

Code No. 27201

# **Human GIP, Active form Assay Kit - IBL**

#### INTRODUCTION

Incretins are a group of gastrointestinal hormones that cause an increase in the amount of insulin released from the beta cells of the islets of Langerhans after eating and they also inhibit glucagon release from the alpha cells of the Islets of Langerhans. GIP, typical incretin like GLP-1, was isolated and sequenced from intestinal mucosa as "gastric inhibitory peptide" in 1970, and then it was renamed as "glucose-dependent insulinotropic peptide". It has been reported that GIP receptor is expressed in cells such as beta cell of pancreas, adipocyte or osteoblastic cell, and it plays essential roles in reserving of ingested nutrients within the body in each cell, and the control of GIP signal can lead to improvement of metabolic syndrome (ref. 1 - 3).

It is rapidly inactivated to GIP (3-42) from active form of GIP (1-42) by DPP-IV in blood.

This ELISA kit can measure only active form of Human GIP (1-42).

#### **PRINCIPLE**

This kit is a solid phase sandwich ELISA using 2 kinds of highly specific antibodies. Tetra Methyl Benzidine (TMB) is used as a coloring agent (Chromogen). The strength of coloring is proportional to the quantities of active form human GIP.

#### **MEASUREMENT RANGE**

0.31 - 20.07 pmol/L (1.6 - 100 pg/mL)

#### **INTENDED USE**

For research use only, not for use in diagnostic procedures.

■ This IBL's assay kit is capable for the quantitative determination active form human GIP in EDTA-plasma.

DPP-IV inhibitor has to be added when collecting samples, or use purpose-made blood collection tubes in order to preserve GIP. (eg. BD<sup>TM</sup> Blood Collection System for Preservation of Plasma GLP-1, GIP, Glucagon and Ghrelin, BD Diagnostics)

### KIT COMPONENT

1	Precoated plate : Anti-GIP (C) Rabbit IgG Affinity Purify	96Well x 1
2	Labeled antibody Conc.:	
	(30X) HRP conjugated Anti- GIP (N) (6A1A) Mouse IgG Fab' Affinity Purify	0.4mL x 1
3	Standard : Human GIP (1-42)	0.5mL x 2
4	EIA buffer*	30mL x 1
5	Solution for Labeled antibody*	12mL x 1
6	Chromogen : TMB solution	15mL x 1
7	Stop solution*	12mL x 1
8	Wash buffer Conc.*	50mL x 1

## **OPERATION MANUAL**

# 1. Materials needed but not supplied

- Plate reader (450nm)Graduated cylinder and beaker
- Deionized water
- · Refrigerator (as 4°C)
- · Graph paper (log/log)

Micropipette and tip

- Paper towel
- Tube for dilution of Standard
- Incubator (37°C ± 1°C)
- Washing bottle for precoated plate
- Disposable test tube for "2, Labeled antibody Conc." and "6, Chromogen"

# 2. Preparation

- Preparation of wash buffer
  - "8, Wash buffer Conc." is a concentrated (40X) buffer. Adjust the temperature of "8, Washing buffer Conc." to room temperature and then, mix it gently and completely before use. Dilute 50 mL of "8, Wash buffer Conc." with 1,950 mL of deionized water and mix it. This is the wash buffer for use. This prepared wash buffer shall be stored in refrigerator and used within 2 weeks after dilution.
- 2) Preparation of Labeled antibody
  - "2, Labeled antibody Conc." is a concentrated (30X). Dilute "2, Labeled antibody Conc." with "5, Solution for Labeled antibody" in 30 times according to required quantity into a disposable test tube. Use this resulting solution as Labeled antibody.

# Example)

In case you use one strip (8 well), the required quantity of Labeled antibody is 800  $\mu$ L. (Dilute 30  $\mu$ L of "2, Labeled antibody Conc." with 870  $\mu$ L of "5, Solution for Labeled antibody" and mix it. And use the resulting solution by 100  $\mu$ L in each well.)

This operation should be done just before applying labeled antibody

The remaining "2, Labeled antibody Conc." should be stored at 4°C in firmly sealed vial.

