Intracellular pathogens fall into two broad categories: those that reside within membrane-bound phagosomes and those that escape their phagosomes and reside freely in the host cell cytoplasm. Examples of intracellular pathogens widely accepted as residing within membrane-bound phagosomes are *Legionella pneumophila* (13), *Chlamydia psittaci* (28), and *Coxiella burnetii* (3). Examples of intracellular pathogens that have been reported to lyse the phagosomal membrane and to multiply freely within the host cell cytoplasm include *Trypanosoma cruzi* (19, 23), *Listeria monocytogenes* (9), *Shigella* spp. (6, 20), and some species of *Rickettsia* (27). For *Mycobacterium tuberculosis*, some investigators have reported an exclusively intraphagosomal location (1, 5, 29), whereas other investigators have reported various degrees of intracytoplasmic localization (16, 18). Recently, Teitelbaum et al. (24) have reported that *Mycobacterium bovis* BCG resides in a phagosome that is permeable to relatively low-molecular-weight molecules (fluorescent dextran of 70,000 Da) and smaller (25, 30, 31, 32). We have hypothesized that *M. tuberculosis* resides in an isolated phagosome that is relatively impermeable to cytoplasmic constituents.

Knowledge of whether *Mycobacterium tuberculosis* resides within a relatively impermeable membrane-bound vacuole or is free within the cytoplasm within its host cell is central to an understanding of the immunobiology of this intracellular parasite but is a matter of controversy. To explore this issue, we assessed the accessibility of medium-size protein molecules (Fab fragments of 50,000 Da) to *M. tuberculosis* within human macrophages. We infected the macrophages with wild-type or green fluorescent protein-expressing *M. tuberculosis*, microinjected Fab fragments directed against a major surface antigen of *M. tuberculosis* into the host cell, and assayed the accessibility of the bacteria to the Fab fragments by both immunofluorescence microscopy and immunogold electron microscopy. Whereas microinjected intact immunoglobulin G molecules against cytoplasmic early endosomal antigen 1 readily stained this antigen, microinjected Fab fragments against *M. tuberculosis* did not stain the bacterium within its phagosome. In contrast, microinjected Fab fragments against *Listeria monocytogenes*, an intracellular bacterium known to permeabilize its phagosomal membrane, strongly stained this bacterium. Our study shows that *M. tuberculosis* resides in an isolated phagosome that is relatively impermeable to cytoplasmic constituents.
described previously (5). Polyclonal rabbit anti-L. monocytogenes antiserum was prepared from rabbits injected with formalin-killed L. monocytogenes strain 10403S cells three times 3 weeks apart. The first immunization was performed with complete Freund’s adjuvant (Sigma Chemical Co.), and subsequent immu- nizations were performed with incomplete Freund’s adjuvant. Anti-LAM IgG and anti-L. monocytogenes IgG were purified from the immune rabbit sera by protein A chromatography and enzymatically digested with papain (11). FC fragments and intact IgG were removed by protein A chromatography, and the Fab fragments were further purified by Superdex 75 gel filtration (Pharmacia Fine Chemicals, Inc.). Texas Red-D was covalently conjugated to the purified Fab fragments with the Texas Red-X coupling kit (Molecular Probes) according to the manufacturer’s directions. A rabbit polyclonal antibody to Texas red was purchased from Molecular Probes. The anticytomegaloviral reactivity of this com- mercial antisera was completely removed by five consecutive overnight incu- bations with acetone-treated M. tuberculosis bacterial pellets (10.1 [vol/wt vol]). This and all other antisera were cleared of aggregates by ultracentrifugation (100,000 × g for 1 h) and filtration (0.2-μm-pore-size filter) prior to use. Protein A-coated gold conjugates (5, 10, and 15 nm) were provided by G. Posthuma (Utrecht University, Utrecht, The Netherlands). Texas red-conjugated goat anti- rabbit IgG, Oregon green-conjugated goat anti-mouse IgG, and Alexa fluor 350-conjugated goat anti-mouse IgG were purchased from Molecular Probes. Rhodamine-conjugated goat Fab antibody fragments directed against rabbit IgG were purchased from ICN Pharmaceuticals. All animal studies were approved by the University of California, Los Angeles (UCLA) Institutional Review Board.

