A simple and sensitive enzymatic method for cholesterol quantification in macrophages and foam cells

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Abstract A precise and sensitive method for measuring cellular free and esterified cholesterol is required in order to perform studies of macrophage cholesterol loading, metabolism, storage, and efflux. Until now, the use of an enzymatic cholesterol assay, commonly used for aqueous phase plasma cholesterol assays, has not been optimized for use with solid phase samples such as cells, due to inefficient solubilization of total cholesterol in enzyme compatible solvents. We present an efficient solubilization protocol compatible with an enzymatic cholesterol assay that does not require chemical saponification or chromatographic separation. Another issue with enzyme compatible solvents is the presence of endogenous peroxides that interfere with the enzymatic cholesterol assay. We overcame this obstacle by pretreatment of the reaction solution with the enzyme catalase, which consumed endogenous peroxides resulting in reduced background and increased sensitivity in our method. Finally, we demonstrated that this method for cholesterol quantification in macrophages yields results that are comparable to those measured by stable isotope dilution gas chromatography with mass spectrometry detection. In conclusion, we describe a sensitive, simple, and high-throughput enzymatic method to quantify cholesterol in complex matrices such as cells. — Robinet, P. Z. Wang, S. L. Hazen, and J. D. Smith. A simple and sensitive enzymatic method for cholesterol quantification in macrophages and foam cells. J. Lipid Res. 2010. 51: 3364–3369.

Supplementary key words cell cholesterol assay • cholesterol solubilization • catalase

Cholesterol accumulation in macrophage foam cells is one of the earliest histological features of atherosclerosis in the arterial intima (1). In order to perform studies of macrophage cholesterol loading, metabolism, storage, and efflux, one requires a precise, accurate, and sensitive method for measuring cellular free and esterified cholesterol levels. The most commonly used assays to quantify cholesterol levels can be separated into two groups: 1) analytical methods such as gas-liquid chromatography or liquid chromatography coupled with flame ionization or mass spectrometry detection and quantification (2–4); and 2) enzymatic assays, which can be colorimetric (5, 6) or fluorometric (7). However, the chromatographic and mass spectrometry methods require specialized equipment and chemical saponification of cholesterol esters; furthermore, these methods are time consuming and require processing samples one at a time. In contrast, the colorimetric and fluorometric enzymatic methods are simple, sensitive, and relatively fast assays that can process many samples at once. The principal of a sensitive fluorometric assay for the measurement of total cholesterol is shown below, with free cholesterol measured by omitting the first esterase step, and cholesterol mass in cholesterol esters calculated as the difference between the total and free cholesterol contents.

\[
\text{Cholesterol esters} \xrightarrow{\text{Cholesterol Esterase}} \text{Cholesterol} + \text{fatty acids}
\]

\[
\text{Cholesterol} + \text{O}_2 \xrightarrow{\text{Cholesterol Oxidase}} \text{Cholest-4-ene-3-one} + \text{H}_2\text{O}_2
\]

\[
\text{H}_2\text{O}_2 + \text{ADHP} \xrightarrow{\text{HRP}} \text{resorufin (fluorescent)}
\]

The enzymatic method is the principal method used for measuring cholesterol levels in plasma, where cholesterol is solubilized in an aqueous solution through its incorporation in lipoproteins. However, in order to accurately measure the cholesterol content of cells or tissue by this method, one must identify a suitable solvent that allows good recovery of free and esterified cholesterol in solution and that is compatible with the enzymes and substrates. Previously, the use of the enzymatic cholesterol assay for cells and tissues has been limited due to the inadequate quantitative recovery of free cholesterol and cholesterol esters in enzyme compatible solvents; in fact, a methods paper has argued that the enzymatic method cannot be used for this purpose due to poor extraction and solubili-

Abbreviations: AcLDL, acetylated low density lipoprotein; ADHP, 10-acetyl-3,7-dihydroxyphenoxazine; BCA, bicinchoninic acid; NP40, nonidet P-40.

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zation, which led to an underestimation of the cholesterol mass compared with an analytical method (8). Here, we have identified a suitable and simple solvent system that yields excellent cholesterol recovery. Although this solvent initially led to high background in the enzymatic cholesterol assay, we discovered that pretreatment of this solvent with the enzyme catalase consumed endogenous peroxides, thus resulting in our ability to measure cellular cholesterol content by the simple fluorometric enzymatic assay. We demonstrate that this method for cholesterol quantification in macrophages yields results that are comparable to those measured by gas chromatography-mass spectrometry (GC-MS).