3) Preparation of Standard

Put just <u>0.5 mL</u> of deionized water into the vial of "3, Standard" and mix it gently and completely. This solution is 40.13 pmol/L Human Active GIP standard.

) Dilution of Standard

Prepare 8 tubes for dilution of "3, Standard". Put 230  $\mu L$  each of "4, EIA buffer" into the tube.

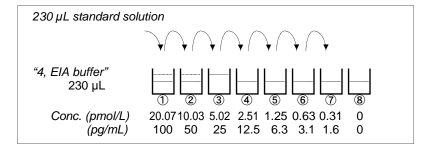
Specify the following concentration of each tube."

i ube- i	20.07 pmo/L	
Tube-2	10.03 pmol/L	
Tube-3	5.02 pmol/L	
Tube-4	2.51 pmol/L	
Tube-5	1.25 pmol/L	
Tube-6	0.63 pmol/L	
Tube-7	0.31 pmol/L	
Tube-8	0 pmol/L	(Test Sample Blank)

Put 230  $\mu L$  of Standard solution into tube-1 and mix it gently. Then, put 230  $\mu L$  of

tube-1 mixture into tube-2. Dilute two times standard solution in series to set up 7 points of diluted standard between 20.07 pmol/L and 0.31 pmol/L. Tube-8 is the test sample blank as 0 pmol/L.

See following picture.



#### 5) Dilution of test sample

It is recommended to dilute test samples more than 4-fold with "4, EIA buffer" in advance.

Most of the fasting blood levels of GIP would be less than measurement range.

#### 3. Measurement procedure

All reagents shall be brought to room temperature approximately 30 minutes before use. Then mix it gently and completely before use. Make sure of no change in quality of the reagents. Standard curve shall be prepared simultaneously with the measurement of test samples.

	Test Sample	Standard	Test Sample Blank	Reagent Blank
Reagents	Test sample 100 µL	Diluted standard (Tube 1-7) 100 µL	EIA buffer (Tube-8) 100 μL	EIA buffer 100 μL
	Incubation for 6	60 minutes at 37°	°C with plate lid	
	4 times (was	sh buffer more th	an 350 μL)*	
Labeled Antibody	100 μL	100 μL	100 μL	-
Incubation for 60 minutes at 4°C with plate lid				
	5 times (wash buffer more than 350 μL)*			
Chromogen	100 μL	100 μL	100 μL	100 μL
Incub	Incubation for 30 minutes at room temperature (shielded)			lded)
Stop solution	100 μL	100 μL	100 μL	100 μL
Read the plate at 450nm against a Reagent Blank within 30 minutes after addition of Stop solution.				

- 1) Determine wells for reagent blank. Put 100  $\mu$ L each of "4, EIA buffer" into the wells.
- 2) Determine wells for test sample blank, test sample and diluted standard. Then, put 100  $\mu$ L each of test sample blank (tube-8), test sample and dilutions of standard (tube-1-7) into the appropriate wells.
- 3) Incubate the precoated plate for 60 minutes at 37°C after covering it with plate
- 4) Wash the plate with the prepared wash buffer and remove all liquid.\*
- 5) Pipette 100 µL of labeled antibody solution into the wells of test samples, diluted standard and test sample blank.
- 6) Incubate the precoated plate for 60 minutes at 4°C after covering it with plate lid
- 7) Wash the plate with the prepared wash buffer and remove all liquid.\*
- 8) Take the required quantity of "6, Chromogen" into a disposable test tube. Then, pipette 100  $\mu$ L from the test tube into every well. Please do not return the rest of used chromogen in the test tube into "6, Chromogen" bottle in order to avoid contamination.
- 9) Incubate the precoated plate for 30 minutes at room temperature in the dark.
  The solution of Chromogen will turn blue.
- 10) Add 100 µL of "7, Stop solution" to all wells. Mix the solution by tapping the side of precoated plate. The solution will turn yellow by addition of "7, Stop solution".
- 11) Remove any dirt or drop of water on the bottom of the precoated plate and confirm there is no bubble on the surface of the solution. Then, run the plate reader and conduct measurement at 450 nm against a reagent blank. The measurement shall be done within 30 minutes after addition of "7, Stop solution".