Bacteria. M. tuberculosis Erdman strain (ATCC 35801), a highly virulent strain, was obtained from the American Type Culture Collection (Manassas, Va.). The organism was passaged through guinea pig lung to maintain virulence, strain, was obtained from the American Type Culture Collection (Manassas, Va.). The organism was passaged through guinea pig lung to maintain virulence,

Preparation of bacteria for infection of phagocytic cells. Preparations of bacteria were grown in 10% FBS and 10% autologous serum. Twelve hours later, the cells were infected with 10.1 [vol/wt vol] Texas red-conjugated anti-LAM Fab (2 mg/ml) in PBS at 37°C. The suspension was electroporated at 750 V and 25 μF, incubated on ice for 15 min, washed three times to remove extracellular Texas red-conjugated Fab, and incubated for 8 h with RPMI 1640 containing 10% HI FBS and 5% autologous human serum (monocytes) or AB serum (THP-1 cells). The cells on the Cytox-1 beads were incubated with GFP<sub>UV</sub>-M. tuberculosis at an MOI of 30:1 for 30 min and washed three times to remove nonengaged bacteria, and the suspension was transferred to 1-cm-diameter wells containing glass coverslips pretreated with poly-L-lysine (0.1 mg/ml; Sigma Chemical Company). Many of the cells moved from the Cytox-1 to the poly-L-lysine-coated coverslips, facilitating subsequent microscopic visualization. At 1 day postinfection, the cells were fixed in 4% paraformaldehyde–0.1 M sodium phosphate, pH 7.4, containing 6% sucrose, washed with PBS, mounted, and viewed by epifluorescence microscopy as described above.

In a control experiment, 0.2 ml of THP-1 cells on Cytox-1 beads was mixed with 0.2 ml of 2-ng/ml Texas red-conjugated anti-LAM Fab and electroporated as described above. The suspension was incubated on ice for 15 min, washed, and cultured in RPMI 1640 containing 10% FBS and 5% AB serum at 37°C in air containing 5% CO₂. Two days later, the beads were washed in PBS, transferred to a conical 15-ml tube, and sonicated with a 2-mm probe tip at setting 5 with a 50% work cycle with continuous cooling on ice for three 5-min periods (with 5-min periods of cooling between the periods of sonication) in 1.0 ml containing 50 mM HEPES, 0.15 M NaCl, 0.1% bovine serum albumin (BSA), 0.01% fish skin gelatin, 10 μg of soy bean trypsin inhibitor/ml, 10 μg of pepstatin/μl, 10 μg of leupeptin/ml, and 0.05% sodium azide. Particular material of the cell lysate was sedimented by ultracentrifugation (100,000 × g for 1 h), and the supernatant was concentrated to 0.2 ml with a Centriplus 20-concentrator (PL-10; Mil- lipore). The capacity of the Texas red-Fab to bind to M. tuberculosis in the final supernatant was determined (Wako Chemicals, Richmond, Va.).

