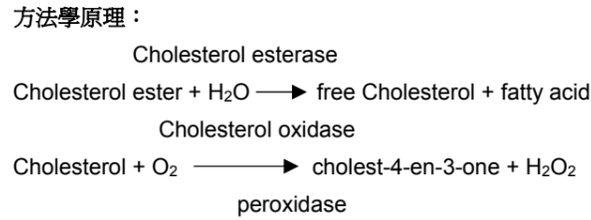




台塑膽固醇試劑 (CHOL) - Enzymatic method

效能：
用於臨床實驗體外定量分析人體血清或血漿中總膽固醇的含量。

臨床意義：
高總膽固醇血症是冠心病的主要危險因素之一。病理狀態下高或低總膽固醇血症的有原發的和繼發的兩類。高總膽固醇原發的如家族性高膽固醇血症、混合性高脂蛋白血症；繼發的見於腎病綜合症、甲狀腺機能減退、糖尿病。低總膽固醇血症原發的如家族性的無 β 或低 β 脂蛋白血症；繼發的如營養不良、肝硬化等。



反應形成紫紅色錯合物而引起 505nm 吸光值的增加。

- 試劑：**
- 產品規格：
詳見外盒包裝標示。
 - 成份與濃度：
- | | 成份 | 濃度 |
|------------------|----------------------|------------|
| R ₁ : | Phenol | 10 mmol/L |
| | Sodium cholate | 50 mmol/L |
| | 4-AA | 1.4 mmol/L |
| | Cholesterol esterase | 200 U/L |
| | Cholesterol oxidase | 180 U/L |
| | Peroxidase | 3000 U/L |

保存溫度：
2-8℃ 避光保存，請勿冰凍。

檢體：
新鮮無溶血的血清、肝素抗凝血漿。檢體採集後必須儘快離心處理。

- 操作步驟：**
- 測定主波長：505 nm 測定副波長：660nm
溫度：37℃ 比色杯光徑：1.0 cm
 - 本試劑盒為液態單試劑，可直接上機使用。

加入物	標準管	測定管	空白管
試劑 R ₁ (ml)	1.0	1.0	1.0
標準液 (ml)	0.01	----	----
檢體 (ml)	----	0.01	----
ddH ₂ O (ml)	----	----	0.01

各管立即搖勻後，於 37℃ 中放置 10 分鐘，在 505nm 處，以空白管調“零”點，分別檢測各管的吸光值。

結果計算

$$\text{總膽固醇 (mg/dL)} = \frac{A_{\text{檢體}}}{A_{\text{標準}}} \times \text{總膽固醇標準濃度 (mg/dL)}$$

參考值：
150~250 mg/dL

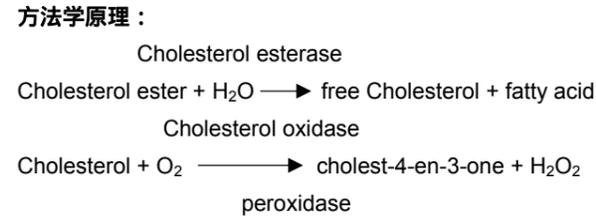
- 注意事項：**
- 本試劑請用專用標準品 (calibrator) 不另外提供質控血清 (control) 建議質控血清為 i o- Rad Lyphochek control。
 - 建議各實驗室建立獨立之品管系統，並定義專屬之參考值範圍。
 - 本檢驗試劑限由醫師或醫檢師臨床使用。
 - 本試劑線性可達 0.0m g/dL，當檢體中膽固醇濃度超過 400m g/dL 時可用生理食鹽水稀釋檢體，結果乘以稀釋倍數。
 - 為保證結果的準確性，必須在檢體加入後 30 分鐘內檢測吸光值。抗壞血酸(Vitamin C)會干擾此試劑的測定。
 - 以上操作步驟適用於手工操作及一般半自動及全自動生化分析儀。
 - 本品操作時請穿戴手套及必要之防護措施，操作中若不慎沾上，應用水或肥皂水清洗。(詳細溶液物化性請洽詢經銷商索取物質安全資料表
 - 用畢應按醫療事業廢棄物處理。(詳細溶液物化性請洽詢經銷商索取物質安全資料表
 - 有效期限見試劑盒上標籤所示。
 - 經專業人員建議，試劑與檢體用量可根據所用分析儀的要求按比例調整，其吸光值不變，不影響監測結果。
 - 試劑特性及參數設定請參見第四頁。



台塑胆固醇试剂 (CHOL) - Enzymatic method

效能：
用于临床实验体外定量分析人体血清或血浆中总胆固醇的含量。

临床意义：
高总胆固醇血症是冠心病的主要危险因素之一。病理状态下高或低总胆固醇血症的有原发的和继发的两类。高总胆固醇原发的如家族性高胆固醇血症、混合性高脂蛋白血症；继发的见于肾病综合症、甲状腺机能减退、糖尿病。低总胆固醇血症原发的如家族性的无 β 或低 β 脂蛋白血症；继发的如营养不良、肝硬化等。



