Increased Synthesis of
Phospholipid during Phagocytosis

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ABSTRACT Incorporation in vitro of $^{32}$P-labeled lysoleicinith (LPC) or lysophosphatidylethanolamine (LPE) into respectively lecitin (PC) and phosphatidylethanolamine (PE) of rabbit granulocytes and alveolar macrophages was compared in the absence and in the presence of ingestable particles. Maximal synthesis of PC by intact cells occurred at added LPC concentrations of less than 0.05 mmole/liter, i.e., at levels found in plasma. Accumulation of PC-$^{32}$P proceeded linearly for at least 30 min and varied directly with cell concentration. While per cell granulocytes and macrophages converted comparable amounts of medium LPC to cellular PC, per milligram of protein, the granulocytes were approximately four times more active than the much larger macrophages. After 30 min newly synthesized PC-$^{32}$P represented as much as 5% of total granulocyte PC. For macrophages this fraction did not exceed 1%. Addition of polystyrene or zymosan particles to the cell suspension resulted in up to 3-fold stimulation of incorporation of LPC-$^{32}$P or LPE-$^{32}$P into their respective diacyl derivatives. This stimulation did not occur when the cells were homogenized. Breakdown of LPC to water-soluble products during phagocytosis of polystyrene particles was the same as at rest. By use of doubly labeled LPC, the mechanism of PC synthesis by the two cell types has been identified as direct acylation of medium LPC, both at rest and during engulfment. Evidence presented in the case of granulocytes suggests that the increased translocation of medium LPC-$^{32}$P during phagocytosis and its conversion to PC represents net synthesis.

The findings indicate that LPC, a normal constituent of plasma, can serve as substrate in PC synthesis by phagocytic cells. This mechanism of PC synthesis can account for appreciable addition of membrane PC, especially by granulocytes. It is proposed that stimulation of this pathway provides building blocks for increased membrane formation during phagocytosis.

INTRODUCTION

Ingestion of particles by phagocytic cells is accompanied by striking morphologic alterations involving both outer and intracellular membranes (2). These changes have suggested formation of new membrane. However, no convincing evidence for increased (net) synthesis of macromolecular membrane constituents has yet been obtained. Thus, leucine-$^{14}$C incorporation into protein of polymorphonuclear leukocytes is not stimulated during phagocytosis (3), and labeling of phospholipids with acetate-$^{14}$C is only modestly increased (3, 4). Furthermore, labeling of phospholipids with $^{32}$P while enhanced during engulfment of particulate matter is only significantly greater in minor phospholipid species (5, 6).

Recently Lands (7) and Erbland and Marinetti (8, 9) have demonstrated in liver homogenates that lyso compounds can be directly converted to their diacyl derivatives. It has subsequently become apparent that enzymes carrying out these

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synthetic activities occur in numerous mammalian tissues including rabbit polymorphonuclear leukocytes and alveolar macrophages (10–16). These enzymes, when concurring with phospholipases that produce lysocompounds, are considered to contribute greatly to the dynamic metabolism of cellular glycerophosphatides (17).

Lysocompounds such as lysolecithin (LPC) and lysophosphatidylethanolamine (LPE) are normal components of plasma (18–21). The studies of Stein and Stein have shown that a considerable portion of i.v. injected radioactively labeled LPC and LPE can be recovered as lecithin (PC) and phosphatidylethanolamine (PE) in many tissues (12). Plasma lysocompounds may therefore be important precursors of cellular phosphoglycerides.

This report concerns an examination in vitro of utilization of LPC and LPE for synthesis of corresponding diacyl products by two types of phagocytic cells, polymorphonuclear leukocytes and alveolar macrophages, both obtained from rabbits. The results indicate that the conversion of medium monoacylphosphoglycerides to cellular diacyl compounds provides an appreciable source of cell lipid. During phagocytosis this conversion is markedly stimulated.

METHODS

Cells. Polymorphonuclear leukocytes were collected from sterile peritoneal exudates produced in rabbits as described before (22). After a cell count the granulocytes (more than 95% of all cells) were resuspended in Hanks' solution or, in some experiments, in ascitic fluid plus 20% homologous serum, to give a final cell concentration of 2 x 10^6 cells per 0.5 ml of complete incubation mixture. This cell number corresponds to approximately 1 mg of protein. In a number of experiments higher cell concentrations were used.

Alveolar macrophages were obtained from BCG-treated rabbits according to the procedure of Myrvik, Leake, and Oshima (23) as modified by Cohn and Wiener (24). After a cell count the macrophages were sedimented at 100 g for 10 min, the saline used for collection and washing discarded, and the cells resuspended in Hanks' solution to give a final cell concentration of 1 x 10^6 cells per 0.5 ml of incubation mixture, corresponding to approximately 2 mg of protein.

Homogenates of both cell types were prepared in 0.25 M sucrose in a glass homogenizing tube, using a motor-driven teflon pestle. Homogenization of granulocytes was carried out for three periods of 30 sec. For macrophages at least three 1 min periods of homogenization were required for complete disruption as assessed by phase-contrast microscopy.

