The Human Alveolar Macrophage: Isolation, Cultivation in Vitro, and Studies of Morphologic and Functional Characteristics

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ABSTRACT Human alveolar macrophages were lavaged from surgically resected lungs and from lungs of normal subjects. Macrophages that had been purified by glass adherence were maintained in tissue culture for as long as 54 days. After 3-4 wk in vitro they underwent transformation into multinucleated giant cells. These aged cells had more than 30 times the phagocytic capacity that the same group of cells had had after 1 day in vitro.

Phagocytosis of heat-killed Candida albicans was inhibited by iodoacetate, sodium fluoride, potassium cyanide, and low partial pressures of oxygen, suggesting that these cells require both oxidative and glycolytic energy sources for maximal particle ingestion. Alveolar macrophages and monocyte-derived macrophages killed Listeria monocytogenes with similar efficiency, but neutrophils were more efficient than either of the other cell types. Bacterial killing is probably not dependent upon myeloperoxidase in the monocyte-derived macrophage or in the alveolar macrophage since histochemical stains for peroxidase do not stain either cell type. C. albicans blastospores, which are killed by neutrophils and monocytes that contain myeloperoxidase, were not killed by human alveolar macrophages during the 4 hr of observation.

Large cells with supernormal phagocytic capacity were recovered from patients with postobstructive pneumonia and from one patient with recurrent bacterial pneumonia, indicating that macrophage function can be altered in certain disease states.

Human alveolar macrophages are unique human phagocytes in their dependence on an oxygen tension greater than 25 mm Hg for maximal phagocytosis. Carbon dioxide tensions as high as 70 mm Hg did not alter phagocytosis when the pH of the medium was held constant. These data suggest that the increased susceptibility to pneumonia of patients with chronic bronchitis or atelectasis may be in part related to suboptimal phagocytosis by macrophages in areas of the lung with depressed oxygen tension.

INTRODUCTION

The alveolar macrophage of a variety of species has been implicated as the most important mechanism by which the lungs are protected from aerosols of microorganisms (1-4). To our knowledge, however, only one study of the function of human alveolar macrophages has been published (5).

In 1961 Myrvik, Leake, and Fariss (6) developed a method of obtaining rabbit alveolar macrophages by transtracheal lavage after removal of the heart and lungs. Voisin, Guillaume, Van-Moorleghem, and Aerts (7) maintained guinea pig alveolar macrophages in tissue culture for as long as 2 wk in order to study their changing phagocytic and metabolic characteristics. In 1967 Finley, Swenson, Curran, Huber, and Ladman (8) devised a method for washing alveolar macrophages from human subjects, and subsequently Pratt, Finley, Smith, and Ladman (9) reported electron microscopic studies of cells retrieved by this technique. The method, however, was found to be unsuitable for use in patients with abnormal lung function, and the number of cells recovered was too small to allow large-scale studies. We therefore developed a method for retrieving alveolar macrophages from surgically removed human lungs. Alveolar macrophages obtained by this method and by the method of Finley (8) were maintained in tissue culture for prolonged periods. The long-term in vitro survival of cells obtained from the surgical specimens has enabled us to study the metabolic requirements for...
phagocytosis, the capacity of the cells to kill bacteria, and the function of the cells in certain disease states.

**METHODS**

**Cell retrieval.** The age, lung disease (if present) and smoking status of donors of the macrophages are given in Table I. Cells were recovered from surgically removed lungs as follows: a balloon-tipped catheter was placed into the largest available bronchus under aseptic conditions. Hanks' balanced salt solution containing 10% fetal calf serum, 10 units of heparin per ml, 50 units of penicillin per ml, and 50 µg of streptomycin per ml (fetal calf serum-Hanks) was injected repeatedly into the lung with a 50-ml syringe until 50-ml volumes of medium could be recovered by gravity-drainage or by aspiration for each 50 ml injected.

Cells were retrieved from the lungs of awake, normal volunteers by the method of Finley (8). Briefly, a No. 19 Metras catheter was placed into a segmental bronchus under fluoroscopic control. The balloon surrounding the catheter was inflated and the cells were retrieved by lavaging the segment of lung with three 100-ml portions of 0.9% saline. The aspirated solution was added to an equal volume of fetal calf serum-Hanks.

