



HDL-C

High Density Lipoprotein Cholesterol Test Kit (Enzymic Method)

Instructions for Use

Version: A/3

REF HP-HDL-C-25

Manufacturer

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Product Name

General Name: High Density Lipoprotein Cholesterol Test Kit (Enzymic Method)

Specification

Package Specification

25 Tests/Kit

Intended Use

The High Density Lipoprotein Cholesterol (HDL-C) test kit is intended for in vitro quantitative determination of High Density Lipoprotein Cholesterol (HDL-C) concentration in human serum using the Hipro immunoassay analyzer. Clinically, it is mainly used for auxiliary diagnosis of hypercholesterolemia, coronary heart disease and atherosclerosis.

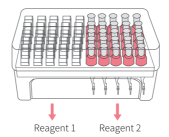
Test Principle

R1 blocks LDL cholesterol and very LDL cholesterol in the sample. Esterified high-density lipoprotein cholesterol was hydrolyzed by cholesterol lipase to generate cholesterol, and cholesterol was oxidized by cholesterol oxidase. The resulting H2O2 reacted with 4-aminoantipyrine and HDAOS to generate quinone dye. The change of light intensity produced by the reaction was proportional to the content of high-density lipoprotein cholesterol in the sample. Determination of light intensity through the standard curve to calculate the content of the sample to be measured.

Component

The Hipro HDL-C test kit consists of two reagents R1 and R2, as shown on figure 1.

Figure 1



Name	Content	Quantity	
Reagent 1	Buffer solution (PH 7.0)	100mmol/L	
(R1)	4-aminoantipyrine	1mmol/L	
	MgCl2	100mM	
Reagent 2 (R2)	Buffer solution (PH 7.0)	100mmol/L	
	HDAOS	1mM	
	Cholesterol lipase	1KU/L	
	Cholesterol oxidase	1KU/L	
	Peroxidase	4KU/L	

Storage&stability

Store the test kit at 2°C-8°C until the expiration date indicated on the label. The test kit is stable for one year when unopened.

Use up the test kit within one month after opening the package.

Do not freeze the test kit.

Do not mix different lots of the test kit.

Special Instrument Requirements

HP-AFS/1 Automatic Immunoassay System, HP-AFS/3 Automatic Immunoassay System.

Specimen type

The specimen type of Hipro HDL-C test kit is human serum.

- -Take blood samples in observance of the standard precautions for the withdrawal of biological fluids.
- -In the serum samples, ensure adequate coagulation of the samples prior to centrifugation.
- -Test tubes must be kept closed in a vertical position.
- -Do not use samples that have remained at room temperature for more than $8\,\mathrm{hours}.$
- -Hermetically seal and refrigerate the samples (from $2^{\circ}\text{C-8}^{\circ}\text{C}$) if dosage is not performed within 8 hours.
- -Freeze the sample at -20°C or lower temperature if dosage is not performed within 72 hours.
- -Due to possible evaporation effects, samples on the analyzer should be measured within 2 hours.
- -Do not use samples and controls stabilized with azide.

Storage

Serum is stable if stored at 2°C-8°C for up to 72 hours. For longer storage, aliquot, cap tightly, and freeze at -20°C for up to 30 days. Avoid repeated freezing and thawing.

Required volume

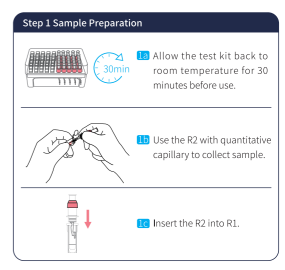
There is a quantitative capillary on R2 for required sample volume. The capillary should be fully collected.

Procedures

HP-AFS/1&HP-AFS/3 Automatic Immunoassay System

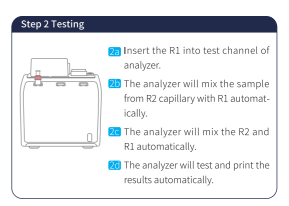
Note:

- Please read user manual of HP-AFS/1 and HP-AFS/3 before use.
- The analyzer will finish the self check after start-up.
- It is recommend to do analyzer calibration monthly and for each new lot of test kit.



Note:

- Please update the standard curve with the barcode on the R1 cuvette if a new lot test kit is to be used.
- The capillary of the R2 should be fully filled.



Calibration

Every Hipro HDL-C test kit contains specific information for calibration of the particular reagent lot, in the barcoded label

on cuvette.

Renewed calibration is recommended as follows:

- -after 1month(28days) when using the same reagent lot
- -before using new lot of reagent

Quality Control

It is recommended that each laboratory routinely uses quality control materials and establish its own control ranges. The control intervals and limits should be adapted to each laboratory's individual requirements. Multi-level controls should be tested in each HDL-C run. The HDL-C values obtained with the quality control material should not repeatedly fall outside the control ranges established in each laboratory. Suitable or published control material can be used.