# **SPECIAL ATTENTION**

- Test samples should be measured soon after collection. For the storage of test samples, store them frozen and do not repeat freeze/thaw cycles. Thaw the test samples at a low temperature and mix them completely before measurement.
- 2) Test samples should be diluted with "4, EIA buffer", as the need arises.
- 3) Duplicate measurement of test samples and standard is recommended.
- 4) Use test samples in neutral pH range. The contaminations of organic solvent may affect the measurement.
- Use only wash buffer in this kit for washing the precoated plate.
   Insufficient washing may lead to the failure in measurement.
- 6) Remove the wash buffer completely by tapping the precoated plate on paper towel. Do not wipe wells with paper towel.
- 7) "6, Chromogen" should be stored in the dark due to its sensitivity against light. Avoid contact of Chromogen with metals.
- 8) Measurement should be done within 30 minutes after addition of "7, Stop solution".



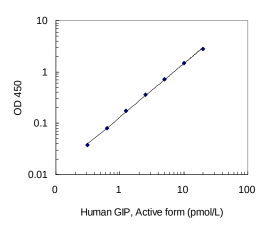
#### **CALCULATION OF TEST RESULT**

Subtract the absorbance of test sample blank from all data, including standards and unknown samples before plotting. Plot the subtracted absorbance of the standards against the standard concentration on log-log graph paper. Draw the best smooth curve through these points to construct the standard curve. Read the concentration for unknown samples from the standard curve.

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#### Example of standard curve

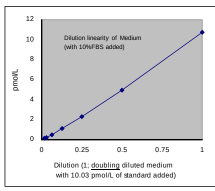
Conc. (pmol/L)	Absorbance (450nm)
20.07	2.832
10.03	1.520
5.02	0.765
2.51	0.397
1.25	0.209
0.63	0.116
0.31	0.073
0 (Test Sample Blank)	0.035

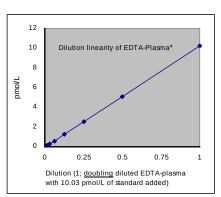


The typical standard curve is shown above. This curve can not be used to derive test results. Please run a standard curve for each assay.

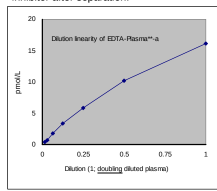
#### PERFORMANCE CHARACTERISTICS

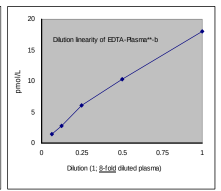
### 1. Dilution linearity





\*The plasma sample was collected by usual method with EDTA and added with DPP-IV inhibitor after separation.





\*\*The plasma sample was separated from after meal blood collected with EDTA and DPP-IV inhibitor added.

# 2. Added Recovery Assay

u	ded Recovery Assay				
	Specimen	Additive Amount (pmol/L)	Theoretical Value (pmol/L)	Measured Value (pmol/L)	%
	*	10.03	10.03	9.97	99.3
	*Human Plasma (EDTA) (x2)	5.02	5.02	5.02	, -
	(LDTA) (XZ)	2.51	2.51	2.42	
		10.03	10.03	10.39	103.5
	10% FBS added RPMI-1640 (x2)	5.02	5.02	5.15	102.6
	( )	2.51	2.51	2.38	95.0

\*The plasma sample was collected by usual method with EDTA and added with DPP-IV inhibitor after separation.