Cells retrieved by either method were centrifuged at room temperature at 125 g for 10 min, suspended in a small volume of fetal calf serum-Hanks, and centrifuged again at the same speed. Viability was assessed by exclusion of 0.5% Trypan Blue dye. The cell pellet was then suspended at 106 viable macrophages per ml in McCoy's 5A medium (Grand Island Biological Co.) with 30% human AB serum, and 1-ml volumes were distributed into Leighton tubes (Bellco Glass, Inc.) containing cover slips. This medium was used in all subsequent experiments. All of the macrophages that would adhere had done so by 4 hr. Cells washed-off after 4 hr failed to adhere to another glass surface and did not phagocitize heat-killed C. albicans in suspension cultures.

**Morphology.** Cells pelleted on glass slides with a cytocentrifuge (Shandon Scientific Co., London) and cells adhering to cover slips were fixed in 100% methanol and stained with Giemsas. Viable cells were observed by phase contrast microscopy, and freshly obtained unfixed cells were stained for peroxidase by the method of Goodpasture as modified by Beacom (10).

Minimum and maximum cell diameters were measured with a micrometer eyepiece, and the cell area was calculated using the formula for area of an ellipse.

Cells viewed with the electron microscope were prepared for examination immediately after retrieval or after they had been allowed to adhere to Melinex plastic cover slips (Imperial Chemical Industries, New York) in Leighton tubes for different periods. Cells from the initial cell suspensions were centrifuged at 250 g for 10 min, fixed at 4°C overnight in 3% distilled glutaraldehyde, buffered to 7.4 with 0.1 M sodium cacodylate, and postfixed with 1% osmium tetroxide. Dehydration, embedding, and sectioning were accomplished by standard methods (11). Sections were examined with a Siemens's Elmiskop 1A electron microscope (Siemens America, Inc, New York). Macrophages adhering to the Melinex plastic film were processed by the method of Finket (12).

**Phagocytosis.** In all phagocytosis experiments, 3 X 106 heat-killed C. albicans cells (or Aspergillus fumigatus spores) were added in 0.1 ml volume to each of two

<table>
<thead>
<tr>
<th>Donors</th>
<th>Non-smokers</th>
<th>Tobacco smokers</th>
<th>Marijuana smokers</th>
</tr>
</thead>
<tbody>
<tr>
<td>Average age, yr</td>
<td>56.0</td>
<td>52.1</td>
<td>25.0</td>
</tr>
<tr>
<td>Cigarettes smoked per day (average)</td>
<td>---</td>
<td>10-60</td>
<td>3-20  (Range)</td>
</tr>
<tr>
<td>Malignant tumor of bronchus</td>
<td>2 (1)*</td>
<td>---</td>
<td>---</td>
</tr>
<tr>
<td>Aspergilloma</td>
<td>---</td>
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</tbody>
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* Number of patients with postobstructive pneumonia.

Leighton tubes containing alveolar macrophages. After incubation at 37°C, the cover slips were retrieved, washed in warm medium, fixed, and stained. The number of C. albicans was counted in 25 cells in each quadrant of each cover slip. The results were expressed as the number of C. albicans taken up by 100 macrophages.

**Inhibition of phagocytosis.** Experiments were carried out on cells within 1 wk after isolation. Sodium iodoacetate (1 X 10-4 M), sodium fluoride (2 X 10-4 M), potassium cyanide (1 X 10-4 M), or 2,4-dinitrophenol (2 X 10-4 M) were each incubated with macrophages for 1 hr. C. albicans blastospores were then added to the control and treated tubes, and the phagocytosis experiment was carried out as described.

The effect of iodoacetate, cyanide, and 2,4-dinitrophenol on the rate of oxygen consumption by the cells was measured with a Gilson Oxygraph (Gilson Medical Electronics, Inc, Middleton, Wis.).

The effect of high carbon dioxide and low or absent oxygen concentration on phagocytosis was determined similarly. The gas and C. albicans were added through needles in the tops of the tubes so that the gas environment of the cells was not disturbed during the experiment. All gases were warmed to 37°C and humidified before they were passed over cells. PH was held constant by adding sodium bicarbonate to the medium. Control tubes were equilibrated with 5% carbon dioxide in air. The gas tension and pH of the fluid bathing the cells were measured after each experiment. Phagocytosis was allowed to proceed for 30 min.

