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Cholesterol metobolism of macrophages in relation to the presence of *Mycobacterium leprae*

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Abstract. Macrophages phagocytose *Mycobacterium leprae* and live bacilli inside such macrophages alter the lipid metabolism. There is increased accumulation of cholesterol ester in the bacteria infected cells. This increase appears to be due to the decreased level of esterase enzyme that could hydrolyse cholesterol esters. Associated with decreased level of this enzyme is the reduced amount of protein synthesis. Increased cholesterol ester may be responsible for conversion of macrophages into foamy cells in the presence of *M. leprae*.

Keywords. Mycobacterium; cholesterol; cholesterol ester level; macrophage.

Introduction

Cholesterol is an important lipid component of macrophages and its role has been identified in various structural and functional aspects of the macrophages (Day, 1967; Brown et al., 1980). Macrophages can phagocytose foreign particles, live or otherwise. Accordingly leprosy causing bacilli, Mycobacterium leprae, are also taken in by the macrophages. Such an uptake could be shown in vitro in macrophage cultures as well as in vivo in human tissues from lepromatous leprosy patients. In leprosy patients there is a tendency to develop foamy type of cells and these cells are primarily macrophages which have high lipid contents. Such foamy macrophages are present at sites where there is an infiltration of bacilli and immune competant cells (Yamamoto et al., 1958; Imaeda, 1960; Skinsness, 1970). It was therefore of interest to study lipid metabolism of macrophages in relation to the presence of *M. leprae* which are easily phagocytosed by the macrophage Kondo and Kanai (1976) had shown that M. tuberculosis would induce accumulation of cholesterol ester in macrophages and indicated that this ester could be a major component of the foamy droplets seen in tissue macrophages during tuberculosis inflammation. With this background information it was thought that the behaviour of macrophages in relation to M. leprae need to be studied with special reference to lipid metabolism. Since a major alteration was indicated primarily in cholesterol in some of our preliminary experiments, it was chosen as a component of interest.

Material and methods

Microorganism

Mycobacterium leprae were obtained from lepromatous tissue of bacillary positive, treated or untreated patients. Bacilli were prepared as per the method of Ambrose *et al.* (1978). Such isolated bacilli were acid-fast and free from other contaminating bacteria. These do not grow in normal mycobacterial media. The bacilli were counted and 5×10^6 bacilli were added to each leighton tube containing cultured macrophages. *M. leprae* obtained from infected armadillo tissue were also used in some of the experiments. The tissue was supplied by Dr. E. Storrs, Florida, USA.

Collection of macrophage from peritoneal cavity

Macrophages from Swiss white mice were obtained by injecting 5 ml of Eagle's minimum-essential medium + 20% inactivated human AB serum into the peritonial cavity, after killing the animal by cervical dislocation. The peritoneal fluid was collected after agitating the cavity and 0.7 ml of the fluid was added to each leighton tube.

Lipid synthesis by macrophages

The macrophages obtained from the peritoneal fluid adhered to the leighton tubes. The medium was changed every 24 h; thereby removing nonadherant cells. After 3 days of such culturing, esterase positive adherant cells were predominately distributed as a uniform layer in the leighton tubes. There were no contaminating neutrophils and non-adherant lymphocytes were not present in significant numbers. Such tubes were infected with *M. leprae* (5×10^6 bacilli/leighton tube). The control tubes did not receive the *M. leprae* inoculum. After 24 h of phagocytosis, the excess bacilli were removed and the macrophages were incubated with [¹⁴C]-acetate or [³H]-cholesterol for 4 days for studying uptake and synthesis of lipids.

Extraction and separation of lipids

Following incubation, the macrophages were scrapped off the surface of the glasstubes, the cells counted and the lipids were extracted according to Dole's method (Doles, 1956). Lipids were separated by thin-layer chromatography on 20×20 cm plates of silicia-Gel G (Chemical division, Glaxo Laboratory). The plates were developed at room temperature in a solvent media of hexane: ether: acetic acid (80 : 20 : 1) to separate the lipids. Identification of the lipid spots was made by stainining with iodine vapours, using standard lipids as reference.

Determination of incorporation into lipids

Following identification of the lipid spots, the silica gel corresponding to each spot was carefully scrapped into vials after evaporation of Iodine and full decolourisation. Scintillation fluid (10 ml) was added to each vial and radioactivity monitored by using Kontron MR-300 automatic scintillation counter.