Radioactive substrates. PC and PE were biosynthetically labeled by incubating rat liver slices with orthophosphate-P as described in detail before (15, 16). The LPC-P and LPE-P were obtained by breakdown of the corresponding P-labeled diacylphosphatides with Crotalus adamanteus phospholipase A as recently reported (14). Radiochemical purity of the P lysocompounds as determined by thin-layer chromatography on silica gel G (see below) was at least 98% for LPC and at least 95% for LPE.

Incubation procedure. Radioisotopically labeled LPC or LPE of known specific radioactivity in chloroform: methanol (2:1) was taken to dryness in a glass tube, and a solution of bovine albumin in phosphate buffer (0.05 mole/liter, pH 7.4) or rabbit serum was added. Homogeneous dispersion of the labeled lysocompound in the protein solution was obtained by vigorous agitation on a Vortex mixer (Scientific Industries Inc., Springfield, Mass.).

The 20 mg of bovine albumin or the 0.1 ml of rabbit serum added to the incubation mixture contained respectively approximately 20 (16) and 10 μmole of LPC, Ascitic fluid contained less than 5 μmole/0.5 ml. No meaningful experiment could be designed to determine whether only part or all of this LPC is available for cellular uptake. Arbitrarily, therefore, only the added LPC (from 15 to 60 μmole) has been included in the determination of medium substrate concentration. This most likely means that all calculations of conversion of LPC to PC and glycerylphosphorycholine (GPC) represent estimates that are too low, especially in experiments at lower added LPC levels. However, this limitation in exact quantification of conversion does not seriously affect the conclusions arrived at.

Incubation mixtures of 0.5 ml contained 0.1 ml of the labeled albumin solution (final albumin concentration 4 g/100 ml) and usually 2 x 10^6 granulocytes or 1 x 10^5 macrophages suspended in Hanks' solution. Other additions were as indicated in the text or in the legends.

To elicit phagocytosis, three types of particles were used: (a) polystyrene latex particles of diameter 1.1 μm; (b) zymosan particles (yeast cell walls); or (c) starch granules. These particles were suspended in normal A,4 International Chemical and Nuclear Corporation, City of Industry, Calif.


6 Armour Pharmaceutical Co., Kankakee, Ill.

7 Dow Chemical Co., Midland, Mich.


9 A generous gift of Dr. Bernard Hofreiter of the Northern Regional Laboratory, U. S. Department of Agriculture, Peoria, Ill.
saline. Unless stated otherwise, 0.1 ml of the polystyrene solution, representing \(3.6 \times 10^6\) particles of diameter 1.1 \(\mu\) or 0.025 ml of a zymosan suspension in normal saline representing \(3.5 \times 10^7\) particles were added to determine the effect of phagocytosis upon metabolism of lysocompounds. Preliminary experiments indicated that dialysis of polystyrene particles before use (6) did not alter the observed effects, and therefore the unmodified suspension was used. Chloroform:methanol (2:1) extracts of polystyrene particles contain only traces of lipid phosphorus, virtually none of which migrates with reference LPC or LPE.

Extracts of \(3.5 \times 10^7\) zymosan particles contain 4 \(\mu\)moles of LPC, 9 of PC, 7 of sphingomyelin, and 5 of PE. As in the case of LPC associated with albumin, the possible contribution of all or part of the LPC extractable from zymosan particles to the LPC available for cellular metabolism has been ignored.

Incubation was carried out at \(37^\circ\)C in a shaking water bath with air as the gas phase. Control samples in each experiment included reaction mixtures without cells, incubated for varying periods of time, and complete mixtures extracted before incubation (0 min).

Radiochemical assays. Conversion of the labeled lysocompounds to diacyl and decylated derivatives was determined as recently described (14–16). Briefly, after the indicated incubation times biochemical activity was stopped by addition of 5 ml of methanol and 5 ml of chloroform (25). Extraction was then allowed to proceed overnight at room temperature. The lipid extracts were not washed with water so that water-soluble breakdown products of the \(^32\)P-labeled substrate were retained. After filtration and removal of the solvents under reduced pressure, the extracts were taken up in a few drops of methanol and transferred to a silica gel G thin-layer plate. The \(^32\)P-labeled substrates and the reaction products were separated in a solvent system consisting of chloroform:methanol:glacial acetic acid: water (110:56:20:10, v/v). The separated compounds were localized by exposure to iodine vapor and by scanning with a survey meter for radioactivity and were identified by comparison with appropriate reference substances. After disappearance of iodine stains, marked areas were scraped off the plate into counting vials for determination of radioactivity by liquid scintillation counting after 12 ml of toluene containing 4.0 g of BBOT \(^11\) per liter was added. Quenching due to silica gel was minimal and closely similar for each fraction since care was taken to scrape off approximately equal amounts of powder. The radioactivity in each fraction was expressed as per cent of the total radioactivity recovered in all fractions and converted to milli- micro moles on the basis of the known amounts of radioactive substrate added. In most experiments the whole incubation mixture was extracted. Where indicated, cells and medium were separately extracted. In these experiments the extracts were quantitatively transferred to the thin-layer plate to permit determination of total radioactivity and extent of conversion of the monoacyl substrate to diacyl and decylated compounds.