**Bacterial killing experiments.** The ability of alveolar macrophages, polymorphonuclear leukocytes, and monocytes to kill Listeria monocytogenes or Klebsiella pneumoniae type I was tested by a previously described method (13). Bacteria were incubated for approximately 18 hr in trypticase soy broth containing methionine labeled with 35Se, or 3P-labeled sodium phosphate, then washed 7 times to remove free radioactivity. Bacterial concentration was determined by measuring optical density at 620 µm. Approximately 3-7 X 106 organisms were added to each Leighton tube at the
beginning of the experiment. At the end of 90 min the tubes were washed to remove extracellular bacteria and the remaining bacteria were released from the macrophages into the clean medium by sonication. Viability of released bacteria was assessed by a standard pour-plate dilution method and the number of phagocytized bacteria was determined by counting the radioactivity in the medium with a liquid scintillation spectrometer. The results were expressed as the per cent of phagocytosed bacteria killed. Control tubes containing no macrophages were included to assess the efficiency of the wash-out procedure. Specific activity of the bacteria (number of viable bacteria per count per min) was measured in tubes containing bacteria but no macrophages.

When killing of bacteria beyond the 90-min phagocytosis period was measured, an antibiotic to kill the few extracellular bacteria was added after the 90-min wash. At the desired time, the action of the antibiotic was terminated by adding penicillinase (penicillin) or by diluting the antibiotic (streptomycin) to nonbactericidal concentrations.

*Candidal activity.* The ability of the macrophages to kill *C. albicans* was assessed by two methods. In the first method the uptake of methylene blue by phagocytized fungi indicates nonviable organisms (14). The second method assesses morphological changes of intracellular organisms in Giemsa-stained slides (15).

**RESULTS**

*Retrieval procedure.* Lavage of surgical specimens with 1–2 liters of wash solution yielded 0.8 × 10⁶–6.0 × 10⁶ cells, whereas lavage of the lungs of awake volunteers with 300 ml of wash solution yielded 1.0 × 10⁸–8.0 × 10⁹ cells. Pure populations of macrophages could be isolated because other cells in the lavage fluid did not adhere and survive on glass for more than 24 hr.

*Morphology.* Cytocentrifuge preparations were made of cells in suspension soon after they were recovered. From 5 to 35% of the nonerythrocytic cells were peripheral blood leukocytes and the rest were large, mononuclear cells. Three types of large, mononuclear cells could be distinguished morphologically (Fig. 1): for convenience these cells are termed type A, B, and C for this study. Type A cells (Fig. 1A) accounted for 94–98% of the mononuclear cells in the lavage fluid from lung tissue which was not distal to the obstructed bronchus. The cells had a mean diameter of 25 μ, with a range of 10–45 μ. The cytoplasm was stained dark and muddy gray and contained many small, dark blue granules with Giemsa stain but showed no staining with peroxidase stain. The nucleus was oval or irregular and stained dark blue to aqua. Vacuoles were often present, especially at the outer border of the cell. Spontaneous ingestion of autologous red blood cells by viable macrophages was occasionally observed by phase microscopy. Electron micrographs of these cells showed numerous mitochondria and lysosomes and abundant quantities of granular endoplasmic reticulum. Large inclusions were seen, especially in smokers' cells.

About 5% of the mononuclear cells in the lavage fluid were type B. The cells had a mean diameter of 30 μ, with a range of 25–40 μ. The most easily identifiable difference between type A and B cells was seen in the nucleus. The nucleous of the type B cell was red to pink with Giemsa stain and had a nucleus:cytoplasm ratio of about 1:6 or 1:7, whereas the type A cell had a nucleus:cytoplasm ratio closer to 1:3. The cytoplasm of the type B cell contained fewer granules than the type A cell and had large, green-staining inclusion bodies (Fig. 1B).

Type C cells accounted for less than 1% of the mononuclear cells washed from normal lungs but contributed up to 95% of the macrophages washed from the lungs of the patients with postobstructive endogenous lipid pneumonia (Fig. 1C). They were the only variety of alveolar macrophage in which phagocytized intact nucleated cells were observed. Their mean diameter was 40 μ and the cytoplasm was packed with vacuoles. When attached to glass, the vacuoles became less prominent or disappeared completely; the cells then looked like macrophages washed from normal lungs.

