

UMA CO., LTD.

2-19-6 Yokosuka

Matsudo, Chiba, Japan



MEASURE GGT

Reagent for determination of γ -GTP

IFCC Method

↓ 2 - 8°C

IVD *In vitro* Diagnostics

QUALITY MANAGEMENT SYSTEM (BY TUV)

⊗ DO NOT freeze

⌚ 18 months/block from light

ISO 13485:2016

1. PURPOSE OF USE

Providing a quantitative *in vitro* assay for the γ -glutamyl-transpeptidase (γ -GT) concentration in serum or plasma.

2. GENERAL INSTRUCTION

- For *in vitro* diagnostics use only.
- Diagnosis should be made in a comprehensive manner, in accordance with other related test results and clinical symptoms by the doctor in attendance.
- For guaranteed results, usage of this product must comply with the instruction in this manual.
- If you use automatic analyzers, follow their instructions carefully.

SUMMARY

Gamma-glutamyltransferase (GGT) is an enzyme present in the cell surface membrane of many tissues. Only the isoform of GGT present in the liver is detected in serum. It catalyzes the transfer of the gamma-glutamyl group of glutathione to peptides, amino acids, or water to form glutamate. By regulating glutathione levels, this reaction is possibly involved in protection against oxidative stress (e.g., induced by the metabolism of ethanol). A daily consumption of between 80 and 200 g of ethanol, for a period of several weeks, is required for an increase in GGT activity (measured in serum). The half-life of GGT is between 14 and 26 days. Normal GGT values are reported within 2–5 weeks after cessation of alcohol consumption (Rose, 2008). Elevated GGT can be caused by excessive alcohol consumption, but is also seen in cases of liver damage due to cholestasis, and pancreas or kidney damage, and obesity, etc. Upper reference limits of 36 and 61 U/L for females and males, respectively, have been published.

3. MATERIALS REQUIRED BUT NOT INCLUDED

- Saline 0.9 % and high grade purified water
- Micropipet and other basic laboratory equipment.
- MEASURE Multi Calibrator and MEASURE Human Lyo L-1 and MEASURE Human Lyo L-2

4. REAGENT COMPOSITION & PREPARATION

- Reagent R-1: Glycylglycine (Gly-Gly)

Reagent R-1 is ready for use

- Reagent R-2: L- γ -glutamyl -3-carboxy -4-nitroanilide mono-ammonium

Reagent R-2 is ready for use

- Once open, Reagent stored on board the instrument is stable for 30 days with Hitachi 7180 Analyzers.

- Applicable to various automated analyzers.

- Calibrator MEASURE Multi Calibrator (separately sold): Put 5 mL of purified water to the vials of Calibrator (MEASURE Multi Calibrator), leave at room temperature for 45 minutes and sometimes gently invert the vial before use. After reconstituting, Calibrator can be used without dilution.

- Controls MEASURE Human Lyo L-1 and MEASURE Human Lyo L-2 (separately sold): Put 5 mL of purified water to the vials of controls (Lyo L-1 and Lyo L-2); leave at room temperature for 45 minutes and sometimes gently invert the vial before use. After reconstituting, controls can be used without dilution.

5. SAMPLE PREPARATION & STORAGE

- Serum: Wait until the sample is completely coagulated. Take the supernatant to use as specimen.

- Plasma: Treat blood sample by anticoagulant (Li-heparin and K2-EDTA); leave it to stand for 3 hours or centrifuge at 2000 rpm for 2 minutes; take the plasma layer (supernatant) and use as a specimen.

- Analyze samples soon after collection.

- Stability

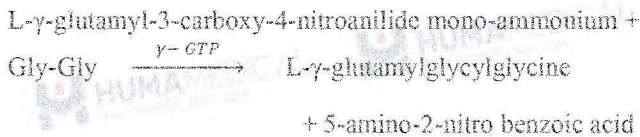
- 8 hours at 15 - 25°C
- 3 days at 2 - 8°C
- 30 days at < -20°C

- See interferences section for details about possible sample interferences.