Reference Value

The normal reference range should be 0.90mmol/L \sim 2.19mmol/L.

The value is indicative only and may differ from other published values as a result of differences in methods and in the population being studied. It is recommended that each laboratory establish its own reference range.

Interpretation

Results ≤ 0.90 mmol/L indicates the risk of coronary heart disease, results≥2.19 mmol/L were more common in primary hyperlipoproteinemia. Further examination and treatment are recommended.

Limitations

When bilirubin≤100µMol/L, hemoglobin≤5g/L, triglyceride ≤ 0.1%, VC≤ 0.5g/L, it has no interference to this determination.

Performance Characteristics

1. Linearity range:

 $0.1 \text{mmol/L} \sim 3.90 \text{mmol/L}, r \ge 0.995.$

The linear deviation within the range of [0.1-1.0]mmol/L should not exceed ± 0.1 mmol/L;

The linear deviation within the range of (1.0-3.90]mmol/L should not exceed \pm 10%.

2. Limit of Blank

Reagent blank absorbance< 0.05.

3. Precision

Within-lot precision (CV) ≤ 4%

Between-lot precision(R) ≤ 10%

4. Analytical sensitivity

When 1.00mmol/L sample was determined, the absorbance difference ($\triangle A/min$) > 0.01

5. Accuracy

Relative deviation (B) shall not exceed $\pm 10\%$

6 Precision

Test the control material by High Density Lipoprotein Cholesterol Test Kit (Direct Method-Selective Inhibition Method).2 times per day for 20 days (n=80) according to EP5-A2 of CLSI. The data as below:



HP-AFS/3 Automatic Immunoassay System					
Sample	Mean	Within-Run		Between-Run	
Sample	mmol/L	S.D.	%C.V.	S.D.	%C.V.
Control 1	0.76	0.021	2.8	0.023	3.0
Control 3	1.62	0.056	3.5	0.055	3.4

D.

HP-AFS/1 Automatic Immunoassay System						
Sample	Mean mmol/L	Within-Run		Between-Run		
Sample		S.D.	%C.V.	S.D.	%C.V.	
Control 1	0.80	0.025	3.1	0.023	2.9	
Control 3	1.68	0.054	3.2	0.056	3.3	

4. Methodology comparison

Compared to HDL-C LTIA(x) by test the same samples, the relative data as below:

HP-AFS/3 Automatic Immunoassay System					
Site No.	Sample Type	No.of Assays	Regression Line	Coefficient correlation	
1	plasma	50	Y= 1.03X-0.16	0.97	

The concentration of sample is about 0.1mmol/L \sim 3.90mmol/L.

Precautions and limitations

For in vitro diagnostic use

Only experienced laboratory personnel should use this test and handling all laboratory reagents should be based on normal precautions required.

⚠ Safety precautions

1.Do not smoke, eat or apply cosmetics in areas in which patients' samples or kit reagents are handled.

2.Laboratory gloves should be worn while handling patients' samples or disposing of solid or liquid wastes.

3. Disposal of all waste material should be in accordance with local guidelines.

Potential biohazard warning

Some reagents of Hipro test kits contain material of animal origin, even if they are certified as deriving from healthy animals, it is recommended to handle them with the same precaution used for potentially infectious samples.

↑ Limitations-interference

Bilirubin ≤100μmol/L, hemoglobin ≤5g/L, triglyceride ≤0.1%, VC≤0.5g/L have no influence on the determination. For diagnostic purposes, the results obtained from this assay should always be used in combination with the clinical examination. patient medical history, and other findings.

Risk Phrases

R 36/38 Irritating to eyes and skin

Safety Phases

S 26 In case of contact with eyes, rinse immediately with plenty of water and seek for medical advice.

S 37 Wear suitable gloves

S 60 This material and /or its container must be disposed of as hazardous waste.

SYMBOLS USED ON LABELS

Symbol	Usage	Symbol		
	Use-By date	\%	Do not freeze	
LOT	Batch code	80	Biological risks	
ш	Manufacturer	(3)	Do Not Reuse	
210 \$ 870	Temperature Limit			
\sum	Contains sufficient for <n> tests</n>			
	Do not use if package is damaged			
[]i	Consult Instructions for use			
*	Keep Away from Sunlight			
IVD	In Vitro Diagnostic Medical device			
EC REP	Authorized Representative in the European Community			

References

Shang Hong, Wang Yusan, Shen Ziyu et al. National Clinical Laboratory Operating Procedures (4th edition) [M]. Beijing: People's Medical Publishing House, 2015:320-323.

Approval Date&Revision Date

Approval Date: Sept 9,2015 Revision Date: Jul 15, 2021 Revision Date: Jan 1, 2023 Revision Date: Dec 22,2023 尺寸:24*25cm展开尺寸,横向三折页再垂直方向两次对折