反应形成紫红色错合物而引起 505nm 吸光值的增加。

- 试剂：**
- 产品规格：
详见外盒包装标示。
 - 成份与浓度：
- | | 成份 | 浓度 |
|------------------|----------------------|------------|
| R ₁ : | Phenol | 10 mmol/L |
| | Sodium cholate | 50 mmol/L |
| | 4-AA | 1.4 mmol/L |
| | Cholesterol esterase | 200 U/L |
| | Cholesterol oxidase | 180 U/L |
| | Peroxidase | 3000 U/L |

保存溫度：
2-8℃ 避光保存，請勿冰凍。

檢體：
新鮮無溶血的血清、肝素抗凝血漿。檢體採集後必須儘快離心處理。

- 操作步驟：**
- 測定主波長：505 nm 測定副波長：660nm
溫度：37℃ 比色杯光徑：1.0 cm
 - 本試劑盒為液態單試劑，可直接上機使用。

加入物	標準管	測定管	空白管
試劑 R ₁ (ml)	1.0	1.0	1.0
標準液 (ml)	0.01	----	----
檢體 (ml)	----	0.01	----
ddH ₂ O (ml)	----	----	0.01

各管立即搖勻後，於 37℃ 中放置 10 分鐘，在 505nm 處，以空白管調“零”點，分別檢測各管的吸光值。

結果計算

$$\text{總膽固醇 (mg/dL)} = \frac{A_{\text{檢體}}}{A_{\text{標準}}} \times \text{總膽固醇標準濃度 (mg/dL)}$$

參考值：
150~250 mg/dL

- 注意事項：**
- 本試劑請用專用標準品 (calibrator) 不另外提供質控血清 (control) 建議質控血清為 i o- Rad Lyphochek control。
 - 建議各實驗室建立獨立之品管系統，並定義專屬之參考值範圍。
 - 本檢驗試劑限由醫師或醫檢師臨床使用。
 - 本試劑線性可達 0.0m g/dL，當檢體中膽固醇濃度超過 400m g/dL 時可用生理食鹽水稀釋檢體，結果乘以稀釋倍數。
 - 為保證結果的準確性，必須在檢體加入後 30 分鐘內檢測吸光值。抗壞血酸(Vitamin C)會干擾此試劑的測定。
 - 以上操作步驟適用於手工操作及一般半自動及全自動生化分析儀。
 - 本品操作時請穿戴手套及必要之防護措施，操作中若不慎沾上，應用水或肥皂水清洗。(詳細溶液物化性請洽詢經銷商索取物質安全資料表
 - 用畢應按醫療事業廢棄物處理。(詳細溶液物化性請洽詢經銷商索取物質安全資料表
 - 有效期限見試劑盒上標籤所示。
 - 經專業人員建議，試劑與檢體用量可根據所用分析儀的要求按比例調整，其吸光值不變，不影響監測結果。
 - 試劑特性及參數設定請參見第四頁。



MeDiPro CHOLESTEROL TEST (CHOL) - Enzymatic method

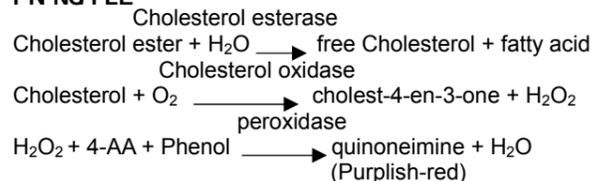
INTENDED USE

For the use of quantitative determination of cholesterol in serum or plasma.

CLINICAL SIGNIFICANCE

Total cholesterol in the blood is elevated in idiopathic hypercholesterolemia, primary and secondary hyperlipidemias, diabetes mellitus, nephrotic syndrome, hypothyroidism and biliary obstruction. Pregnancy may also be accompanied by a moderate increase of cholesterol level. Decrease of cholesterol is seen in patients with severe hepatitis and occasionally in severe anemia or infection.

PRINCIPLE



REAGENT

- Package: please see the reagent box label shown.
- Components:

Component	Conc.
Phenol	10 mmol/L
Sodium cholate	50 mmol/L
4-AA	1.4 mmol/L
Cholesterol esterase	200 U/L
Cholesterol oxidase	180 U/L
peroxidase	3000 U/L

STORAGE

The standard is stable up to the end of the indicated expiration date. If stored at 2-8 °C, reagent should be protected from light and contamination should be avoided.

Do not freeze the reagent!

SPECIMEN COLLECTION AND PREPARATION

Serum or heparinized plasma is recommended. The use of oxalate, citrate, fluoride and EDTA might result in slightly lower cholesterol values.

PROCEDURES

- Main wavelength : 505 nm
Sub. wavelength : 660nm
Reaction Temperature : 37°C
Optical path length : 1.0 cm
- This kit contains single reagent, ready to use.

