# HIGH SENSITIVITY SOLUBLE IL1R3/IL1RAP (HUMAN) **ELISA KIT**

FOR THE QUANTITATIVE DETERMINATION OF **HUMAN SOLUBLE IL1R3/IL1RAP CONCENTRATIONS IN SERUM, EDTA PLASMA** 



ALWAYS REFER TO LOT SPECIFIC PROTCOL PROVIDED WITH EACH KIT FOR **INSTRUCTIONS. PROTOCOL MUST BE** READ BEFORE USING THIS PRODUCT.

FOR RESEARCH USE ONLY.NOT FOR USE IN **DIAGNOSTIC PROCEDURES.** 

## PRODUCT INFORMATION: THIS KIT IS FOR ONE TIME USE ONLY.

ELISA NAME	HIGH SENSITIVITY		
	IL1R3/IL1RAP (HUMAN)		
	ELISA KIT		
Catalog No.	SK00118-19		
Formulation	96 T		
Lot No.			
Standard	15.6 -1000 pg/mL		
range			
Sensitivity	7 pg/mL		
Sample	100 μL		
Volume			
Dilution	200 ~400 (Optimal dilutions		
Factor	should be determined by		
	each laboratory for each		
	application)		
Sample Type	Serum, EDTA Plasma		
Specificity	Human Soluble IL1R3/IL1RAP		
Calibration	Human Soluble IL1R3/IL1RAP		
	HEK293 derived		
Intra-assay	4 - 6%		
Precision			
Inter-assay	8 - 10%		
Precision			
Storage	2 – 8° C for 1 month. See		
	page 2-3 for detail		
This kit contain	s sufficient materials to run		
approximately 40 samples duplicated			
provided that assay is rup according to			

provided that assay is run according to protocol.

**Order Contact:** 

AVISCERA BIOSCIENCE, INC. 2348 Walsh Ave., Suite C Santa Clara, CA 95051

**USA** 

Tel: (408) 982 0300 (408) 982 0301 Fax:

Email: Sales@AvisceraBioscience.com

Info@AvisceraBioscience.com

www.AvisceraBioscience.com www.AvisceraBioscience.net

### **DESCRIPTION**

This High Sensitivity Soluble IL1R3/IL1RAP (Human) ELISA Kit contains the necessary components required for the quantitative measurement of recombinant human soluble (extracellular domain) IL1R3/IL1RAP derived from HEK293 cells and/or natural human IL1R3/IL1RAP from serum and EDTA plasma in a sandwich ELISA format.

This immunoassay contains recombinant human IL1R3/IL1RAP derived from HEK293 cells and antigen affinity purified antibodies raised against this protein. Results from this immunoassay have shown to accurately quantify recombinant and natural IL1R3 /IL1RAP in samples.

#### **ASSAY OVERVIEW**

This assay employs the quantitative sandwich ELISA format. The plate is pre-coated with an antibody specific for human IL1R3/IL1RAP. The capture antibody can bind to the human IL1R3/IL1RAP in the standard and samples. After washing the plate of any unbound substances, the biotinylated antibody against human IL1R3/IL1RAP is added to the wells. After another washing of the plate, Streptavidin-HRP Conjugate is added. After the last wash to remove any unbound enzyme, a substrate solution (TMB) is added to the wells and color develops in direct proportion to the amount of human IL1R3/IL1RAP bound in the standard solutions or samples. A standard curve can be established and sample values can be read off the standard curve.

## **PROCEDURAL LIMITATIONS**

\_FOR RESEARCH USE ONLY. NOT FOR USE IN DIAGNOSTIC PROCEDURES.

\_This ELISA kit should not be used beyond the expiration date on the kit label.

\_Do not mix reagents with those from other lots or sources

\_It is important that the Dilution Buffer selected for the standard curve be consistent with the samples being assayed.

\_Each laboratory must determine the optimal dilution factors for the samples being assayed.

\_Any modifications in buffers, pipetting technique, washing technique, incubation time or temperature, as well as kit age can cause a change in signal.

\_Not all interfering factors have been tested in the immunoassay, therefore the possibility of interference cannot be excluded.

### **COMPONENTS PROVIDED**

DESCRIPTION	CODE	QUANTITY
<b>IL1R3 Microplate</b> – 96 well microplate coated with	118-19-	1 plate
antibody specific for human IL1R3/IL1RAP.	01	
IL1R3 Standard – refer to lot of lyophilized	118-19-	1 vial
recombinant human IL1R3/IL1RAP (HEK293).	02	
Detection Antibody Concentrate – refer to lot lot of 10-fold concentrate	118-19-	1 vial
of lyophilized biotinylated antibody against human IL1R3/IL1RAP.	03	
Positive Control— one vial of lyophilized recombinant human IL1R3/IL1RAP.	118-19-	1 vial
Streptavidin-HRP	04	
Conjugate – 120 μL/vial of 100-fold concentrated solution of Streptavidin-	SAHRP	1 vial
HRP conjugate.		
<b>Dilution Buffer</b> – 40 mL of buffered solution with preservative.	DB12	2 bottles
Antibody Diluent Solution – 12 mL of buffered solution with preservative.	DB11B	1 bottle
Wash Buffer – 50 mL of 10-fold concentrated buffered surfactant with preservative.	WB01	1 bottle
TMB Substrate Solution  – 11 mL of TMB substrate solution.	TMB01	1 bottle
Stop Solution – 11 mL of 0.5M HCl.	S-STOP	1 bottle
Plate Sealer	EAPS	1 piece
Plastic Pouch	P01	1 piece