The population of cells that adhered to glass appeared to be homogeneous and resembled type A cells. Type B cells either did not adhere to glass or phagocytose, or they did adhere but changed their morphologic characteristics after adherence. Type C macrophages did adhere to glass and phagocytize and probably represented type A cells that had been modified by the environment distal to the obstructed bronchus. Adherent cells transformed into multinucleated, large, vacuolated cells after 3–4 wk in vitro (Fig. 1D). Experiments were performed on cells during the 1st wk after isolation, except for experiments specifically designed to study late-stage cells.

*Phagocytosis.* Cells were maintained in vitro for as long as 54 days. The capacity of the cells to ingest heat-killed *C. albicans* in 15 min increased as the cells aged in vitro (Fig. 2). The increase in phagocytic capacity was proportional to the square root of the area of the cells (Fig. 3) and was therefore directly proportional to length of the cell perimeter. Uptake of *C. albicans* by cells on the 1st day after isolation was also related to...
cell size but rate of change of capacity with measured cell area was greater for these cells than for cells that had been in vitro longer than 1 wk. The number of phagocytized particles did not increase when higher concentrations of C. albicans were added. On the 1st day after isolation the mean uptake of heat-killed C. albicans by alveolar macrophages from nonsmoking subjects was 3.27 C. albicans per cell (±0.6) (Fig. 4). Uptake of C. albicans by macrophages from smokers of marijuana and tobacco cigarettes was not significantly different from the uptake of normal cells (P > 0.1 by Student's "t" test). Alveolar macrophages from four patients phagocytized higher numbers of C. albicans.

The phagocytic capacity of cells from all four patients was outside the 99% tolerance limits of the phagocytic capacity of the macrophages from the other subjects, but the increased uptake was not greater than predicted for their size (Fig. 3). Three of these subjects had "postobstructive" or "lipoid" pneumonia. The fourth subject, a marijuana smoker, had suffered repeated episodes of pneumonia, most likely from aspiration during his frequent semistuporous periods.

When particle uptake was observed at intervals after the addition of phagocytic particles (either heat-killed C. albicans or heat-killed A. fumigatus), the cell capacity to ingest particles rose to a maximum between 30 and 45 min and declined thereafter (Fig. 5). The decline after 45 min was caused in part by particle digestion, as judged by the observation of C. albicans fragments inside large vacuoles, and in part by loss of particle-laden cells from the cover slips, as judged by the decreasing number of cells per cover slip.

Phagocytosis by these cells was partially opsonin dependent. Studies of phagocytosis of A. fumigatus were carried out with normal pooled serum in the culture medium or with serum from two patients with aspergillosis. The serum of both patients contained precipitating antibodies of the IgG class, and one serum also contained skin-sensitizing antibody of the IgE class. (IgG and IgE were demonstrated with specific antisera to each class of antibody.) Phagocytosis of A. fumigatus was greater in 1-day-old cells incubated with either of the two antibody-containing sera (506.6 A. fumigatus/100 cells ±SE 55.5 and 911.4 A. fumigatus/100 cells ±SE 51.1, respectively) than in the cells incubated with the normal serum (162 A. fumigatus per 100 cells ±SE 11.6). These differences were significant at the 0.1% level for both sera. Phagocytosis of C. albicans was not increased by these two sera.

The rate of oxygen consumption was measured before and after the addition of inhibitors or after the addition of C. albicans in two experiments on two different

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**Figure 2** Increase in capacity of alveolar macrophages to ingest C. albicans with increase in cell age in vitro. Cells were incubated with 10^7 heat-killed C. albicans blastospores for 15 min. Increasing the concentration of blastospores did not increase phagocytosis further. The line is the calculated regression line for samples from one individual studied at varied times after isolation. Four similar experiments also showed a linear increase of phagocytosis with cell age. Vertical lines indicate standard errors of the mean.

