step! Refer to data sheet EA150 for complete description of use.
- 3. With ANTIGENIX coating stabilizer ( **recommended** ) aspirate plate but **DO NOT WASH**. For extended storage -Dry plate in humidity controlled chamber or similar. ( see data sheet EA150). With standard block reagent, aspirate plate and wash 3X with 300 uL wash buffer.

# PROTOCOL:

STANDARD/SAMPLE: Dilute a portion of the standard (store unused standard in aliquots, high concentration, frozen -20 Deg. C.) from 2.0 ng/mL (1:2, 8 point serial dilution). Immediately add 100 uL of standard or sample to each well in duplicate. Incubate at room temp. for aprrox. 90 minutes.

**DETECTION:** Aspirate and wash plate 4 times. **Dilute** detection (Tracer) antibody in diluent to concentration of 0.20 ug/mL. (approx. 1: 250 from 50.0 ug/mL) **Add** 100 uL per well. Incubate

at room temperature for 1 hour.

Note: detection antibody can be used in approximate range of  $0.15 - 0.30 \, \text{ug/mL}$ , you may need to optimize for subsequent plates.

STREPTAVIDIN-HRP: Aspirate and wash plate 4 times. Dilute Streptavidin-HRP conjugate approx. 1:2000 in diluent (follow recommended dilution of manufacturer). (May need to optimize) Add 100 uL per well, incubate 30 minutes at room temperature.

**SUBSTRATE:** Aspirate and wash plate 4 times. Add 100 uL substrate solution to each well. (follow directions from manufacturer) **Incubate** at room temp. for color development. STOP color development after approx. 10-15 minutes. Read plate at 450 nm within 30 minutes.

NOTE: reliable standard curves are obtained when O.D. readings do not exceed 0.30 units for the zero standard concentration, or 1.5 units for the highest standard concentration.

RESEARCH USE ONLY -NOT For DIAGNOSTIC USE

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