## **STORAGE**

**Unopened Kit:** Store at  $2-8^{\circ}$  C for up to 1month. For longer storage for up to 10 months, unopened Standard, Positive Control, Detection Antibody Concentrate, Dilution Buffer and Antibody Diluent Solution should be stored at -20° C. Streptavidin-HRP Conjugate and TMB Substrate Solution should be stored only at  $2^{\circ}$  8°C. Do not use kit past expiration date.

ADDITIONAL MATERIALS REQUIRED

- Microplate reader capable of absorbance measurement at 450 nm.
- Microplate shaker (250 300 rpm).
- Microplate washer or manifold dispenser.
- 100 mL and 500 mL graduated cylinders.
- Multi-channel Pipette, Pipettes and pipette tips.
- Deionized or distilled water.

### **PRECAUTION**

This kit should be handled by those persons who have been trained in and can follow the principles of good laboratory practice. Wear protective clothing such as laboratory overalls, safety glasses and gloves. Care should be taken while handling solutions in this kit to avoid contact with skin or eyes, especially with the stop solution because it contains diluted hydrochloric acid. Wash immediately with water in case of contact on skin or eyes.

## SAMPLE COLLECTION AND STORAGE

**Plasma** - Collect plasma using EDTA as an anticoagulant. Centrifuge for 15 minutes at 1000 x g within 30 minutes of collection. Assay immediately or aliquot and store samples at  $\leq$  -20° C. Avoid repeated freeze-thaw cycles.

**Serum** – Use a serum separator tube (SST). Allow blood to clot for 30 minutes. Centrifuge at  $1000 \times g$  for 15 minutes and collect serum. Assay samples immediately or aliquot and store at  $\leq$  -20° C. Avoid repeated freeze-thaw cycles.

Optional: Use Aprotinin (enzyme inhibitor) (Aviscera Bioscience's Catalog No: 00700-01-25) for ALL sample collection to prevent sample degradation. 0.5 TIU per mL of sample solution.

## SAMPLE PREPARATION

EDTA Plasma or Serum samples may require at least a 200-fold  $^{\sim}$  400-fold dilution. A 20-fold dilution is 10  $\mu L$  sample + 190  $\mu L$  1x Dilution Buffer. A suggested 200-fold dilution is 30  $\mu l$  of 20-fold diluted sample solution + 270  $\mu l$  Dilution Buffer. A suggested 400-fold dilution is 15  $\mu l$  of 20-fold diluted sample + 285  $\mu l$  Dilution Buffer.

Optimal dilutions should be determined by each laboratory for each application with a sample pretest.

Use polypropylene test tubes.

### REAGENT PREPARATION

Bring all reagents to room temperature before use.

Wash Buffer – If crystals have formed in the concentrate, warm bottle in a water bath until the crystals have completely dissolved. Dilute 50 mL of Wash Buffer Concentrate into 450 mL distilled or deionized water to make 500 mL of 1x Wash Buffer.

Dilution Buffer (DB12) - Dilution Buffer (DB12) is highly viscous, warm in 30 - 37° C water bath until liquid flows more freely.

IL1R3/IL1RAP Standard – Reconstitute the IL1R3/IL1RAP Standard vial with refer to lot of Dilution Buffer (DB12). Allow the stock standard to sit for at least 15 minutes with gentle agitation until completely dissolved prior to making standard dilutions. Pipette 250  $\mu$ L of Dilution Buffer (DB12) into tubes #2 to #7. Use the stock solution to produce a dilution series (below). Mix each tube thoroughly before the next transfer. The 1000 pg/mL standard serves as the high standard. The Dilution Buffer (DB12) serves as the zero standard (0 pg/mL).

TUBE	STANDARD	DILUTION BUFFER	CONCENTRATION
Stock	powder	1mL	XXXXX
#1	Refer to lot	Refer to lot	1000 pg/mL
# 2	250 μL of 1	250 μL	500 pg/mL
#3	250 μL of 2	250 μL	250 pg/mL
# 4	250 μL of 3	250 μL	125 pg/mL
# 5	250 μL of 4	250 μL	62.5 pg/mL
# 6	250 μL of 5	250 μL	31.25 pg/mL
# 7	250 μL of 6	250 μL	15.6 pg/mL

**Positive Control** - Reconstitute the Positive Control with refer to lot of **Dilution Buffer (DB12)** to prepare working solution.