MATERIALS AND METHODS

Reagents

[14C]Cholesterol, [3H]cholesterol, and [3H]cholesterol olate were purchase from Perkin Elmer. Bovine liver catalase, cholesterol oxidase from Streptomyces sp., cholesterol esterase from Pseudomonas sp., horseradish peroxidase (HRP), deuterated cholesterol [3H 2,2,3,4,4,6], and Sylon® HTP (HMDS+TMCS+Pyridine) were purchased from Sigma-Aldrich. 10-acetyl-3,7-dihydroxyphe- 

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Fluorometric cholesterol assay

Cholesterol loading and extraction. RAW 264.7 mouse macrophages were plated in 6-well dishes unless otherwise specified and cholesterol loaded by incubation with the specified concentration of AcLDL for 24 h. Unloaded cells were used as a control. Total cell cholesterol was extracted from the cells using 1 ml hexane/isopropanol (3:2, v:v). The extracts were transferred to microfuge tubes and dried. After solvent extraction, any residual solvent remaining on the cells was evaporated at room temperature. Then the protein from the same wells was dissolved by addition of 0.2 ml sodium hydroxide. The plate was incubated at 37°C for 3 h then rocked for 5 min at room temperature. The protein lysates were transferred to microfuge tubes and protein concentration was determined using the BCA assay.

Free cholesterol quantification. Cholesterol standards, samples, and blanks were dissolved in isopropanol:NP40 (9:1, v:v) and treated similarly, using 1 ml to redissolve each well of cells from a 6-well dish. In a black 96-well plate, 10 µL of a 100 µM catalase solution was distributed in each well and 40 µL of each sample was mixed followed by 15 min incubation at 37°C in order to eliminate any peroxides present in reagents or samples. Then 150 µL of reagent A (0.1 M potassium phosphate buffer, pH 7.4, 0.25 M NaCl, 5 mM cholic acid, 0.1% Triton X-100, 0.3 U/ml cholesterol oxidase, 1.3 U/ml HRP, and 0.4 mM ADHP) was added and mixed in each well. The plate was incubated at 37°C for an additional 15 min and fluorescence was read at an excitation wavelength of 530 nm and an emission wavelength of 580 nm. The cholesterol mass of the 40 µM aliquot of the unknown samples was determined by linear regression using the fluorescence emission of the blanks and the 40 µM cholesterol standards (20 to 800 ng range). The final cholesterol mass in the 1 ml samples was calculated by multiplying by 25.

Total cholesterol quantification. The same procedure was followed for total cholesterol except that reagent A was supplemented with 0.67 U/ml cholesterol esterase, yielding a final concentration of 0.5 U/ml.

Cholesterol esters quantification. The cholesterol mass in cholesterol esters was determined by subtracting the free cholesterol values from the total cholesterol values.

GC-MS

Total cholesterol extraction and cholesterol esters saponification. Macrophages were scraped from each well of a 6-well plate and were resuspended in 400 µL water and 100 µL of 1 µg/ml deuterated internal cholesterol standard ([3H 2,2,3,4,4,6]cholesterol) in isopropanol. Two milliliters of hexane/isopropanol (3:2) and 20 µL of acetic acid were then added to the cell suspension. After vortexing and centrifugation, cholesterol and its derivatives present in the organic phase were collected. The aqueous phase was reextracted by the addition of 1ml hexane followed by vortexting and centrifugation. The hexane layer was collected and combined with the previous organic phase. The remaining aqueous phase containing the extracted cells was used to determine protein concentration by the BCA assay. The combined organic extract was divided into two parts. One part was dried under nitrogen for free cholesterol quantification. The other part was dried under nitrogen and cholesterol esters were saponified in 100 µL 0.5 M potassium hydroxide in methanol for 1 h at 37°C followed by adding 100 µL of 1 M hydrochloric acid, 300 µL water, and 1 ml isopropanol/hexane/acidic acid (40:10:1, v:v:v). The total cholesterol was extracted by two successive additions of 1 ml hexane and dried under nitrogen.