The radioactive water-soluble products formed during incubation with LPC-\(^32\)P or LPE-\(^32\)P were identified by established techniques (26, 27) as GPC and GPE. Phosphorus determination on phospholipids fractionated by thin-layer chromatography was carried out directly on the silica gel by the method of Parker and Peterson (28). Protein contents of cell suspensions and homogenates were determined by the method of Lowry, Rosebrough, Farr, and Randall (29).

Assessment of "binding" of LPC to polystyrene and zymosan particles. A small and constant amount of LPC-\(^32\)P and increasing concentrations of unlabeled LPC in albumin solution (20 mg in 0.1 ml) were mixed in conical glass centrifuge tubes, with 0.1 ml of polystyrene particles \(3.6 \times 10^5\) particles of 1.1 \(\mu\) diameter and 0.3 ml of Hanks' solution. The particles were sedimented by centrifugation. The supernatant fluid was removed and transferred to a counting vial. The particles were resuspended in 0.3 ml of Hanks' solution, sedimented again, and the wash transferred to a counting vial. The remaining particles were quantitatively transferred with normal saline to a counting vial. After addition of 10 ml of chloroform:methanol (2:1), the contents of the three vials, representing one test concentration of LPC, was taken to dryness, 12 ml of liquid scintillation mixture in toluene added, and the samples counted.

Similar experiments were carried out with \(2 \times 10^6\) zymosan particles, except that 0.1 ml of rabbit serum was used in place of albumin.

RESULTS

Protection by albumin against membrane-lytic effect of LPC. Recent studies of Switzer and Eder have shown that plasma LPC occurs almost exclusively in association with albumin (20).

The importance of binding of LPC to albumin for the protection of polymorphonuclear leukocytes against the lytic properties of the lysocompound is illustrated in Table I. In the absence of albumin, even low LPC concentrations in the medium produce leakage of K\(^+\) from a suspension of granulocytes. Increasing LPC concentrations result in the appearance of more K\(^+\) into the medium. This evidence of damage to the cell is accompanied by morphologic signs of structural changes. Examination by phase-contrast microscopy revealed swollen cells. With higher LPC concentrations the swelling was more marked, and cell outlines became unclear until actual disintegration of cells was noted. By contrast, in the
TABLE I

<table>
<thead>
<tr>
<th>Lysolecithin</th>
<th>Medium (K⁺)</th>
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<tr>
<td>mmole/liter</td>
<td>mg/100 ml</td>
</tr>
<tr>
<td>0</td>
<td>5.0</td>
</tr>
<tr>
<td>0.2</td>
<td>7.4</td>
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<tr>
<td>0.4</td>
<td>8.0</td>
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<tr>
<td>0.6</td>
<td>8.4</td>
</tr>
<tr>
<td>1.0</td>
<td>9.0</td>
</tr>
</tbody>
</table>

A fresh suspension of granulocytes in Hanks' solution was divided into 10 equal portions each containing 3 x 10⁷ cells. Incubation was carried out at 37°C for 30 min in the presence of the indicated concentrations of lyssolecithin (LPC), with or without 20 mg of bovine serum albumin, in a total volume of 0.5 ml. After 20 min a drop of each suspension was taken for microscopic examination. After 30 min cells and medium were separated by centrifugation. The K⁺ concentrations in the medium were determined by flame photometry.

A fresh suspension of physiologic concentrations of albumin (4 g/100 ml) none of these changes occurred until the LPC level in the medium was 1 mmole/liter. Because of these results all following experiments were carried out with albumin in the incubation mixture.

I ncorporation of medium LPC⁻³²P complexed to albumin into PC of granulocytes and macrophages. While albumin protects against the membrane-lytic properties of LPC, presumably because of complex formation between the two molecular species (16, 20), the apparent binding does not prevent metabolic utilization of LPC. Fig. 1 compares the formation of PC⁻³²P from albumin-bound LPC as a function of time, by intact granulocytes and macrophages and by homogenates of both cell types. PC synthesis per milligram of protein is distinctly greater by intact granulocytes than by intact macrophages. Homogenates of both cell types under these experimental conditions are usually more active than whole cells. Whereas synthesis by intact cells proceeds in linear fashion for at least 30 min, PC formation by homogenates levels off after 10 min.

Fig. 2 shows the effect of increasing LPC concentrations on PC synthesis and further demonstrates the difference in activity between intact cells and homogenates. To fit the results in one graph, the units on the ordinate expressing PC synthesis per milligram of protein have been enlarged 10-fold as compared to Fig. 1. Both intact cell systems manifest maximal activity at low substrate concentrations. Especially at higher substrate levels PC synthesis by homogenates is far greater than by intact cells. This is particularly striking in the case of granulocyte homogenates (15, 16).