**Figure 3** Phagocytosis with size. C. albicans uptake and the length of and major and minor axes of cells were measured on the same coverslips. Areas were calculated using the formula for an ellipse. Open squares represent studies on cell populations from two individuals measured from 7 to 54 days after isolation (correlation coefficient r = 0.957). Each closed circle represents a different subject studied on the first day after isolation (correlation coefficient r = 0.879). The upper four points represent cells from patients with postobstructive pneumonia. Lines are calculated regression lines. Points are plotted on a log log scale.
groups of cells. Sodium iodoacetate, $1 \times 10^{-4} \text{ M}$, did not change the rate of resting oxygen consumption of the cells. Potassium cyanide, $1 \times 10^{-4} \text{ M}$, reduced the rate of oxygen consumption to levels too low to be measured by the Oxygraph. At this concentration of cyanide the cells were still able to exclude Trypan Blue dye. The addition of 2,4-dinitrophenol, $2 \times 10^{-4} \text{ M}$, increased the rate of oxygen consumption 4-fold. No increase was observed after addition of *C. albicans*. Phagocytosis was studied after exposure of the cells to metabolic inhibitors before adding the test particles. Partial inhibition of phagocytosis was demonstrated when glycolysis, cytochrome electron transport, or total oxidative metabolism was inhibited (Table II).

The effect of high carbon dioxide and low oxygen tensions on phagocytosis is shown in Table III. Phagocytosis was inhibited by an oxygen tension of 25 mm Hg. No greater inhibition occurred when high CO$_2$ tension was added to the low oxygen tension. No inhibition was noted when the cells were exposed to high carbon dioxide and normal oxygen tension.

**Microbicidal activity.** The killing of *L. monocytogenes* by alveolar macrophages, polymorphonuclear leukocytes, and monocyte-derived macrophages is shown in Fig. 6. Within 1½ hr after the onset of phagocytosis, alveolar macrophages killed 64% of ingested *L. monocytogenes*, whereas polymorphonuclear leukocytes killed 98%. Killing of *L. monocytogenes* by alveolar macrophages was nearly complete within the first 5 hr. Macrophages derived from monocytes resembled alveolar macrophages in their ability to kill *L. monocytogenes*. In similar studies, *K. pneumoniae* were eliminated from all three groups of cells within 1½ hr. Electron micrographs of phagocytized bacteria showed electron-dense granules at the periphery of the phagosomes within the phagosomal membrane. This observation suggests that degradation of lysosomes into phagosomes occurs in alveolar macrophages as it does in other phagocytic cells. In contrast to their ability to kill *L. monocytogenes* and *K. pneumoniae*, we were unable to detect killing of ingested *C. albicans* during periods of observation up to 4 hr.

**DISCUSSION**

Until recently the alveolar macrophage has been inaccessible for study in man. Retrieval of macrophages from awake subjects has three major drawbacks: (a) The number of cells recovered is small; (b) The cells survive poorly in suspension cultures, precluding long-term studies; and (c) Only lungs from subjects with normal pulmonary function can be lavaged safely. We over-

![Figure 4](image-url)  
**Figure 4** Phagocytosis of *C. albicans* by alveolar macrophages from normal (nonsmoking), tobacco smoking, and marijuana smoking subjects, and patients with postobstructive or lipoid pneumonia 24 hr after isolation. Tubes of cells were incubated, in duplicate, with $10^7$ heat-killed *C. albicans* blastospores for 15 min. Vertical broken line indicates range. Vertical solid line indicates standard error. n is the number of subjects.

![Figure 5](image-url)  
**Figure 5** Phagocytosis of *C. albicans* by 4-day-old alveolar macrophages. Maximum particle ingestion at 45 min was followed by a decline. Cells were incubated, in duplicate, with $10^7$ heat-killed *C. albicans* blastospores and tested at the intervals shown. Similar curves were obtained when *A. fumigatus* was used as the test particle or when older or younger cells were tested.
TABLE II
Effects of Metabolic Inhibitors on Phagocytosis by Alveolar Macrophages*

<table>
<thead>
<tr>
<th>Metabolic inhibitor of</th>
<th>Inhibitor (concentration)</th>
<th>Mean per cent inhibition</th>
<th>( P ) *</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glycolysis</td>
<td>Sodium iodoacetate (1 × 10^{-4} M)</td>
<td>50.6</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Glycolysis</td>
<td>Sodium fluoride (2 × 10^{-4} M)</td>
<td>25.7</td>
<td>&lt;0.025</td>
</tr>
<tr>
<td>Cytochrome electron transport</td>
<td>Potassium cyanide (1 × 10^{-4} M)</td>
<td>19.7</td>
<td>&lt;0.025</td>
</tr>
<tr>
<td>Oxidative phosphorylation</td>
<td>Dinitrophenol (2 × 10^{-4} M)</td>
<td>9.1</td>
<td>&gt;0.10</td>
</tr>
<tr>
<td>Oxidative metabolism</td>
<td>Nitrogen (100%)</td>
<td>39.8</td>
<td>&lt;0.02</td>
</tr>
</tbody>
</table>

* Cells incubated with inhibitor for 1 hr before adding \( \text{C. albicans} \). Phagocytosis examined 15 min after adding \( \text{C. albicans} \).
‡ Probability estimated by Student’s “t” test for matched controls.