Detection Antibody - Reconstitute the Detection Antibody Concentrate with refer to lot of Antibody Diluent Solution (DB11B) to produce a 10-fold concentrated stock solution. Allow the concentrated solution to sit for at least 5 minutes until completely dissolved. Pipette refer to lot of Antibody Diluent Solution (DB11B) into a 15 mL centrifuge tube and transfer refer to lot of 10-fold concentrated stock solution to prepare working solution.

Streptavidin HRP Conjugate - Pipette 10.89 mL of Dilution Buffer (DB12) into a 15 mL centrifuge tube and transfer 110  $\mu$ L of 100-fold concentrated stock solution to prepare working solution (protect from light). DO NOT FREEZE.

## **ELISA PROTOCOL**

Bring all reagents and samples to room temperature before the start of the assay. Blank, standard dilutions, positive control and samples should be assayed in duplicate. ELISA Protocol may need further optimization.

- 1. Prepare all reagents, standard dilutions, positive control and samples as directed previously.
- 2. Add 100  $\mu$ L per well of **Dilution Buffer (DB12)** to Blank wells (BL).
- Add 100 μL per well of Standard dilutions 7 to 1 and or optional standard dilution (O), samples, or positive control (P). See page 5 for more information. Cover with plate sealer and incubate for 2 hours on microplate shaker at room temperature.
- 4. Aspirate and wash each well with 300 μL of 1x Wash Buffer four times. After the last wash, aspirate any remaining 1x Wash Buffer, invert the plate and blot against clean paper towel(s).
- 5. Add 100  $\mu$ L per well of **Detection Antibody** working solution. Cover with plate sealer and incubate for 2 hours on microplate shaker at room temperature.
- 6. Repeat the aspiration and wash as in step 4.
- 7. Add 100 µL per well of Streptavidin HRP Conjugate working solution. Cover with plate sealer and incubate for 60 minutes on microplate shaker at room temperature. Protect from light.
- 8. Repeat the aspiration and wash as in step 4.
- 9. Add 100  $\mu$ L per well of **Substrate Solution**. Incubate for refer to lot on microplate shaker at room temperature. **Protect from light.**
- 10. Add 100  $\mu$ L per well of **Stop Solution**. The color in the wells should change from blue to yellow. If the color change does not appear uniform, gently tap the plate to ensure thorough mixing.
- 11. Read plate using a microplate reader set to 450 nm within 3 minutes.

## **CALCULATION OF RESULTS**

Create a standard curve by plotting the log of the known concentrations of the standard dilutions (x-axis) versus the log of its corresponding O.D. (y-axis) and draw the best fit line through the points. It is

recommended to use computer software capable of generating a log-log or 4-parameter curve fit to more accurately quantify the standard dilutions.

If samples have been diluted, the concentration read from the standard curve must be multiplied by the dilution factor (200 or 400).

#### SPECIFICITY

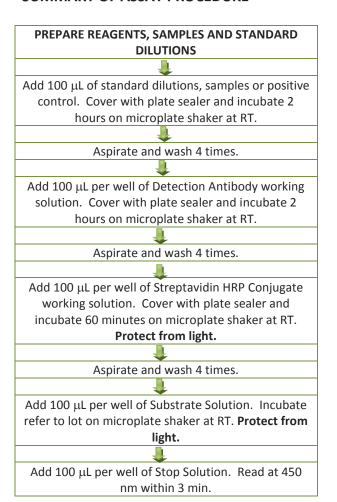
Protein	Cross-reactivity (%)	
Human Soluble IL1R3	100	
/IL1RAP		
Human Soluble IL1R4	0	
/ST2		
Human Soluble IL1R1	0	
Human IL1R2	0	
Human IL-33	0	

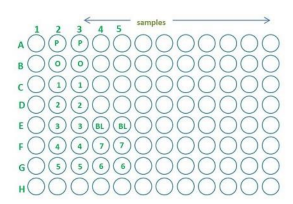
#### TYPICAL STANDARD CURVE

This standard curve is for demonstration only. A new standard curve should be made for each set of samples assayed.

STANDARD	AVERAGE OD450 (CORRECTED)		
(PG/ML)			
Blank	0 (refer to lot)		
15.6	0.042		
31.25	0.088		
62.5	0.198		
125	0.382		
250	0.773		
500	1.256		
1000	2.251		

### **SUMMARY OF ASSAY PROCEDURE**





## Sample Test

The research samples were diluted by Dilution Buffer (DB12). Its linearity and recovery was assayed by High Sensitivity Soluble IL1R3/IL1RAP (Human) ELISA Kit, code SK00118-19.

Sample Type	Dilution Factor	Assayed (pg/mL)	Final (ng/mL)	Recovery (%)
Human EDTA Plasma	100 X	804.965	80.496	100
Human EDTA Plasma	200 X	416.836	83.367	104
Human EDTA Plasma	400 X	207.059	82.823	102
Human Serum	100 X	837.946	83.794	100
Human Serum	200 X	415.391	83.078	99.2
Human Serum	400 X	216.128	86.451	103