While relative to homogenates intact cells manifest modest PC-forming activity, the radioactive PC formed from medium LPC represents as much as 5% of the total PC content of granulocytes (see Table VI) in 30 min. For macrophages, which are less active per milligram of protein than the granulocytes and which contain more PC (16), the contribution of the conversion of exogenous LPC to cellular PC is less than 1% in 30 min.

Fig. 3 shows that PC synthesis increased linearly with increasing numbers of cells.

Varying the pH of the incubation medium between 6.0 and 8.0 had little effect on PC synthesis by either cell type.

E ffect of ingestion of particles upon incorporation of medium LPC into cell PC. Since the preceding results indicated that albumin-bound LPC constitutes an appreciable source of cell PC, especially for granulocytes, we examined the effect of phagocytosis upon utilization of medium LPC as a precursor of cellular PC.

Fig. 4 shows the mean values of at least four paired experiments at each time interval, in which incorporation of medium LPC⁻³²P into PC of granulocytes and macrophages was compared in the absence and presence of polystyrene or zymosan particles. The former are effectively taken up in a medium that contains albumin only (30), whereas engulfment of zymosan requires addition of whole serum (31). Control values for PC synthesis by granulocytes were the same in medium that contained albumin or serum, whereas macrophages formed more PC in the presence of 20% serum. Therefore, results obtained in macrophage experiments with polystyrene and zymosan particles are depicted by separate graphs. It is evident that ³²P-labeling of PC is stimulated by addition of either polystyrene latex or zymosan particles. The results of these experiments are given as millimicromoles of PC synthesized per 2 x 10⁷ cells.
It may be noted that PC synthesis by an equal number of granulocytes and macrophages in the absence of particles is comparable. However, as was apparent from Figs. 1–3, PC-forming activity per milligram of protein by the much larger macrophages is about one-fourth of that by granulocytes. The extent of stimulation of PC labeling by addition of particles in individual experiments with granulocytes ranged from 25 to 100% for polystyrene and from 100 to 300% for zymosan, at time intervals of up to 60 min. Although stimulation by both types of particles was also clearly evident in macrophage experiments, its magnitude was less than seen with granulocytes when zymosan or starch granules (Fig. 5) were used. After 30 min PC synthesis by granulocytes in the

![Figure 1](image1.png)

**Figure 1** Kinetics of lecithin formation by intact and homogenized cells. Assays were carried out at pH 7.4 and at a lysolecithin (LPC) concentration of 0.1 mmole/liter, as described in Methods. Approximately equal amounts of protein were added to each incubation mixture, either as intact cells (in Hanks' solution) or as homogenate (prepared in 0.25 M sucrose). Incubation mixtures containing homogenate were fortified with 2 μmoles of adenosine triphosphate (ATP), 0.1 μmole of coenzyme A (CoA), 2 μmoles of MgCl₂, and 2.5 μmoles of phosphate buffer (pH 7.4) in a total volume of 0.5 ml. Activities of homogenates were the same in phosphate buffer or in Hanks' solution. (Omission of ATP and CoA slightly reduces activity of granulocyte homogenates, but considerably reduces activity of macrophage homogenates) 12 (15).

12 Since optimal synthesis of lecithin (PC) by homogenates required addition of ATP and CoA, the effect of these agents on whole cells was examined. Addition of ATP and CoA in the indicated concentrations does not affect PC formation by intact granulocytes but stimulates intact macrophages by an average of 30% in five paired experiments (unpublished observations). In none of the experiments shown was ATP or CoA added to intact cell preparations.

![Figure 2](image2.png)

**Figure 2** Effect of increasing lysolecithin concentrations on lecithin synthesis by intact and homogenized cells. For experimental conditions see Fig. 1.
absence of particles was 2.8 ± 0.4 μmole (mean ± se) per 2 × 10⁶ cells and 4.0 ± 0.5 μmole in the presence of polystyrene in 18 paired experiments (P < 0.01). These values were 3.1 ± 1.0 and 8.2 ± 1.1 μmole per 2 × 10⁶ cells in five paired experiments with zymosan (P < 0.005). In six macrophage experiments the control values were 2.5 ± 0.3 μmole per 2 × 10⁷ cells with albumin and 3.1 ± 0.5 μmole per 2 × 10⁷ cells with serum in the medium and upon addition of polystyrene or zymosan particles, 3.8 ± 0.5 and 4.75 ± 0.8 μmole per 2 × 10⁷ cells respectively. These differences were not statistically significant (P > 0.05). However, when the values obtained in the presence of particles were expressed as percent of control in each experiment, the P value was < 0.005 for polystyrene particles and < 0.001 for zymosan particles.

The effect of increasing numbers of particles upon stimulation of incorporation of LPC-³²P to PC is shown in Fig. 5. Results are given as percent of control values after incubation for 30 min. Unlike polystyrene particles, the much larger zymosan granules elicited a nearly maximal stimulatory effect at concentrations as low as 1.4 × 10⁸ particles (0.01 ml).