Microinjection and immunofluorescence microscopy. Monolayers of differentiated THP-1 cells or human monocytes on CELLlocate glass coverslips were incubated with 30 μg/ml of GFP<sub>UV</sub>-M. tuberculosis at an MOI of 30:1 in RPMI 1640 containing 10% fresh AB serum (THP-1 cells) or autologous serum (monocytes) and 10% HI FBS. Monolayers were washed with culture medium, incubated in fresh medium at 37°C for an additional 1 to 3 days, fixed in 0.4% paraformaldehyde in 0.1 M Pipes, pH 7.4-6.4% sucrose (Pipes-sucrose) for 30 min, washed twice in Pipes-sucrose buffer, and incubated for 10 min at room temperature in Pipes-sucrose containing 10 mM glycine. Extracellular M. tu- bercolosis cells were stained by incubation with 30 min with CS35 mouse anti-LAM (10 μg/ml) in Pipes-sucrose-containing 1% BSA and 5% goat serum, washed, and incubated with goat anti-mouse Alexa 350 (1:50; Molecular Probes) in the same buffer for 30 min. Coverslips were washed with Pipes-sucrose and micro-injected with mouse anti-EEA1 IgG (0.2 mg/ml; Transduction Laboratories) and rabbit anti-LAM Fab (0.2 mg/ml; either unconjugated or conjugated to Texas red) in PBS. Microinjections were performed with Femtotip micropipetted needle (Eppendorf/Brinkmann Instruments) and a Microinjector 5252 controlled by a Micromanipulator 5171 (Eppendorf/Brinkmann Instruments) and moni- tored with an Axiovert 135 fluorescence microscope (Zeiss). The cells were allowed to incubate at 4°C for the last 10 min after microinjection. There was an additional interaction between the intracellular antigen and microinjected antibody. The microinjected cells were then fixed for 30 min in 2% paraformalde- hyde in 0.1 M Pipes, pH 7.4-6.4% sucrose, permeabilized in acetone at −20°C for 30 min, air dried completely, rehydrated by being swirled briefly in three-con-
seekers containing 50 ml of PBS, blocked with 5% goat serum–1% BSA in PBS for 30 min at room temperature, and incubated with a combination of Oregon green-conjugated goat anti-mouse IgG and Texas red-conjugated goat anti-rabbit IgG (both diluted 1:500) in 5% goat serum–1% BSA in PBS. Cells were washed, postfixed in 2% paraformaldehyde, mounted, and viewed by epifluorescence microscopy as described above.

As a positive control, human peripheral blood monocytes or THP-1 cells were infected with GFPUV-M. tuberculosis and fixed and permeabilized with acetone as described above, air dried, rehydrated in PBS, blocked with 5% goat serum–1% BSA in PBS, incubated with the antibody mixture used for microinjection diluted 50-fold, and finally incubated with Oregon green-conjugated goat anti-mouse IgG and Texas red-conjugated goat anti-rabbit IgG. In these experiments, we found that the microinjected antibody solutions retained their capacity to stain M. tuberculosis and EEA1 in the permeabilized cells when diluted 50-fold (i.e., final Fab and IgG concentrations of 4 μg/ml).

Immunoelectron microscopy. Plastic coverslips were scratched with a sterile 25-gauge needle to facilitate subsequent orientation in microinjection, embedding, and thin sectioning of a designated area of the monolayers. Human peripheral blood monocytes or THP-1 cells were cultured on plastic coverslips as described above for 2 days in RPMI 1640 with 10% autologous human serum or 10% FBS containing 0.16 nM PMA prior to infection with M. tuberculosis or L. monocytogenes at an MOI of 30:1 for 30 min at 37°C. The monolayers were washed three times with RPMI 1640 and grown for an additional 1 day (M. tuberculosis) or 3 to 4 h (L. monocytogenes) in RPMI 1640 with 10% FBS. The monolayers were fixed with 0.4% paraformaldehyde in 0.1 M PIPES, pH 7.40, containing 6% sucrose for 30 min at room temperature, washed with the same buffer, and microinjected with rabbit anti-LAM Fab-Texas red (2 mg/ml) or rabbit anti-L. monocytogenes Fab-Texas red (2 mg/ml) (1:50) in 5% goat serum in PBS. After 1 h at 4°C, the cells were fixed with 0.5% glutaraldehyde–2% paraformaldehyde in 0.1 M PIPES, pH 7.4, containing 6% sucrose for 2 h at 4°C, washed with PBS, and dehydrated in a graded series of ethanol (70, 80, 90, 95, and 100%). The coverslips were immersed for 1 min in propylene oxide to denature and immobilize proteins, rinsed briefly in 100% ethanol, and embedded in medium-grade LR White embedding resin (Polysciences), and ultrathin sections were collected on Formvar-coated nickel grids. Nonspecific antibody sites on the thin sections were blocked by incubating the grids for 30 min at room temperature on drops of 1% BSA–0.1% fish skin gelatin in 50 mM HEPES buffer, pH 7.4, containing 0.3 M NaCl and 0.05% sodium azide (blocking buffer). Immungold double labeling was performed by a modification of the sequential protein A-gold (PAG) technique as described by Slot et al. (22). Sections were incubated for 1 h at room temperature with a rabbit anti-Texas red antibody (Molecular Probes) diluted 1:500 in blocking buffer, washed on seven consecutive drops of 50 mM HEPES buffer, pH 7.4, containing 0.3 M NaCl and 0.05% sodium azide (washing buffer), and stained with 15 nm PAG in blocking buffer. Free protein A and Fc sites are destroyed by incubating the sections with 2% glutaraldehyde for 10 min. Aldehydes were quenched with 10 mM glycine in wash buffer, and the sections were incubated with rabbit anti-LAM or rabbit anti-L. monocytogenes followed by staining with 5 nm PAG. Sections were stabilized with 2% glutaraldehyde for 10 min, washed in water, and stained with saturated uranyl acetate. Consecutive phagosomes within microinjected cells were photomicrographed with a JEOL 100 CX II electron microscope. Measurements of gold particles per unit area were made with a Nano NIC 2110 digitizer tablet and SigmaScan software (Jandel Science Co.).