# 3. Intra - Assay

 a rioday				
Mean Value (pmol/L)	SD (pmol/L)	CV (%)	n	
54.7	2.75	5.0	24	
13.2	0.52	4.2	24	
4.9	0.25	5.1	24	

# 4. Inter - Assay

E	er - Assay				
	Mean Value (pmol/L)	SD (pmol/L)	CV (%)	n	
	11.50	0.63	5.5	6	
	2.56	0.15	5.9	6	
	0.98	0.07	7.1	6	

#### 5. Specificity

Substance	Cross-Reactivity
Human GIP (1-42)	100 %
Human GIP (3-42)	< 0.1%
Human Glucagon	< 0.1%
Human GLP-1 (7-36) amide	< 0.1%
Human GLP-2	< 0.1%

#### 6. Sensitivity

#### 0.24 pmol/L

The sensitivity for this kit was determined using the guidelines under the National Committee for Clinical Laboratory Standards (NCCLS) Evaluation Protocols. (National Committee for Clinical Laboratory Standards Evaluation Protocols, SC1, (1989) Villanova, PA: NCCLS.)

#### PRECAUTION FOR INTENDED USE AND/OR HANDLING

- 1. All reagents should be stored at 2 8°C. All reagents shall be brought to room temperature approximately 30 minutes before use.
- 2. "3, Standard" is lyophilized products. Be careful to open this vial.
- 3. "7, Stop solution" is a strong acid substance. Therefore, be careful not to have your skin and clothes contact "7, Stop solution" and pay attention to the disposal of "7, Stop solution".
- 4. Dispose used materials after rinsing them with large quantity of water.
- 5. Precipitation may occur in "2, Labeled antibody Conc.", "4, EIA buffer" or "8, Wash buffer Conc.", however, there is no problem in the performance.
- 6. Wash hands after handling reagents.
- 7. Do not mix the reagents with the reagents from a different lot or kit.
- 8. Do not use expired reagents.
- 9. This kit is for research purpose only. Do not use for clinical diagnosis.

### STORAGE AND THE TERM OF VALIDITY

Storage Condition : 2 - 8°C
The expiry date is specified on outer box.

## REFERENCE

- Miyawaki K, Yamada Y, Yano H, Niwa H, Ban N, Ihara Y, Kubota A, Fujimoto S, Kajikawa M, Kuroe A, Tsuda K, Hashimoto H, Yamashita T, Jomori T, Tashiro F, Miyazaki J, Seino Y. Glucose intolerance caused by a defect in the entero-insular axis: a study in gastric inhibitory polypeptide receptor knockout mice. Proc Natl Acad Sci U S A. 1999 Dec 21:96(26):14843-7.
- Miyawaki K, Yamada Y, Ban N, Ihara Y, Tsukiyama K, Zhou H, Fujimoto S, Oku A, Tsuda K, Toyokuni S, Hiai H, Mizunoya W, Fushiki T, Holst JJ, Makino M, Tashita A, Kobara Y, Tsubamoto Y, Jinnouchi T, Jomori T, Seino Y. Inhibition of gastric inhibitory polypeptide signaling prevents obesity.Nat Med. 2002 Jul;8(7):738-42.
- 3. Tsukiyama K, Yamada Y, Yamada C, Harada N, Kawasaki Y, Ogura M, Bessho K, Li M, Amizuka N, Sato M, Udagawa N, Takahashi N, Tanaka K, Oiso Y, Seino Y. Gastric inhibitory polypeptide as an endogenous factor promoting new bone formation after food ingestion. Mol Endocrinol. 2006 Jul; 20(7):1644-51
- 4. Hansotia T, Maida A, Flock G, Yamada Y, Tsukiyama K, Seino Y, Drucker DJ. Extrapancreatic incretin receptors modulate glucose homeostasis, body weight, and energy expenditure. J Clin Invest. 2007 Jan;117(1):143-52.

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Made in Japan.

# IBL Incretin-related Products:

IBL Incredin-related Products.		
製品番号	製品番号製品名	
27201	Human GIP, Active form Assay Kit - IBL	96 Well
27202	Rat GIP, Active form Assay Kit - IBL	96 Well
27203	Human GIP, Total Assay Kit - IBL	96 Well
27204	Mouse GIP, Total Assay Kit - IBL	96 Well
27205	Rat GIP, Total Assay Kit - IBL	96 Well
27764	Mouse GIP, Active form Assay Kit - IBL	96 Well
27784	GLP-1, Active form Assay Kit - IBL	96 Well
27788	GLP-1 (9-36/37) Assay Kit - IBL	96 Well



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