Macrophageholesterol

 $[{}^{3}$ H]-Cholesterol was obtained as crude tritiated product from Bhabha Atomic Research Centre, Trombay and purified by repeated thin layer chromatography and 0.16 μ Ci was used for uptake studies. $[{}^{14}$ C]-Cholesterol oleate (Sp. Activity 50.8 μ Ci/m mol) was obtained from Radiochemical Centre, Amersham and 0.1 mCi was added per leighton tube. $[{}^{14}$ C]-Acetate 0.5 μ Ci (Sp. activity 56.7 μ Ci/m mol) obtained from Bhabha Atomic Research Centre, Bombay, was added in each leighton tube used for incorporation experiment.

Estimation of esterase activity, was carried out according to the method of Vahouny *et al.* (1968). Labelled [¹⁴C]-cholesterol oleate (0.2 μ Ci) in 50 μ l acetone was added *via* microsyringe beneath the surface of 2 ml of enzyme preparation containing 200 μ mol of potassium phosphate buffer pH 7.4. Incubations were carried out for 1 h at 30 C in a metabolic shaker. Reaction was stopped after 2 h by adding the extraction mixture containing solvents. Lipids were extracted and separated by thin layer chromatography and radioactivity associated with oleic acid and cholesterol ester spots were recorded.

Results

About 60% of the macrophages adherent to the glass had bacilli as demonstrated by the presence of acid fast *M. leprae* inside them. The macrophages in culture with or without *M. leprae* are able to incorporate [¹⁴C]-acetate into lipids (table 1). However, macrophages with *M. leprae*, incorporated much less [¹⁴C]-acetate into cholesterol. A similar lowered incorporation is also seen, when heat killed *M. leprae* are used as a control. An interesting feature, however, was that the radioactivity associated with cholesterol ester was higher in *M. leprae* infected cultures as compared to the uninfected or those infected with killed bacilli. Data in table 2 present the ratio of acetate incorporated in cholesterol and the ester and it is seen the ratio of cholesterol ester to cholesterol in each experiment which is higher in the infected macrophages and this increase is statistically significant (P<0.01). This is reflected in the ratio of ester to cholesterol in all the five experiments (table 1).

When labelled cholesterol was used in incorporation studies it was observed that the total cholesterol uptake was significantly lowered in the bacteria (M. *leprae*) infected cultures (table 3) the ratio of cholesterol ester to cholesterol was also higher in the infected culture compared to the controls.

The increased incorporation into cholesterol ester fraction (monitored by calculating the ratio of incorporation into cholesterol ester: cholesterol) appears to be correlated to size of the *M. leprae* inoculum (figure 1). It is to be noted that only part of the added bacilli get phagocytosed. The uptake of labelled cholesterol and conversion of the label to the ester form have also shown to exhibit different kinetic patterns depending upon whether the macrophage culture is infected or not. While cholesterol uptake increases linearly up to 4 days, conversion to ester appears to be proceeding at maximum rate by 3 days itself. [¹⁴C]-Labelled cholesterol uptake by infected cultures showed marked reduction between day one and three of culture, relatively conversion to cholesterol ester did not show much change (figure 2).

<0.01 (a-c)

p value <0.25 (a-b)

Table 1. Acetate incorporation into cholesterol and cholesterol-ester in macrophages with and without M. leprae. (com incornorated 10⁶ macronhages in 5 senarate experiments)

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		Cholestero	1			Chole	esterol ester	
Expt. No.	Macrophage	Macrophage killed M. lep	c+ rae	lacrophage + M. <i>leprae</i>	Macrophage	e Mac kille	rrophage + d <i>M. leprae</i> b	Macrophage – M. <i>leprae</i> c
01 m 4 m	0.39 0.11 0.11 0.89 0.23	0.34 0.08 0.073 0.71		0.27 0.26 0.11 0.81 0.19	0.085 0.13 0.16 0.19 0.048		0.07 0.065 0.043 0.10 0.073	0.25 0.38 0.14 0.35 0.094
Tabl	 a. [³H]-Choicsterol Cholesterol level 	uptake* by norm t in	al and infected Ch	macrophages and iolesterol ester lev	l level of choleste vel in	rol ester synthe Ratio of rad	esised inside the ioactivity in ester 10 ⁶ marconhau	cells. r/cholesterol in
xpt. Macrop	hage Macrophage + heat-killed M. leprae	Macrophage+ M. <i>leprae</i>	Macrophage	Macrophage + heat-killed - M. <i>leprae</i>	Macrophage+ M. <i>leprae</i>	Macrophage a	Macrophage + heat-killed M. leprae b	Macrophage + M. leprae c
1 465 2 258 3 205(4 697)	1 1651 4 1633 0 1647 2 3527 2 2297	1673 1159 658 814 2676	361 394 528 572 570	156 145 230 224 408	200 400 302 881	0.078 0.15 0.26 0.08 0.21	0.094 0.09 0.14 0.16 0.18	0.12 0.34 0.46 0.265 0.33
* as com 1	adioactivity/10 ⁶ macro	the second in the second	omine our of the other					