RESULTS

M. tuberculosis is not accessible to Fab fragments electroporated into live macrophages prior to infection. To determine whether M. tuberculosis permeabilizes its phagosome to molecules of 50,000 Da, we prepared rabbit anti-mycolobacterial LAM Fab fragments and covalently coupled these to the fluorescent probe, Texas Red-X (anti-LAM Fab-Texas red). We electroporated the anti-LAM Fab-Texas red into the human THP-1 macrophage-like cell line and human peripheral blood monocytes and, 8 to 12 h later, infected the cells with GFPUV-M. tuberculosis. One day later, the macrophages were fixed and examined by fluorescence microscopy to determine whether the intracellular M. tuberculosis cells were stained by the intracellular Texas red-conjugated anti-LAM Fab antibody fragments. Of 200 consecutive GFPUV-M. tuberculosis cells within Texas red-positive macrophages, none of the M. tuberculosis cells stained positive for Texas red. This result indicated that intracellular M. tuberculosis was inaccessible to the Fab fragments. However, it remained possible that the intracytoplasmic Texas red-Fab was degraded within the macrophages.

To explore this possibility, we determined whether the Texas red-Fab retained anti-LAM activity 2 days after electroporation into the macrophages. Macrophages adherent to CytoFLEX-1 beads and containing the probe were lysed by sonication, and the supernatant fluid was examined for its capacity to stain GFPUV-M. tuberculosis. Immunofluorescence microscopy and immunoelectron microscopy confirmed that the antibody retained its activity (data not shown). Thus, intracellular M. tuberculosis is inaccessible to Fab fragments electroporated into live macrophages prior to infection.