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**RESULTS**

**Cholesterol extraction and recovery**

We first validated that hexane:isopropanol (3:2) yielded quantitative recovery of total \[^3\text{H}\]cholesterol from cholesterol loaded RAW264.7 cells. The extracted dpm represented 98.6 ± 2% (N = 4 ± SD) of the total cells lysate control. For the enzymatic cholesterol assay, the dried hexane:isopropanol extract must then be redissolved in a solvent compatible with the enzymes and substrates. Cullen et al. (8) reported that isopropanol, ethanol, or 10% triton X-100, all suitable for enzyme activity, allowed at best 50% free cholesterol recovery and 20% for cholesterol esters. We tested the suitability of isopropanol:NP40 (9:1) for sterol extraction by adding it to a dried cell hexane:isopropanol extract containing known amounts of \[^{14}\text{C}\]cholesterol and \[^{3}\text{H}\]cholesterol oleate. We observed that the isopropanol:NP40 mixture yielded 102 ± 3% and 96 ± 7% recovery of free cholesterol and cholesterol esters, respectively (N = 4, ± SD).

**Enzymatic assay optimization**

**Catalase.** One of the limitations for the use of our enzymatic method resides on the apparent contamination of the isopropanol:NP40 mixture with endogenous peroxides, a common problem with some solvents, resulting in high fluorescence background of sample blanks. In order to counteract this problem, we pretreated the blanks, standards, and samples with a final concentration of 20 U/ml of catalase for 15 min at 37°C before cholesterol quantification. This greatly reduced the background fluorescence by at least 75%. Standards curves were prepared in presence or absence of catalase and the results are shown in Fig. 1. Doubling the amount of catalase to 40 U/ml did not alter the results (data not shown); thus, 20 U/ml was retained as the standard dose. The data showed that catalase treatment not only reduced background but also improved the sensitivity of the assay as shown by the higher slope obtained with this treatment.

**Sensitivity and reproducibility.** We wanted to determine the lowest number of unloaded cells that our technique was sensitive enough to detect. We measured total cholesterol in different numbers of cells recovered from a T75 flask (10⁴ to 10⁵ cells) and found we could detect both protein (4.88 ± 0.02 µg) and total cholesterol (88 ± 10 ng) in 8 × 10⁴ cells, yielding 18.1 ± 2.0 µg total cholesterol/mg cell protein. To determine the reproducibility of this assay, one unloaded and one AcLDL loaded (100 µg/ml) cell extract were subjected to total and free cholesterol assays using six technical replicates each. The coefficient of variation for the technical replicates of these four assays ranged from 2.2% to 4.4%, thus showing excellent technical reproducibility. In this experiment, the unloaded and AcLDL loaded cells yielded 22.4 and 128.6 µg total cholesterol/mg cell protein, respectively.

**Esterase efficiency.** In order to quantify total cholesterol, cholesterol esters need to be hydrolyzed into free cholesterol by cholesterol esterase. Thus, we needed to make sure that the concentration of cholesterol esterase as well as the incubation time were sufficient to achieve the hydrolysis of all cholesterol esters into free cholesterol. Using the cholesteryl oleate standard, we assayed 400 ng cholesterol (cor-

![Fig. 1. Standard curves for the determination of cholesterol.](image-url)
responding to 672 ng cholesteryl oleate) in the presence of 0.5 U/ml esterase. We obtained 393.5 ± 9.9 ng corresponding to 98.3 ± 2.5% of the theoretical amount; thus, the esterase dose used could efficiently convert up to 672 ng of cholesteryl oleate into free cholesterol in our assay.

Comparison of the enzymatic and the GC-MS assays

To compare the efficiency and accuracy of the enzymatic assay with the GC-MS assay, we performed three separate experiments with unloaded cells in which duplicate or triplicate wells were assayed for cholesterol by each method. The total cholesterol levels were 30.0 ± 8.9 and 32.0 ± 8.5 µg/mg cell protein for the enzymatic and GC-MS assays, respectively (Fig. 2, not significant). In these unloaded cells, the bulk of the cholesterol was detected as free cholesterol with similar values in both assays (Fig. 2). Although the overall cholesterol esters levels were low in these unloaded cells, they were somewhat higher in the GC-MS assay; however, this difference was not significant (Fig. 2).