Macrophage cholesterol

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Figure 1. The influence of the number of bacteria, exposed to macrophages, on the ratio of the level of cholesterol ester to cholesterol. The uptake studies were carried out as described in the text using labelled cholesterol.



Figure 2. The level of cholesterol and cholesterol ester in macrophages incubated with and without *M. leprae* for various incubation periods in the presence of labelled cholesterol.

(O), Cholesterol in macrophage; (\bullet) cholesterol in macrophage plus *M. leprae*; (\triangle), cholesterol ester in macrophage; (\blacktriangle), cholesterol ester in macrophage plus *M. leprae*.

Since the increased level of ester could be due to reduction in degradation of synthesized ester, the cholesterol ester hydrolytic activity in the infected macrophages was determined. This was done by incorporating into macrophage cultures, $[^{14}C]$ -cholesterol oleate and monitoring the level of hydrolysis in uninfected and infected cultures. The level of hydrolysis is virtually double in uninfected cultures as compared to infected cultures (table 4). This is due to lower level of esterase enzyme in the infected cultures. Using macrophage lysate as a source of the enzyme and $[^{14}C]$ -cholesterol oleate as substrate, the ester hydrolytic activity of infected and control macrophages was determined. The activity was lowered by 50% in the infected macrophages (table 4). Thus the macrophage lysate after proper incubation, with or without *M. leprae* was prepared and used to

Table 4. Hydrolytic activity in the macrophages towards cholesterol oleate added to them with or without M. leprae. Radioactivity (cpm) associated with the compounds as calculated/10⁶ macrophages.

	Cholesterol	ester level in	Oleic acid	(released)	Per cent h	ydrolysis*
Expt. No.	Macrophage	Macrophage + M. leprae	Macrophage	Macrophage + M. leprae	Macrophage	Macrophage+ M. leprae b
0 το 4 το	1844 1086 1020 1360 1082	2040 1700 1810 2192 1999	300 156 165 770 225	156 128 150 382	14 12.5 14 36 17.3	7.1 7.6 14.8 9
* Per cei p value	nt hydrolysis is calcula :-<0.0125 (a-b)	tted as oleic acid released	to total radioactivity (ester + oleic acid)		

Macrophage cholesterol

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(lysate).
extract
nacrophage
n the 1
hydrolytic activity i
Ester
Table 5.

**[]	lacro- nage+ <i>leprae</i> b	2.9 2.2 9
lýsis tot	M.P.	
Hydro	Macro- phage a	16 16.3 14.3 47
	0- :+ rae	
activity	Macr phage M. lep	3.2 2.6 2.8
Specific	facro- shage	5 44.3 4.3
	N 4	
ein (µg)	Macro- phage + M. leprae	205 360 112 250
l'otal prote	acro- iage	30 20 25
	M Iq	1003
liberated 3m)	Macro- phage + <i>M. leprae</i>	665 920 274 660 6 the ester
Olcic acid (as cr	lacro- bhage	1680 2950 956 594 paration of
	N H	TLC se
ester in the mixture *	Macro- phage + M. leprae	39540 40940 18174 9011 mined after
Cholesterol reaction	Macro- phage	37590 23060 14674 6800 cpm as deter
	Expt. No.	- 0 0 4

p value <0.0125 (a-b)

** Since the lysate came from different number of macrophages the hydrolysis is standardised to µg protein and 10⁸ macrophages.

Specific activity: Oleic acid (cpm) liberated/ μ g protein.

Total hydrolysis: Olcic acid (cpm) liberated/µg protein/10⁸ macrophages.