M. tuberculosis is not accessible to Fab fragments microinjected into macrophages after infection as assessed by immunofluorescence microscopy. We considered the possibility that the intracytoplasmic anti-LAM Fab-Texas Red might alter the capacity of M. tuberculosis to permeabilize its phagosome and the possibility that M. tuberculosis might permeabilize its phagosome at times later than 1 day after infection. To explore these possibilities, we examined the capacity of anti-LAM Fab to bind to M. tuberculosis when microinjected 1 to 3 days after infection. Human macrophage-like THP-1 cells and peripheral blood-derived monocytes were allowed to adhere to glass coverslips, infected with GFPUV-M. tuberculosis, fixed gently with 0.4% paraformaldehyde, and microinjected with unconjugated or Texas red-conjugated anti-LAM Fab antibody fragments. As a positive control, mouse IgG directed against EEA1 was microinjected simultaneously with the anti-LAM Fab. After fixation and permeabilization, the microinjected antibodies were immunolocalized with a goat anti-rabbit and a goat anti-mouse IgG-conjugated secondary antibodies. For each of days 1, 2, and 3 following infection, we examined over 100 consecutive M. tuberculosis-infected microinjected cells and found none that stained positive for Texas red. We observed intense staining of the cytoplasmically exposed EEA1 by the microinjected mouse antibody (Fig. 2a2, a3, and b2), despite the absence of staining of the M. tuberculosis cells by the rabbit Fab (Fig. 3 and b3). These results were obtained regardless of whether the second antibody used to immunolocalize the rabbit anti-LAM Fab was an intact goat IgG molecule or a Fab fragment. Thus, although the microinjected antibodies were capable of diffusing throughout the cells and binding to cytoplasmically exposed EEA1, the intracytoplasmically injected anti-LAM antibodies were unable to stain M. tuberculosis. These results indicate that the phagosome is not permeable to Fab molecules, which are 50,000 Da in size. In control experiments using cells fixed and permeabilized prior to incubation with antibodies, both EEA1 and mycobacterial LAM were stained by the microinjected antibody solution, even when diluted 50-fold (Fig. 1d1 to d3 and e1 to e3). Microinjection generally results in approximately a 10-fold dilution of the microinjected solution within the microinjected cells.

We considered the possibility that permeabilization of the cells with acetone at −20°C prior to staining with the secondary antibodies might extract LAM and its associated Fab frag-
Microinjected Fab fragments detect EEA1, but not GFP UV-M. tuberculosis, in human macrophages. Monolayers of THP-1 cells and human peripheral blood monocytes were infected with GFP UV-M. tuberculosis and fixed 1 or 3 days after infection, as indicated. (a and b) THP-1 cells (a) and monocytes (b) were microinjected with rabbit anti-LAM Fab fragments and mouse anti-EEA1 IgG, postfixed for 30 min with 2% paraformaldehyde, permeabilized with acetone, air dried, rehydrated in a large volume of PBS, blocked with 57% goat serum–1% BSA in PBS, and stained with Texas red-conjugated goat anti-rabbit IgG and Oregon green-conjugated goat anti-mouse IgG in PBS with 57% goat serum–1% BSA. Coverslips were washed, mounted in Molecular Probes antifade mounting medium, and viewed by fluorescence microscopy. White and black arrows indicate the corresponding positions in micrographs within the same column. The GFP UV-M. tuberculosis cells are visualized by excitation with either violet (a1 and b1) or fluorescein (a2 and b2) filters. The microinjected cells show abundant Oregon green staining of EEA1 (a2 and b2) but no staining of M. tuberculosis by the microinjected anti-LAM Fab (a3 and b3). (c) Monocytes were fixed 3 days after infection with GFP UV-M. tuberculosis and microinjected with rabbit anti-LAM Fab and mouse anti-EEA1 IgG as in panels a and b. The monocytes were then microinjected a second time with rhodamine-conjugated goat Fab directed against rabbit IgG and Oregon green-conjugated goat anti-mouse IgG. Again, EEA1 stains well in the microinjected cells (c2) but the GFP UV-M. tuberculosis cells are not stained by the microinjected anti-LAM Fab (c3). (d and e) As positive controls, monocytes were fixed and permeabilized 3 days after infection with GFP UV-M. tuberculosis, incubated with a 1:50 dilution of the rabbit anti-LAM Fab and mouse anti-EEA1 IgG solution used for microinjection, washed, and stained with secondary antibodies as for panels a and b. In panels c, to confirm the stability of the staining of LAM by anti-LAM Fab fragments, after incubation with primary antibodies, we postfixed the monolayers for 30 min and then subjected them to a second treatment with acetone at −20°C, air dried them, rehydrated them, and washed and incubated them with secondary antibodies (as was done after microinjection of primary antibodies in panels a and b). In these acetone-permeabilized control cells, both LAM on the M. tuberculosis cell wall (e3) and EEA1 (e2) are readily stained. Magnification, ×157.