In contrast to the reproducible increase in conversion of medium LPC into cellular PC during phagocytosis, concomitant breakdown of LPC to water-soluble GPC is not enhanced upon addition of polystyrene particles (Fig. 6).

On the other hand incorporation of another monoacylphosphoglyceride, namely LPE-³²P, into its diacyl derivative by both cell types is stimulated during incubation with polystyrene particles (Fig. 7).

Verification of actual phagocytosis of the polystyrene particles was obtained by electron microscopy of thin sections of granulocytes incubated under the usual experimental conditions for 15

2222  P. Elsbach
Intracellular localization of poly styrene particles, surrounded by a membrane-like structure, was readily apparent.\(^{13}\)

Addition of particles to the cell suspension might in some manner change the physicochemical properties of substrate and (or) enzyme to produce the observed stimulation. If this were so, addition of particles may be expected to stimulate PC synthesis by homogenates. Table II shows that, in contrast to the reproducible stimulation of PC synthesis by intact cells, homogenates are unaffected by addition of particles.

Further, homogenates of cells that had ingested particles were no more active than homogenates of resting cells.

\(^{13}\) We are greatly indebted to Dr. D. Zucker-Franklin who kindly performed the electron microscopy.

<table>
<thead>
<tr>
<th>Table II</th>
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<tbody>
<tr>
<td><strong>Effect of Addition of Particles upon Lecithin Synthesis by Intact Granulocytes and Homogenized Cells</strong></td>
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<tr>
<td></td>
</tr>
<tr>
<td>Intact cells</td>
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<tr>
<td></td>
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<tr>
<td>Homogenates</td>
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</tbody>
</table>

Suspensions of intact granulocytes in Hanks’ solution and freshly prepared homogenates were incubated for 30 min with 3.6 × 10^6 polystyrene particles or with 3.5 × 10^7 zymosan particles. Synthesis of lecithin (PC) was assayed as described in Methods. The results are given as per cent of control values in each experiment. The values listed are means ± SE of the indicated number of experiments (n). Homogenization in 0.25 M sucrose or in Hanks’ solution gave the same results.

Studies on the nature of increased incorporation of medium LPC-\(^{32}\)P into cellular PC during engulfment of particles

**Adsorption of LPC to polystyrene and zymosan particles.** If LPC-\(^{32}\)P were adsorbed to the particles, increased labeling of PC might reflect increased entry of LPC along with engulfed particles. Fig. 8a shows that increasing concentrations of added LPC result in a linear increase in the quantity of LPC associated with particles washed free of medium (see Methods). This association is distinctly greater for zymosan than for polystyrene particles. Fig. 8b depicts a similar experiment carried out at one LPC concentration (0.075 mmole/liter) but with a varying number of zymosan particles. It demonstrates a linear increase in “binding” with an increasing number of particles.

The extent of “binding” by the usual number of polystyrene particles (3.6 × 10^6) and of zymosan particles (3.5 × 10^7) introduced to each incubation mixture is less than 2 mM moles at the LPC concentration range used in these experiments (0.03–0.12 mmole/liter). Since only a limited portion of the added particles is engulfed during incubation (32), increased delivery of substrate during phagocytosis through “binding” of the LPC to ingested particles can only account for a fraction of the observed stimulation of PC synthesis (even if it is assumed that the adsorbed LPC is available to cellular enzymes). Moreover, several experiments were carried out with up to 10^9 cells per incubation mixture instead of the usual 2 × 10^7 granulocytes or 1 × 10^7 macrophages. Under these conditions the absolute increase in PC formation in the presence of particles far exceeded the amount adsorbed, since both PC...
synthesis and its stimulation in the presence of particles increased proportionately with increasing cell number.

**Biochemical mechanism of PC formation from medium LPC by granulocytes.** In mammalian tissues LPC can be directly converted to PC in two ways: (a) by acylation of LPC according to the pathway of Lands (7), or (b) by a transfer reaction in which one LPC molecule donates its fatty acid to another LPC (8, 9, 13-15). Current evidence suggests that LPE does not serve as substrate in such a transfer reaction, and that the conversion of LPE to PC occurs only as a result of direct acylation (16). Previous studies in this laboratory have indicated that homogenates of polymorphonuclear leukocytes contain enzymes of both pathways, whereas in macrophage homogenates conclusive evidence was obtained only for the operation of pathway a (14-16).

With a mixture of palmitic acid-1-14C-labeled LPC and LPC-32P, it can be determined which of the two pathways mainly accounts for synthesis of PC. If insertion of an unlabeled fatty acid by acylation represents the mechanism of PC synthesis, the 14C/32P ratio in the newly formed PC and in the LPC substrate should remain the same. On the other hand, in the transfer reaction a fatty acid-1-14C from LPC occupies the free -OH group, resulting in a 14C/32P ratio in PC formed twice that of LPC. Intermediate values for 14C/32P may be expected if both pathways operate simultaneously.

