

Lipase Estimation Kit

High-Q Lipase



(Enzymatic/Colorimetric Method)

Intended Use:

Kit for the quantitative determination of Lipase in human Serum and Plasma.

Summary:

Lipases are enzymes which hydrolyze glycerol esters of long fatty acids. The enzyme and its cofactor colipase are produced in the pancreas, lipase being also secreted in small amounts by the salivary glands as well as by gastric, pulmonary and intestinal mucosa. Bile acids and colipase form micellar complexes with the lipids and bind lipase on the substrate / water interface. Determination of lipase is used for investigation of pancreatic disorders. In acute pancreatitis the lipase concentrations rise to 2-50 fold the upper reference limit within 4-8 hours after the beginning of abdominal pain peaking at 24 hours and decrease within 8 to 14 days. Elevated lipase values can also be observed in chronic pancreatitis and obstruction of the pancreatic duct.

Method:

A synthetically produced lipase substrate (1.2-o-dilaurylrac-glycero-3-glutaric acid-(6-methylresorufin) ester) is added to a microemulsion which is specifically split by lipase in the presence of colipase and bile acids. The combination of lipase and bile acids make this specific and reliable for pancreatic lipase without any reaction due to lipolytic enzymes or esterases. The reagent composition has been thoroughly optimized so there are no serum matrix effects. The generated methylresorufin-ester is spontaneously degraded to methylresorufin. The absorbance by this red dye is directly proportional to the lipase activity in the sample.

Principle: Lipase catalyses the reaction

1,2-o-Dilauryl-rac-glycero-3-glutaric acid(6-methylresorufin) ester

Lipase / Colipase

--> 1,2-o-Dilauryl-rac-glycerin + Glutaric acid-(6methylresorufin)-ester

Spontaneous degradation Glutaric acid-(6-methylresorufin)-ester <-Glutaric acid + Methylresorufin

The increase in absorbance is determined photometrically.

Reagents: **Components and Concentrations**

Reagent 1:

Goods Buffer (pH 8.0) Taurodesoxycholate Desoxycholate Calcium chloride Colipase Detergent Preservative

Reagent 2:

Tartrate Buffer (pH 4.0) Taurodesoxycholate Lipase Substrate Coemulgator Stabilizer Preservative

50 mmol/L 4.3 mmol/L 8.0 mmol/L 15 mmol/L 2.2 mg/L

7.5 mmol/L 17.2 mmol/L 0.65 mmol/L

Storage Instructions and Reagent Stability:

The reagents are stable up to the end of the indicated expiry date if stored at

2-8 °C and when the contamination is avoided. Do not freeze the reagents!

Reagent Preparation:

The reagents are ready to use. Do not shake.

Specimen:

Serum or Heparin Plasma 20 - 25 °C Stability : 7 days at 7 days at 4 - 8 °C 1 Month at -20 °C

Discard contaminated specimens.

Test Procedure:

System Parameters		
Reaction Type (Mode)	:	Kinetic
Reaction Direction	:	Increasing
Wave Length	:	578 nm
Flow Cell Temp.	:	37°C
Zero Setting with	:	Distilled Water
Delay Time	:	5 Seconds
Measuring Time	1. I	120 Seconds
Reagent Volume	1.1	500 µl (R1) + 100 µl (R2)
Sample Volume	:	10 µl
Kinetic Factor	:	1150 (Lot Specific)
Linearity	:	300
Units	:	U/L
High Normal	:	64 U/L

Procedure :

Lipase Buffer (R1)	500 μL		
Serum/ Plasma	10 μL		
Mix and incubate for 5 min at 37 °C in an Incubator			
Lipase Substrate (R2)	ι 100 μL		
	I /		

Mix well and aspirate in to the analyzer. After 5 Sec delay measure the change of optical density per 60 seconds during 120 seconds against distilled water at 578 nms as follows:

A°	-	Exactily after 5 Seconds.
A1, A2	-	Exactily after every 60 seconds for 120 seconds.