Microinjection of Texas red-conjugated primary anti-LAM Fab fragments eliminates the need for secondary antibody staining and for acetone permeabilization. When we lightly fixed GFP UV-M. tuberculosis-infected cells 1 to 3 days postinfection and microinjected them with 0.2 mg of 2-mg/ml Texas red-conjugated anti-LAM Fab, we still observed no Texas red staining of GFP UV-M. tuberculosis, despite diffuse cytoplasmic Texas red staining of the microinjected host cells (data not shown). In additional experiments, we lightly fixed human peripheral blood monocytes 3 days after infection with GFP UV-M. tuberculosis and microinjected the cells with a rabbit anti-LAM Fab (0.37 mg/ml) together with mouse anti-EEA1 (0.2 mg/ml), followed by a second microinjection into the same cells with rhodamine-conjugated goat Fab fragments directed against rabbit IgG and Oregon green-conjugated goat anti-mouse IgG. These doubly injected cells (which had no exposure to acetone) showed intense endosomal staining of EEA1 but had no rhodamine staining of GFP UV-M. tuberculosis (Fig. 1c1 to c3).

In these experiments, mouse anti-EEA1 Oregon green fluorescence was employed to identify which cells are microinjected and to demonstrate that the microinjected antibody is able to move throughout the cell and bind to cytoplasmically exposed antigens. The GFP UV fluorescence can be distin-
guished from the Oregon green fluorescence because the former, but not the latter, fluoresces under UV excitation (Fig. 1, top row). However, since both GFP<sub>UV</sub> and Oregon green fluorescence green under visible-blue-light excitation (Fig. 1, middle row), the images are not intended to assess the degree of colocalization of EEA1 and <i>M. tuberculosis</i>.

As a positive control, we permeabilized infected macrophages with acetone before applying antibodies and then stained them with a 50-fold dilution of the primary and secondary antibody solutions removed from the microinjection needles. Under this condition, both EEA1 and <i>M. tuberculosis</i> were positively stained (Fig. 1d1 to d3). In additional control experiments, exposure of acetone-fixed cells to a second round of −20°C acetone treatment had no discernible effect on the fluorescent staining of GFP<sub>UV</sub>−<i>M. tuberculosis</i> or EEA1 (Fig. 1e1 to e3). This result is consistent with the fact that, in the purification of LAM, the addition of −20°C acetone is routinely used to precipitate LAM after it has been solubilized by Triton X-100 detergent (4). In parallel experiments, we also microinjected rabbit anti-LAM IgG molecules into lightly fixed GFP<sub>UV</sub>−<i>M. tuberculosis</i> cells and found no staining of the <i>M. tuberculosis</i> by the anti-LAM IgG (data not shown).

<i>L. monocytogenes</i> but not <i>M. tuberculosis</i> is accessible to microinjected Fab fragments, as assessed by immunoelectron microscopy. To examine at a higher resolution the interaction of microinjected anti-LAM Fab antibody fragments with <i>M. tuberculosis</i>-infected cells, we gently fixed monolayers of human peripheral blood monocytes 3 days after infection with <i>M. tuberculosis</i>, microinjected the cells with a high concentration (2 mg/ml) of Texas red-conjugated anti-LAM Fab fragments, and examined the distribution of the Fab fragments by electron microscopy (Fig. 2A). We quantitated the number of immunogold particles per square micrometer of macrophage cytoplasm and per square micrometer of bacteria and found that the immunogold particles were excluded from <i>M. tuberculosis</i> (Fig. 3). Nonmicroinjected cells and <i>M. tuberculosis</i> cells within nonmicroinjected cells had negligible levels of immunogold staining. None of the <i>M. tuberculosis</i> cells in over 40 consecutively microinjected macrophages in each of two separate experiments showed appreciable staining by the anti-LAM Fab fragments: 160 of 166 <i>M. tuberculosis</i> cells had no immunogold particles; 5 bacteria had a single immunogold particle, and only 1 of the <i>M. tuberculosis</i> cells had two immunogold particles. This level of staining is comparable to the background level of immunogold staining of <i>M. tuberculosis</i> in macrophages that are not microinjected (Fig. 3). As a positive control, we found that even a 200-fold dilution of the Texas red-conjugated anti-LAM Fab (10 µg/ml) yielded intense staining of mycobacteria in thin sections (data not shown). As an additional control, we found that microinjected Texas red-conjugated Fab fragments directed against <i>L. monocytogenes</i> do stain <i>L. monocytogenes</i> in the cytoplasm of host macrophages (Fig. 2B). Of 64 consecutively <i>L. monocytogenes</i> cells in microinjected cells, 59% had more than two immunogold particles. In sharp contrast to the case with <i>M. tuberculosis</i>, we found that the anti-<i>L. monocytogenes</i> immunogold particles were relatively concentrated on the bacteria compared with their concentration in the host cell cytoplasm (Fig. 3).