Table III shows the results of such an experiment. The addition of polystyrene particles resulted in the usual stimulation of PC synthesis. The 14C/32P ratio in the PC formed by the granulocytes and in the LPC substrate remained the same, both in the absence and presence of particles. These findings indicate that direct acylation accounts for synthesis, and that the observed stimulation is not due to appreciable additional operation of the transfer reaction.

**Evidence that increased labeling of PC by granulocytes during phagocytosis represents net increase of cellular PC.** Increased labeling of PC by incorporation of medium LPC-32P in the presence of particles need not represent net addition of cellular PC if newly labeled PC is released into the medium, or if increased conversion of LPC-32P to PC is matched by increased degradation of PC. To examine these possibilities, experiments were carried out with both cell types. Since the relative contribution of medium LPC to formation of cellular PC is substantially greater for granulocytes than for macrophages, it could be predicted that results of experiments on granulocytes would be more conclusive. This turned out to be the case, and therefore emphasis will be placed in this section on the data obtained with granulocytes.

Table IV shows that during incubation of granulocytes for 1 hr the accumulating PC-32P remains associated with the cells, both in the absence and presence of polystyrene particles.

### Table III

<table>
<thead>
<tr>
<th>Time (min)</th>
<th>Lechithin formed</th>
<th>14C/32P in lecithin</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td></td>
<td></td>
</tr>
<tr>
<td>30</td>
<td>4.85</td>
<td>1.15</td>
</tr>
<tr>
<td>60</td>
<td>7.4</td>
<td>1.06</td>
</tr>
<tr>
<td>+ Polystyrene</td>
<td>7.2</td>
<td>1.05</td>
</tr>
<tr>
<td>60</td>
<td>15.0</td>
<td>0.96</td>
</tr>
</tbody>
</table>

Suspensions of granulocytes (5 X 10⁷ cells/tube) in Hanks' solution were incubated with a mixture of palmitic acid-1-14C-labeled lysolecithin (LPC) and LPC-32P (total LPC content 150 µmole in 0.5 ml of incubation mixture) in the absence or presence of 3.6 X 10⁹ polystyrene particles. Labeling of LPC with palmitic acid-1-14C and determination of 14C and 32P radioactivity in LPC and lecithin (PC) were carried out as previously described (14, 15).

### Table IV

<table>
<thead>
<tr>
<th>Time (min)</th>
<th>Cells</th>
<th>Medium</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Control</td>
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</tr>
<tr>
<td>10</td>
<td>0.9</td>
<td>2.2</td>
</tr>
<tr>
<td>30</td>
<td>2.8</td>
<td>5.8</td>
</tr>
<tr>
<td>60</td>
<td>6.2</td>
<td>8.5</td>
</tr>
</tbody>
</table>

Granulocytes (2.5 X 10⁷ cells/tube) were incubated with or without polystyrene particles. At the end of the indicated incubation times cells and medium were separated by centrifugation and each extracted (see Methods).
To examine degradation of cellular PC, this phosphatide was labeled during preincubation of granulocytes for 1 hr with LPC-\textsuperscript{32}P of high specific activity (Table V). In this experiment the lipid extracts were washed with water so that compounds such as GPC were not recovered. However, total counts in cells and medium were roughly comparable for all portions, indicating that no appreciable loss had taken place. It is further evident that the radioactivity in cellular PC, expressed as per cent of total counts, does not decrease during reincubation for 1 hr, even in the presence of zymosan particles. In fact, in this and other similar experiments, a slight increase in PC radioactivity was noted in the presence of particles, matched by a small decrease in LPC radioactivity in the medium. By contrast, cellular LPC radioactivity remained about the same with and without particles. These findings indicate that under the experimental conditions employed no detectable degradation of granulocyte PC-\textsuperscript{32}P takes place during 1 hr.

Phosphorus determinations on major phospholipid fractions of granulocytes incubated with and without particles for 0 min or for periods from

<table>
<thead>
<tr>
<th>Reincubation time</th>
<th>Total cpm recovered</th>
<th>Lecithin Cells Medium</th>
<th>Lysolecithin Cells Medium</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{min}</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
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<tr>
<td>30</td>
<td>9950</td>
<td>86.2</td>
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<tr>
<td>30 + zymosan</td>
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<td>60</td>
<td>9625</td>
<td>86.0</td>
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<tr>
<td>60 + zymosan</td>
<td>10,765</td>
<td>89.6</td>
<td>0.7</td>
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</tbody>
</table>

Preincubation of $6 \times 10^6$ granulocytes was carried out for 60 min in 0.5 ml of ascorbic fluid containing 200,000 cpm of LPC-\textsuperscript{32}P representing 100 umoles. At the end of this period the cells were sedimented by centrifugation for 10 min at 100 g; the supernatant radioactive medium was removed and the cell pellet rinsed twice with nonradioactive ascorbic fluid. The cells were then resuspended in ascorbic fluid in a total volume of 1.8 ml. This suspension was equally distributed over six tubes each of which contained 0.2 ml of homologous serum and, where indicated, $1.4 \times 10^6$ zymosan particles in 0.1 ml saline. After the indicated times of reincubation cells and medium were separated by centrifugation and each extracted in chloroform; methanol (2:1, v/v). The extracts were washed twice with 7 mM CaCl\textsubscript{2} and quantitatively transferred to thin-layer plates (see Methods).