6. MEASUREMENT PRINCIPLE

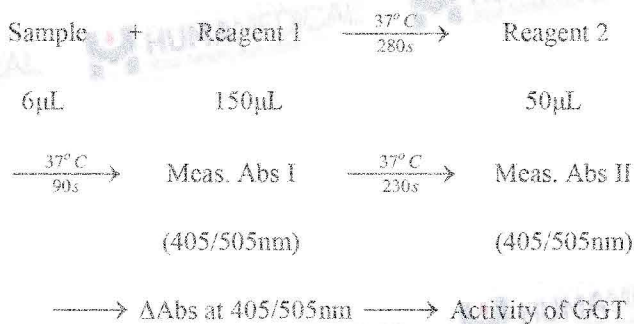
In the presence of glycylglycine, γ -GTP in the sample acts on L- γ -glutamyl-3-carboxy-4-nitroanilide mono-ammonium and generates L- γ -glutamyl-glycylglycine and 5-amino-2-nitro benzoic acid.

Activity of γ -GTP can be determined by measuring the increase of the absorbance of this 5-amino-2-nitro benzoic acid.



7. ASSAY PROCEDURE

This product is compatible with various types of clinical analyzer. An example of the assay procedure is indicated below.



Perform the assay according to the instructions for operating the automated analyzer Hitachi models. Refer to the 13. INFORMATION FOR AUTOANALYZERS for the details of the assay method. Contact HUMA MEDICAL CO., LTD. for information about the parameters for other automated analyzers.

8. CALCULATION & UNIT CONVERSION

Calculation

- Calculate Δ Abs of specimen & standards vs blank
- Plot a calibration curve $GGT = f(\Delta Abs)$
- Calculate GGT in specimen using the curve

(doing same procedure for

Controls)

Unit conversion

$$\text{U/L} \times 0.0167 = \mu\text{kat/L}$$

9. PERFORMANCE & CORRELATION TEST

a. Measuring range

- The assay is linear within an GGT enzyme activity in serum/plasma of 1 - 1500 U/L.
- If the concentration of sample exceeds assay range, dilute the sample with saline and repeat the measurement.

b. Detection Limit

Limit of Blank (LoB)	=	0 U/L
Limit of Detection (LoD)	=	1 U/L
Limit of Quantitation (LoQ)	=	1 U/L

The LoB, LoD and LoQ were determined in accordance with CLSI EP17-A2 requirements.

The LoB is the highest apparent analyte concentration expected to be found when replicates of a blank sample containing no analyte are tested. The LoB corresponds to the concentration below which analyte-free samples are found with a probability of 95%.

The LoD is determined based on the LoB and standard deviation of low concentration samples. The LoD corresponds to the lowest analyte concentration which can be detected (value above the LoB with a probability of 95%).

The LoQ is the lowest analyte concentration that can be reproducibly measured with a total error of 20%. It has been determined using low concentration samples.

c. Performance

- Sensitivity: when using purified water; change in absorbance is 0.001 - 0.050, when using γ -GTP 200U/L solution, change in absorbance is within 0.02 - 0.20 Abs/min.
- Accuracy: When measuring a control sample, the result is within $\pm 5\%$ of assigned value.

d. Precision (on Biolis 30i / SK300)

Representative performance data on the analyzers are given below.

Results obtained in individual laboratories may differ.

Precision was determined using controls followed the CLSI Approved Guideline EP5-A2 with repeatability, reproducibility and total precision (1 aliquot per run, 2 run per day, 20 days). The following results were obtained.

Criterion: CV of Repeatability (aka. Within-run precision) is less than 3% and Total Precision is less than 5%.

Repeatability	Mean U/L	SD U/L	CV %
Control Lyo L-1	36.5	0.36	0.98
Control Lyo L-2	142.7	0.85	0.59

Measure GGT

Reproducibility	Mean	SD	CV
	U/L	U/L	%
Control Lyo L-1	36.5	1.23	3.36
Control Lyo L-2	142.7	3.74	2.62

Total precision	Mean	SD	CV
	U/L	U/L	%
Control Lyo L-1	36.5	1.25	3.43
Control Lyo L-2	142.7	3.78	2.65

e. Correlation Test

Regression equation $y = 0.9961x + 0.3818$ ($n = 50$)

Correlation coefficient $r = 1.000$

Reference Material for Calibration

ReCCS JCCLS CRM-001

10. EXPECTED VALUES

- Male: 13 - 64 U/L

- Female: 9 - 32 U/L

Reference range should be established at each facility and judgement should be based on measurement results in a comprehensive manner together with clinical symptoms and other measurement results.

11. INTERFERENCES

- Icterus: No significant interference of conjugated/free bilirubin concentration up to 20 mg/dL

- Hemolysis: No significant interference of hemoglobin concentration up to 500 mg/dL.