**DISCUSSION**

In summary, we have found that <i>M. tuberculosis</i> within human cells is inaccessible to microinjected Fab fragments. This indicates that the <i>M. tuberculosis</i> phagosome is not permeable to proteins with a size of 50,000 Da or greater. Our results differ from those of Teitelbaum et al. (24), who studied the related but avirulent mycobacterium <i>M. bovis</i> BCG and reported that approximately 30% of phagosomes containing GFP-expressing <i>M. bovis</i> cells in mouse macrophages were permeable to fluorescent dextrans of less than 75,000 Da. Aside from the different species studied, methodological differences may account for the different observations. The fluorescent-dextran technique used by Teitelbaum et al. relied on a determination of the color of the untargeted fluorescent dextran probe in the vicinity of the phagosome. An erroneous result might occur in the case of very tight-fitting phagosomes. It has been observed that <i>M. tuberculosis</i> and <i>Mycobacterium avium</i> phagosomes can be extremely tight (7, 29). Microinjected fluorescent dextrans can be expected to move freely within the cytoplasm and to surround these tight phagosomes. The resolving power of fluorescence microscopy, even confocal microscopy, is limited by the wavelength of light used to detect the fluorescent probes. Judging from published electron micrographs of tight-fitting <i>M. tuberculosis</i> phagosomes, the distance between the mycobacterium and the host cell cytoplasm (50 to 75 nm) may be considerably less than the wavelength of the light used to detect the fluorescent probes (450 nm). This would result in color mixing and potentially misleading results. Further loss in resolution of the fluorescent-dextran technique may occur if the mycobacteria are moving within the microinjected cell. The combined mobility of the fluorescent dye and the bacterial phagosome can be expected to increase the measured degree of color mixing. Note also that the use of fluorescent beads immobilized on a polyclonal antibody coated microscope slide (as described by Teitelbaum et al.) would not define the resolution that is achieved when observing GFP-expressing <i>M. bovis</i> BCG within living macrophages. This is because the beads are stationary, whereas the living specimen is not, and because light is scattered at multiple interfaces of differing refractive indices that are present in the living cell but that are absent from a microscope slide with adherent fluorescent beads. Differences in refractive index occur at the following interfaces: extracellular medium-plasma membrane, plasma membrane-cytosol, cytosol-phagosomal membrane, phagosomal membrane-phagosomal space, phagosomal space-mycobacterial cell wall, and mycobacterial cell wall-GFP-containing bacterial cytoplasm. Refractile structures or organelles between the objective and the plane of focus, for example, granules or lipid globules, can also scatter light, further degrading resolution. Such refractile structures and interfaces are absent when the microscope resolving power is assessed with stationary fluorescent microspheres.