### Table V

<table>
<thead>
<tr>
<th>Reincubation</th>
<th>Total cpm recovered</th>
<th>Lecithin Cells Medium</th>
<th>Lysolecithin Cells Medium</th>
</tr>
</thead>
<tbody>
<tr>
<td>min</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>9550</td>
<td>84.7</td>
<td>1.9</td>
</tr>
<tr>
<td>15</td>
<td>9520</td>
<td>85.6</td>
<td>0.9</td>
</tr>
<tr>
<td>30</td>
<td>9950</td>
<td>86.2</td>
<td>1.0</td>
</tr>
<tr>
<td>30 + zymosan</td>
<td>9515</td>
<td>87.2</td>
<td>0.7</td>
</tr>
<tr>
<td>60</td>
<td>9625</td>
<td>86.0</td>
<td>0.9</td>
</tr>
<tr>
<td>60 + zymosan</td>
<td>10,765</td>
<td>89.6</td>
<td>0.7</td>
</tr>
</tbody>
</table>

LPC, lysolecithin; SPH, sphingomyelin; PC, lecithin; PE, phosphatidylethanolamine. Washed lipid extracts of granulocytes were fractionated by thin-layer chromatography, and lipid phosphorus determinations were carried out on the major phospholipid species separated. LPC, SPH, PC, and PE comprised from 70 to 80% of the total lipid phosphorus applied to the plate. No attempt was made to recover other phospholipid species. Results are given as mean ± sd and fall in the same range as those recently reported for homogenates of granulocytes (16).

### Table VI

**Major Phospholipid Fractions of Granulocytes Incubated with and without Particles**

<table>
<thead>
<tr>
<th>Incubation time</th>
<th>LPC</th>
<th>SPH</th>
<th>PC</th>
<th>PE</th>
</tr>
</thead>
<tbody>
<tr>
<td>\textit{min}</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0</td>
<td>7.5 ± 1.5</td>
<td>41.4 ± 3.7</td>
<td>55.5 ± 1.8</td>
<td>31.5 ± 3.0</td>
</tr>
<tr>
<td>15–60</td>
<td>6.0 ± 1.0</td>
<td>40.5 ± 3.1</td>
<td>56.5 ± 1.8</td>
<td>33.0 ± 1.4</td>
</tr>
<tr>
<td>Polystyrene</td>
<td>10.0 ± 1.3</td>
<td>43.8 ± 5.6</td>
<td>56.3 ± 3.0</td>
<td>37.0 ± 2.2</td>
</tr>
<tr>
<td>15–60</td>
<td>8.1 ± 1.2</td>
<td>45.0 ± 4.4</td>
<td>53.7 ± 2.3</td>
<td>38.0 ± 2.2</td>
</tr>
<tr>
<td>Zymosan</td>
<td>8.2 ± 2.9</td>
<td>36.0 ± 2.3</td>
<td>58.0 ± 5.7</td>
<td>32.0 ± 7.5</td>
</tr>
<tr>
<td>15–60</td>
<td>9.0 ± 1.3</td>
<td>34.1 ± 1.4</td>
<td>54.7 ± 2.1</td>
<td>31.7 ± 2.2</td>
</tr>
</tbody>
</table>

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15 to 60 min revealed no major changes\textsuperscript{14} (Table VI).

We conclude from the evidence thus far available that the transfer of LPC-$^{32}$P from medium to cell PC not only represents net translocation of lipid from medium to cells, but also that the stimulation of conversion observed during phagocytosis by granulocytes reflects net addition of PC.

**DISCUSSION**

Previous investigations in this laboratory have demonstrated in homogenates of granulocytes and macrophages the presence of enzymes of a diacylmonoaoylphosphatide cycle (14-16, 33). Such a cycle has been found in a number of mammalian tissues (10, 34) and explains that the fatty acids of glycerolphosphatides have a more rapid turnover than the glycerolphosphate backbone (17). The operation of the cycle implies that lyso-compounds are produced during phospholipid turnover, and that the level of these naturally occurring membrane lytic agents can be regulated not only by their further degradation to GPC by ubiquitous lysolecithinases, but also by their utilization for the resynthesis of important membrane constituents. While the reconversion of monoacylphosphatide, derived from metabolism of native phospholipids, to the diacyl derivatives cannot give rise to net synthesis of phospholipid, exogenous LPC and LPE, if available as precursors for synthesis of cellular PC and PE, do provide a source of new membrane lipids.

The present in vitro studies carried out with intact cells indicate that LPC and LPE complexed to albumin in the incubation medium indeed serve as substrates in the synthesis of appreciable quantities of phospholipid by granulocytes, and, to a lesser extent, by alveolar macrophages.