	Control	Specimen	Blank
R ₁ (ml)	1.0	1.0	1.0
Control (ml)	0.01	----	----
Specimen (ml)	----	0.01	----
ddH ₂ O (ml)	----	----	0.01

Mix, incubate at 37°C for 10 min, and read the absorbance against reagent blank. $A = A_{505} - A_{660}$

CALCULATION

With standard or calibrator

$$\text{Cholesterol (mg/dL)} = \frac{A_{\text{sample}}}{A_{\text{std./cali.}}} \times \text{conc. Std./cali. (mg/dL)}$$

REFERENCE RANGE

150~250 mg/dL

WARNINGS AND PRECAUTIONS

- This kit offers an optional calibrator, which is sold individually. Bo-Rad Lyphochek control is recommended to use as serum control.
- Each laboratory has to perform the quality control test to assure the results being reliable before running the specimen tests.
- This kit is for professionals and *in vitro* diagnostic use only.
- To ensure the accuracy of result, the absorbance should be measured within 30 minutes after sample addition. Specimen should avoid ascorbic acid which might interfere with the test result.
- The test is developed to determine cholesterol concentrations up to 400 mg/dL. When values exceed this range, samples should be diluted with normal saline and calculate the results by multiplying the dilution factor.
- The above-mentioned procedures are suitable either for the general semi-automatic, full-automatic biochemical analysis instrument or manual operation.
- Since all specimens are potentially infectious, they should be handled with appropriate precautions and practices in accordance with Biosafety level 2 as recommended by USA National Biosafety in Microbiological and Biomedical Laboratories, and in accordance with National or local regulations related to the safety precautions of such materials.
- When using reagent please refer to the local legal requirements.
- Please refer to the manufacturer's safety data sheet and the product labeling for information on potentially hazardous components. MSDS could be obtained from local dealer.
- According to the technical suggestion, the volume of reagent and specimen could be adjusted in a ratio for full-automatic biochemical analysis instrument use. It will not affect the absorbance and the result.



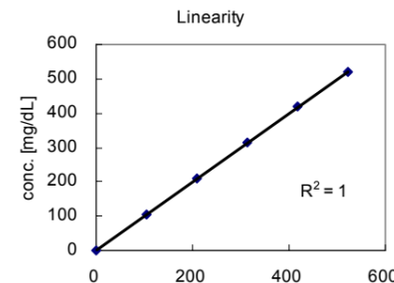
MeDiPro CHOLESTEROL TEST (CHOL) - Enzymatic method

11. Validity please see the reagent box label shown

REAGENT CHARACTERISTICS

1. Precision (Within run)			
N=15	Mean[mg/dL]	SD [mg/dL]	CV[%]
Sample1	243	2.50	1.03
Sample2	128	0.96	0.75
Sample3	131	1.08	0.83

2. Linearity



This kit has a good linearity up to 500mg/dL.

3. Interference

Interference	Influence effect
Hemoglobin	No interference was observed by hemoglobin up to 200mg/dL
Ascorbic acid	Not suitable when sample contain reducing agent
Bilirubin (free form)	No interference was observed by bilirubin up to 12mg/dL
Bilirubin (conjugate form)	No interference was observed by bilirubin up to 16mg/dL
Intrafat	No interference was observed by intrafat up to 1.0%

4. Stability

Expire day	1 year
Open vial stability	30 day

REFERENCE

- Allain, cc, Poon, LS, Chan, LS, et.al. Enzymatic determination of total serum cholesterol. Clin. Chem. 1974, 20:470.
- Bishop ML, Duben-Engelkirk JL, Fody EP. Clinical Chemistry: principles, procedures, correlations. Fourth edition, 2000.

PARAMETER SETUP

Hitachi 7170/917 Applications

TEST	[CHOL]
ASSAY CODE	[1 POINT]: [34]-[0]
SAMPLE VOLUME	[2]
R1 VOLUME	[200]
R2 VOLUME	[0]
WAVELENGTH (nm)	[660][505]
CALIB. METHOD	[Linear]

Hitachi 7150/717 Applications

TEST	[CHOL]
ASSAY CODE	[1 POINT]: [50]-[0]
SAMPLE VOLUME	[2]
R1 VOLUME	[200]
R2 VOLUME	[0]
WAVELENGTH(nm)	[660][505]
CALIB. METHOD	[Linear]

ORDERING INFORMATION

Cat. No.	Product	Package
BC-0014M	MeDiPro CHOLESTEROL TEST	R1 6×20ml
BC-0014A	MeDiPro CHOLESTEROL TEST	R1 4×60ml
BC-0014B	MeDiPro CHOLESTEROL TEST	R1 4×100ml
BC-0014C	MeDiPro CHOLESTEROL TEST	R1 4×300ml
BC-0014D	MeDiPro CHOLESTEROL TEST	R1 4×500ml



FORMOSA BIOMEDICAL TECHNOLOGY CORP.

F-5F, No. 201, Tunghua N. Rd, Taipei, 105, Taiwan Website: <http://www.fbc.com.tw/>
 TEL: +886-2-2712-2211 #7822 FAX: +886-2-2717-8381
 Factory: No. 3, Longchuan Rd, Longtang Village, Jiaosi, Yilan County, 262, Taiwan



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