Calculations:

From absorbance readings calculate Δ A/min and multiply by the corresponding factor 1150 at 578 nm

Lipase Activity (IU/L) = \triangle Abs / Min X **1150** (Kinetic Factor)

Performance Characteristics:

Measuring range

The test has been developed to determine lipase concentrations up to 300 U/L. When values exceed this range samples should be diluted 1 + 1 with NaCl solution (9 g/L) and the result should be multiplied by 2.



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Specificity/Interferences:

No interference was observed by ascorbic acid up to 30 mg/dL, free and conjugated bilirubin up to 60 mg/dL, hemoglobin up to 500 mg/dL and lipemia up to 1000 mg/dL triglycerides.

Sensitivity / Limit of Detection:

The lower limit of detection is 3 U/L.

Precision:

According to protocol EP-5 of the NCCLS (National Committee of Clinical Laboratory Standards)

Within run precision	Mean	SD	CV
n = 40	[U/L]	[U/L]	[%]
Sample 1	13.4	0.24	1.81
Sample 2	58.9	0.60	1.01
Sample 3	103	1.50	1.45
Between day precision	Mean	SD	CV
n = 40	[U/L]	[U/L]	[%]
Sample 1	13.4	0.24	1.81
Sample 2	58.9	0.49	0.80
Sample 3	103	0.65	0.63

Method Comparison:

A comparison between High-Q Lipase (y) and a commercially available colorimetric test (x) using 67 samples gave following results: y = 0.06 x = 1.15 LM tr = 0.000

y = 0.96 x - 1.15 U/L; r= 0.999.

Interference:

The following concentrations were not found to affect the assay: Conjugated Bilirubin 40 mg/dl, Free Bilirubin 70 mg/dl, Haemoglobin 1000mg/dl , Intralipid 800 mg/dl, Triglycerides 1000 mg/dl

Reference Range: 0 - 64 U/L

It is strongly recommended that each laboratory establish its own normal range.

Literature

1. Lorentz K. Lipase. In: Thomas L, editor. Clinical laboratory diagnostics. 1st ed. Frankfurt: TH-Books

Verlagsgesellschaft; 1998. p. 95-7.

2. Moss DW, Henderson AR. Digestive enzymes of pancreatic origin. In: Burtis CA, Ashwood ER, editors.Tietz

Textbook of Clinical Chemistry. 3rd ed.Philadelphia: W.B Saunders Company; 1999.p. 689-708.

- 3. Tietz N, Shuey DF. Lipase in serum the elusive enzyme: an overview. Clin Chem 1993; 39: 746-56.
- Lott J, Patel ST, Sawhney AK, Kazmierczak SC, Love JE. Assays of serum lipase: analytical and clinical considerations. Clin Chem 1986; 32: 1290-1302.

Ordering information:

Ref./Cat. No.	Pack Size
P-LIP - 12	12 ml
P-LIP - 50	50 ml

Presentation (10 ml R1 + 2 ml R2 (4 x 10.5 ml R1 + 4 x 2 ml R2)

Product Features

- Two Liquid Reagents (5 Parts R1+ 1 Part R2).
- Uses 1,2-O-dilauryl-rac-glycero-3-glutaric acid (6'methylresorufin)-ester as Lipase Specfic Substrate.
- Linearity : 300 U/L.
- Measuring wavelength 578 nm.

Symbols used with IVD devices

- Two Step Kinetic Assay : 5 Sec Delay+ 120 Sec Measuring.
- No interference from Conjugated Bilirubin 40 mg/dl, Free Bilirubin 70 mg/dl, Haemoglobin 1000 mg/dl, Intralipid 800mg/dl, Triglycerides 1000 mg/dl, Ascorbic Acid 40 mg/dl

Date of manufacture Manufactured by IVD In vitro diagnostic device Keep away from sunlight (\mathfrak{R}) Do not freeze This way up Reagent Use by (yyyy-mm-dd or mm/yyyy) R CAL **Calibrator Material** LOT Batch code Temperature limitation (store at) CONTROL Control Keep dry Consult instructions for use i Keep away from rain REF Catalog Number info@parikshabio.com www.parikshabio.com



JEU Indicator

Manufactured in India by : Pariksha Biotech Pvt Ltd, Plot no. 1/B-14, SVICE, Balanagar, Hyderabad-500037 Telangana State



AN ISO 13485 Certified Company