Lyso-compounds represent a small but significant portion of the circulating phospholipids (18, 19, 24). The virtually exclusive association of LPC with albumin in the plasma has been reported by Switzer and Eder (20). Intravenous administration of isotopically labeled LPC and LPE is followed by rapid removal from the circulation (12), and the studies of Glomset have revealed a mechanism for repletion of LPC by the enzymatic reaction in serum: PC + cholesterol $\rightarrow$ LPC + cholesterol ester (35). This evidence of rapid turnover in the circulation further suggests the importance of lysocompounds in cellular phospholipid and membrane metabolism. It is therefore of particular interest that ingestion of particles is associated with a distinct stimulation of PC and PE formation from their respective lysoderivatives in the medium. These two glycerolphosphatides represent major phospholipids of both granulocytes and macrophages. The nature of the process of phagocytosis appears to require assembly of new membrane for replacement of portions of the outer membrane that served in the formation of phagocytic vacuoles.

Previous studies concerned with examining synthesis of lipid at rest and during phagocytosis have employed a number of radioactively labeled precursors. In experiments with granulocytes increased labeling of lipid during phagocytosis has been found with acetate-$^{14}$C (3, 4), uniformly labeled glucose-$^{14}$C (3), orthophosphate-$^{32}$P (5), and inositol-2-$^3$H (6), but not with glycerolphosphate-$\alpha^{-32}$P (6) or fatty acid-$^{14}$C (6, 36). The characteristics of labeling with acetate-$^{14}$C were consistent with chain elongation of pre-existing fatty acids, rather than indicative of de novo synthesis (4). A recent report further indicates that acetyl-CoA carboxylase is indeed not demonstrable in homogenates of mixed human leukocytes from peripheral blood (37). Thus, maintenance of lipid content and composition in granulocytes must depend chiefly on utilization of exogenous lipid constituents. As shown previously, granulocytes actively incorporate long-chain free fatty acids into both neutral lipid and phospholipid, a finding which suggests operation of the phosphatidic acid pathway (36, 38). This operation is also suggested by incorporation of orthophosphate-$^{32}$P into a wide spectrum of phosphatides (5, 6). However, the increased incorporation during phagocytosis of orthophosphate-$^{32}$P into phospholipids is restricted to minor acidic phospholipid components of granulocytes (5, 6), and its relevance to increased synthesis of membrane lipids seems doubtful.

\textsuperscript{14} We have no explanation for the higher values for PE and the somewhat lower values for sphingomyelin found respectively in the presence of polystyrene particles and zymosan particles, regardless of incubation time. These differences with respect to control values were not statistically significant.
In studies on macrophages moreover, no stimulation has been observed of incorporation of acetate-\(^{14}\)C or orthophosphate-\(^{32}\)P into lipids during engulfment of particles (30).

These essentially negative findings left unexplained where the phagocytizing cell derives its membrane components for the presumed increased membrane synthesis. We would like to raise the intriguing possibility that the enhanced incorporation during phagocytosis of compounds such as LPC into the quantitatively and metabolically more important membrane lipids may account for formation of new membrane. Since the conversion of monoacyl to diacylglycerolphosphatide involves a single biochemical step, the extent of conversion can be reasonably accurately assessed.

Evidence has been presented suggesting that increased incorporation by granulocytes of medium LPC-\(^{32}\)P into PC reflects net addition of phospholipid. This conclusion is based upon the following findings: (a) while increased labeling of cellular PC took place during incubation with particles, no labeled PC appeared in the medium, indicating that newly synthesized PC remains associated with the cells (Table IV). This finding is in accord with previous experiments in which granulocyte lipids were labeled with acetate-\(^{14}\)C (4) or with fatty acid-\(^{14}\)C (36); (b) no demonstrable breakdown of prelabeled PC of granulocytes occurred during reincubation for 1 hr, either in the presence or in the absence of particles (Table V).

While the stimulation of incorporation of medium lysocompounds into PC and PE of macrophages during incubation with particles is comparable to that observed in granulocytes, its significance in terms of net addition of lipid is more difficult to assess. In 30 min PC synthesis from medium LPC usually is less than 1% of the PC content of macrophages, even during phagocytosis. Current, as yet unpublished, studies indicate appreciable turnover of prelabeled PC-\(^{32}\)P in the same period. It cannot be reliably ascertained therefore whether or not the increased labeling of PC represents net addition of PC molecules.

The mechanism of synthesis of diacylglycerolipids from LPC and LPE by intact granulocytes and macrophages is direct acylation. In homogenates of granulocytes, but not of macro-

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changes in the outer membrane, a ready means of adding building blocks for new membrane synthesis.

Further studies to explore these possibilities must await more satisfactory techniques for obtaining reasonable yields of pure outer membrane fractions without loss of enzymatic activity and the preparation of \(^{3}H\)-labeled monoacylglycerolphosphatides of high enough specific activity to allow thin-section radioautography of cells in various stages of functional activity.

ACKNOWLEDGMENTS

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REFERENCES