In order to compare the enzymatic and GC-MS assays over a broad range of cholesterol loading, we incubated RAW 264.7 cells with 0, 10, 20, 40, 80, or 120 µg/ml AcLDL in duplicate. Using two internal standards for recovery, as described in the Methods section, each well was assayed for total and free cholesterol by both the enzymatic and GC-MS methods. AcLDL incubations led to dose dependent increases in cellular total cholesterol that are mainly due to increases in cholesterol esters (Fig. 3). In this experiment, the GC-MS assay yielded higher values than the enzymatic assay, with 19% higher total cholesterol levels observed in the cells loaded with 120 µg/ml AcLDL. However, in a separate experiment comparing the two methods using cells loaded with 100 µg/ml AcLDL, we observed the opposite bias, with 8% higher total cholesterol values obtained by the enzymatic assay (193 ± 17 µg total cholesterol/mg cell protein) compared with the GC-MS method (178 ± 5 µg total cholesterol/mg cell protein). In the cells treated with varying doses of AcLDL, we plotted the values of total, free, and esterified cholesterol for each well determined by the enzymatic versus GC-MS methods (Fig. 4). The total cholesterol levels were highly correlated ($r^2 = 0.97, P < 0.0001$), with a slope of 1.08 ± 0.06 (Fig. 4A). The bias in this assay was due to the y-intercept, which was 19.3, indicative of a systematic error. The range of free cholesterol levels did not vary as much but they were still well correlated between the two methods ($r^2 = 0.77, P = 0.0002$, slope = $1.01 ± 0.17$, y-intercept = 13.8, Fig. 4B). The cholesterol esters levels were highly correlated between these two methods ($r^2 = 0.97, P < 0.0001$, slope = $1.05 ± 0.06$, y-intercept = 9.0, Fig. 4C).

**DISCUSSION**

In this report, we describe a sensitive and easy method to quantify total and free cholesterol content in cultured cells, which does not require chemical saponification or specialized equipment other than a fluorescence plate reader.
We demonstrated efficient extraction of cellular sterols and good recovery by redissolving in an isopropanol:NP40 solution that is compatible with the enzymatic assay. We also found that pretreatment of the samples and standards with catalase to remove endogenous peroxides in the isopropanol:NP40 solution was beneficial in reducing the background and increasing sensitivity. We determined that there was no need to specifically inactivate the catalase before the enzymatic cholesterol quantification, which is dependent upon hydrogen peroxide generation. A prior study also used catalase treatment before measurement of cholesterol esters by an enzymatic assay, and these investigators also did not need to inactivate the catalase (10). The reason that catalase does not need to be inactivated is most likely due to catalase’s instability at low concentra-

tions at 37°C (11, 12), and the >1000-fold higher affinity of HRP versus catalase for hydrogen peroxide (13, 14).

To evaluate the accuracy of the enzymatic method, we compared it with stable isotope dilution GC-MS, the gold-standard method for free cholesterol and cholesterol ester quantification. On the whole, the results of the high-throughput enzymatic assay were highly correlated with the GC-MS assay. However, in specific assays we could observe one method yielding higher cholesterol levels than the other method. In three separate experiments with unloaded cells, the two methods on average yielded comparable results (Fig. 2), demonstrating that the bias did not consistently affect one assay over the other. In two separate experiments with cholesterol loaded cells, one yielded higher levels for the enzymatic assay, whereas the other yielded higher levels for the GC-MS assay. In the experiment where varying amounts of AcLDL were used to yield a range of cholesterol loading, our analysis showed that the two methods were highly correlated in detecting total, free, and esterified cholesterol (Fig. 4). However, in this study the GC-MS method yielded higher values than the enzymatic method, which was reflected in the y-intercept having values >0, whereas the slope was still ~1. Thus, both methods could comparably detect increasing amounts of total and free cholesterol. We speculate that the y-intercept difference may be due to a systematic error introduced by use of different cholesterol standards, different operators using different pipettors, and different types of calculations used to determine the cholesterol mass. For example, the enzymatic assay uses a cholesterol standard curve in the linear range to calculate the cholesterol mass in unknown samples. However, the GC-MS results are calculated using an internal deuterated cholesterol standard in each sample that is present at <1% of the sample cholesterol level. Thus, a small variation in the integration of the internal standard peak could lead to a systematic variation in the calculated cholesterol content.

In conclusion, we described a sensitive, simple, and rapid method to quantify cholesterol in cultured macrophages. This method allowed the detection of cholesterol in 80,000 cells prior to cholesterol loading. Combined with the use of 96-well plates, this assay can process a large number of samples per day, which may further be augmented through assay automation. This assay may be widely applicable to all types of cells and tissues after determination that the isopropanol:NP40 mixture fully dissolves all cholesterol in the solvent extract.

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