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assay their ability to hydrolyse labelled [14 C]-cholesterol oleate *in vitro*. The level of hydrolysis was determined by the amount of oleic acid released by estimating the amount of radioactivity associated with oleic acid separated in TLC from the incubation mixture (table 5). It is clear from the data that quantum hydrolysis by lysate from *M. leprae* containing macrophage, is extremely low compared to the control lysate. The total level of protein is also lower in the lysate from infected macrophages as compared to the control lysate. Nevertheless if specific activity is determined as cpm oleic acid/unit μ g protein, and cpm oleic acid/ μ g protein/10⁶ macrophages the enzyme level is again indicated as low in infected macrophages.

Discussion

Macrophages from the peritoneal cavity of Swiss albino mice when cultured in vitro show uptake of M. leprae as a characteristic phagocytic feature. Phagocytosis of live bacteria could interfere in macrophage metabolism. Earlier Salgame et al. (1980) have shown that protein synthesis is reduced in the macrophages when M. leprae are present. Present data shows that the lipid metabolism of macrophage are also effected followed bacterial infection resulting specifically in increased cholesterol ester level. This abbreviation in cholesterol metabolism is due to the presence of live *M. leprae* in the macrophage is evident by the fact that heat killed M. leprae has no effect. The increase in ester level could be due to either increased synthesis or decreased degradation. The crude extract of the macrophage with or without M. leprae was assayed for cholesterol ester hydrolytic activity. This showed that the hydrolytic enzyme (esterase) is very low in M. leprae infected macrophage. Interestingly enough such a lowered enzyme level is closely correlated with lowered protein level. Thus we confirm the earlier observation of reduced protein synthesis by Birdi et al. (1979) and the system reported for human macrophage by Salgame et al. (1980). The reduced enzyme activity is probably a reflection of reduced level of protein synthesis, taking place after M. leprae are engulfed by these macrophage. At present we have no data to show the level of cholesterol ester synthetase (ACAT) in these cases, but it appears this enzyme level may not be altered along with lowered protein synthesis. The level of radioactive ester formed when cholesterol is given is not lowered inspite of lower level of cholesterol uptake by M. leprae infected cells. This indicate that the synthetic enzyme level may not play a limiting role. It is also clear from all the experiments that the total uptake of cholesterol by M. leprae infected cells are lower. We do not know whether it is due to permeability changes after phagocytosis or any other factor affecting the uptake.

There are definite evidences for the role of low density lipoprotein (LDL) for facilitating entry of cholesterol into the cells and in our experiments we presume that LDL present in the human serum enables entry of cholesterol into the macrophages. The basic observation of accumulation of cholesterol ester in macrophages in the presence of *M. leprae* has significant biological implication in the tissue macrophages. It is a common observation that in leproma tissues there are macrophage which have *M. leprae* in them attributed to presence of excess lipids espcially esters. Our experiments with mice macrophages show the reason for such foamy macrophages, is due to accumulation of cholesterol esters. Such ester

accumulation being a clear result of *M. leprae* interaction with host cell is also indicated. Confirmation of similar observations with human macrophage is now being done so as to identify the significance of this phenomenon in leprosy infections.

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References

Ambrose, E. J., Khanolkar, S. R. and Chulawalla, R. G. (1978) Lep. India, 50, 131.
Birdi, T. J., Salgame, P. R., Mahadevan, P. R. and Antia, N. H. (1980) Int J. Lep., 48, 178.
Brown, M. S., Ho, Y. K. and Goldstein, J. L. (1980) JBio. Chem., 255, 9344.
Day, A. J. (1967) AdvLipid Res., 5, 185.
Doles, V. P. (1956) J. Clin. Invest., 35, 150.
Kondo, E. and Kanai, K. (1976) J. P. J. Med. Sci. Biol., 29, 123.
Imaeda, T. (1960) Int J. Lep., 28, 22.
Salgame, P. R., Birdi, T. J., Mahadevan, P. R. and Anita, N. H. (1980) Int J.Lep., 48, 172.
Skinsness, P. K. (1970) Int J.Lep., 38, 379.
Vahouny, V. G. and Treadwell, C. R. (1968) Methods of Biochemical Analysis, ed. D. Glick (New York: John Wiley and Sons) 16, 219.
Yamamoto, T., Nishiura, M., Harada, N. and Imaeda, T. (1958) Int J.Lep., 26, 1.

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