- Lipemia (Intralipid): No significant interference triglycerides concentration up to 3000 FTU

- Ascorbic Acid: No significant interference of ascorbic acid concentration up to 50 mg/dL.

- For diagnostic purposes, the results should always be assessed in conjunction with the patient's medical history, clinical examination and other findings. Please use another methods if the result is affected by any factors

12. HANDLING, USAGE & DISPOSAL

Handling

1. Specimen can be potentially positive for infectious agents including hepatitis B virus and HIV. Wear glove and goggle when needed.

2. In case reagents got into skin, eye or mouth by mistake, wash it immediately with plenty of water and consult the doctor if needed.

3. If reagents are spilled, dilute with water and wipe it out. If specimen is spilled, spray 80% of alcohol over the specimen and wipe it out.

Usage

1. Store reagents under specified condition. Do not use after expiration date.

2. Do not use the container and auxiliaries included in this kit for other purposes.

3. Do not mix reagents of different lot for use.

4. Do not add to the reagent being used even if it is the same lot number.

Disposal

1. All specimens, as well as all instruments (e.g. test tubes) that come in contact with the specimens, must be treated by the following methods, or they must be treated according to the manual for infectious medical waste provided in each facility.

* Sterilize with an autoclave, subjecting them to high pressure saturated steam at 121 °C for more than 20 minutes. Do not process waste containing sodium hypochlorite solution with an autoclave.

* Immerse at least one hour in sodium hypochlorite solution (active chloride concentration of over 1000 ppm).

2. This reagent contains sodium azide. Sodium azide can react with lead pipe and/or steel pipe and can generate explosive metal azide. Make sure to use plenty of water at disposal. Concentration of sodium azide in R-2 is 0.05%.

13. INFORMATION FOR AUTOANALYZERS

❖ For Hitachi Model

Calculation Method		Rate
Temperature		37°C
Volume (µL)	Specimen	6
	R1	150
Wavelength (nm)	R2	50
	Main	405
Measurement (cycle)	Sub	505
	Point 1	10
	Point 2	22
Calibration type	Point 3	34
		Linear
Unit		U/L

14. OTHER INSTRUCTIONS AND CAUTION

- Results may differ depending on the sample/reagent ratio. Adjust parameters for different analyzer.
- Perform the QC procedure on the day of determination.

15. PACKING AND KIT CONFIGURATION

Code	Package	Test/Kit*	Test/Kit**
11G012A	1x60mL; 1x20mL	310	540
11G012A2	2x60mL; 2x20mL	620	1080
11G012A3	3x60mL; 3x20mL	930	1620
11G012A4	4x60mL; 4x20mL	1240	2160
11G002A	5x60mL; 5x20mL	1550	2700
11G012A6	6x60mL; 6x20mL	1860	3240
11G012	1x90mL; 1x30mL	470	810
11G012-2	2x90mL; 2x30mL	940	1620
11G002	3x90mL; 3x30mL	1410	2430
11G012-4	4x90mL; 4x30mL	1880	3240
11G012-5	5x90mL; 5x30mL	2350	4050

* For middle-scale automatic analyzers such as: SK300; BS series; BA200; BA400. Chemwell Series; Dirui Series; Biolyzer series, HumanStar 300, Erba Series; Bioelab Series, BX 3010; Pictus P500;...

** For large-scale automatic analyzers such as: CA800; CA400; Randox Imola; Randox Modena+; BM 6010; Biolis50i; SK500; AU Series; Pictus P700; C series; Ci series; HumanStar 600; Kenolab series ...

The above-mentioned test's number are calculated base on technical specifications of each analyzer. The real number of test per kit may higher than the calculation's number.

The above-mentioned test's number cover the loss of the dead volume of reagent bottles but not cover the loss of Calibrator and Control.

Please feel free to contact authorized distributor for further confirmation.

16. REFERENCES

1. Aurelie De Vos, ... Christophe Stove, in Neuroscience of Alcohol, 2019
1. CLSI/NCCLS Evaluation of Precision Performance of Clinical Chemistry Devices, EP05-A2, 2004

2. CLSI EP17 - Evaluation of Detection Capability for Clinical Laboratory Measurement Procedures, 2nd Edition, 2017
3. In house data, UMA Diagnostics

17. MANUFACTURER

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