To determine which metabolic pathways provide energy for phagocytosis, we added metabolic inhibitors to macrophages and found that phagocytosis is partially inhibited by iodoacetate, sodium fluoride, potassium cyanide, and nitrogen atmosphere. Other investigators have published conflicting views regarding the effects of metabolic inhibitors on phagocytosis by animal alveolar macrophages. Oren, Farnham, Kazuhisa, Milofsky, and Karnovsky (16) found a similar pattern of phagocytosis inhibition in alveolar macrophages from guinea pigs. Ouchi, Selvaraj, and Sbarra (17), however, found that only anoxia and concentrations of cyanide that caused the cells to take up trypan blue from the medium (i.e., levels which killed the cells) inhibited phagocytosis by rabbit alveolar macrophages. These investigators further argued that the levels of iodoacetate used by Oren et al. (16) to inhibit glycolysis also inhibited oxidative metabolism. To avoid such criticisms of our experiments, we demonstrated that the concentration of cyanide used in the experiments did not cause the cells to take up trypan blue after 2 hr of exposure and that the levels of iodoacetate used did not change oxygen consumption.

Certain pathologic states were also shown to be associated with abnormal macrophage function. Macrophages from patients with tumors that obstruct a bronchus and from one patient with recurrent bacterial pneumonia had supernormal capacity to ingest particles. We believe that these observations indicate that diseases probably exist in which altered macrophage function may be a part of the disease state.

In our experiments macrophages from smokers had normal phagocytic capacity, and Harris, Swenson, and...
Johnson (5) found no difference in bacterial killing by macrophages from smokers and from nonsmokers. Green and Carolin (18), however, have shown that the ability of rabbit alveolar macrophages to kill bacteria in vitro is inhibited by very high concentrations of cigarette smoke.

After phagocytosis, the second important function of the alveolar macrophage is killing of microorganisms. Our comparative studies of the efficiency with which alveolar macrophages, neutrophils, and monocyte-derived macrophages kill L. monocytogenes and K. pneumoniae showed that alveolar macrophages resemble macrophages derived from monocytes in their ability to kill bacteria. Both cell types fail to stain with histochemical stains for peroxidase. Klebanoff (19) showed that bactericidal activity of neutrophils is partially dependent upon myeloperoxidase. Lehrer, Hanfin, and Cline (20) confirmed that observations by showing that neutrophils from a patient with hereditary myeloperoxidase deficiency had subnormal bacterial killing ability. The microbicidal mechanisms of the human alveolar macrophage are independent of peroxidase but may involve a catalase-dependent mechanism. Gee, Vassallo, Bell, Kaskin, Basford, and Field (21) have shown that catalase-related H2O2 metabolism does occur in rabbit alveolar macrophages.

In order to determine how alveolar macrophages function under conditions that may prevail in diseased lungs, we performed experiments on the effect of the gas environment on phagocytosis. Since an arterial Po2 of about 25 mm Hg is the lowest compatible with brain viability (22), it is reasonable to assume that some alveolar macrophages exist at this oxygen tension, or lower, in patients with severe chronic bronchitis and emphysema or atelectasis. We found that an oxygen tension as high as 25 mm Hg still had a suppressive effect on phagocytosis. Therefore, macrophages probably function suboptimally in areas of the lung in which oxygen tensions are greatly decreased. In contrast, carbon dioxide tensions as high as 70 mm Hg did not affect particle uptake as long as pH was held constant. Unlike other leukocytes, the human alveolar macrophage is therefore highly dependent upon oxidative metabolism for optimal phagocytic activity. These observations suggest that the predisposition of patients with atelectasis (23) or with severe chronic bronchitis and emphysema (24) to pulmonary infection may be related to the suppressive effect of localized hypoxia on alveolar macrophage function. Defense of the lungs by alveolar macrophages is probably most important in these patients because they have impaired bronchial clearance and infected lower respiratory trees.

**ACKNOWLEDGMENTS**

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**REFERENCES**