Furthermore, the different results obtained with different sizes of dextran probe (24) could reflect the different viscosities of the probes injected. Smaller probes are less viscous and greater volumes are likely to be injected into the host cell cytoplasm. This could result in a greater fluorescence signal, a greater osmotic shift, and greater osmotic shrinkage of the phagosome. Smaller volumes of more-viscous higher-molecu-
FIG. 2. *L. monocytogenes* but not *M. tuberculosis* is accessible to cytoplasmically injected Fab antibody fragments as assessed by immunoelectron microscopy. Human peripheral blood mononuclear cells were infected with *M. tuberculosis* (A) or *L. monocytogenes* (B) and fixed after 3 days (A) or 4 h (B). Cells of the monolayer were microinjected with Texas red-conjugated-rabbit anti-LAM Fab or Texas red-conjugated rabbit anti-*L. monocytogenes* Fab. The monolayers were fixed, dehydrated, embedded in LR White resin, and thin sectioned, and the sections were incubated sequentially with rabbit anti-Texas red and 15-nm PAG to localize the microinjected Texas red-Fab conjugates. The sections were blocked and incubated with rabbit anti-LAM IgG or rabbit anti-*L. monocytogenes* and 5-nm PAG. (A) In the *M. tuberculosis*-infected macrophages, the cytoplasm of the microinjected macrophage has copious Texas red-conjugated anti-LAM Fab (15 nm gold; arrowheads) but there is no 15-nm gold associated with the *M. tuberculosis*. The *M. tuberculosis* cells are well stained postsectioning by the 5-nm anti-LAM immunogold (arrows). (B) In the *L. monocytogenes*-infected macrophages, the 15-nm gold (arrowheads) is abundantly associated with the *L. monocytogenes* bacteria (identified postsectioning with 5-nm gold; arrows). Magnification (A and B), ×22,400.
lar-weight dextran probes might be injected into the host cell cytoplasm, resulting in a smaller fluorescence signal and less osmotic shrinkage of the phagosome.

Another potential problem with the use of confocal imaging to assess the integrity of phagosomal membranes in living cells is that the intense laser light of a confocal microscope can cause photo-oxidation and can be toxic to the specimen. Excitation of a fluorophore inevitably produces nonradiative transitions of the fluorophore to the ground state with generation of reactive products, including singlet oxygen, that can oxidize membrane lipids and cause membranes to become permeable (17, 26). Indeed, this principle has been employed deliberately to destroy organelles and to permeabilize cells.

Alternatively, it is possible that the differences between our results and those of Teitelbaum et al. (24) reflect the properties of a putative pore. It is possible that a pore such as that in the M. tuberculosis phagosome is selectively permeable to uncharged dextrans but not to ionically charged Fab fragments or that the formaldehyde fixation employed in our microinjection experiments blocked a channel in such a pore. We believe that this is unlikely, considering the relative dimensions of the 30-Da formaldehyde molecule (4 Å) versus the dimensions of a channel that would be needed to allow passage of a 50,000-Da Fab fragment (40 Å wide, 60 Å long).

Our experiments and those of Teitelbaum et al. (24) have examined the movement of antigens from the cytosol into the phagosome. These experiments do not directly address the movement of mycobacterial antigens from the phagosome into the cytosol, the direction of antigenic flow that would be relevant to the MHC-I antigen-processing and presentation pathway.

The possible existence of macromolecule-sized pores in another mycobacterial species has also been examined. Schaible et al. (21) used immunoelectron microscopy to demonstrate that intracytoplasmic epitope-tagged dextrans (10,000 Da) do not enter the M. avium vacuole. In contrast, intracytoplasmic epitope-tagged high-molecular-weight dextrans enter the Leishmania mexicana vacuole, possibly via an autophagic process (21).

Our findings are not consistent with escape of M. tuberculosis into the cytoplasm of human host cells. Interestingly, when Hess et al. (12) infected macrophages with an M. bovis BCG that expressed recombinant listeriolysin (Hly), they found no evidence that the M. bovis BCG escaped from its phagosome into the cytoplasm, although they did find evidence of Hly outside of the phagosome and evidence of improved MHC-I presentation of cophagocytosed soluble protein by the infected macrophages. The Hly-expressing M. bovis BCG had a reduced capacity to persist in macrophages compared with the parental strain (12). The extraphagosomal, intracytoplasmic compartment may not be a permissive environment for bacteria that have not specifically adapted to intracytoplasmic growth and replication. Goetz et al. (10) microinjected individual bacteria into Caco-2 cells and examined their capacity to grow and multiply within the cytosol. Whereas L. monocytogenes and Shigella were able to grow in the intracytoplasmic compartment, many other bacteria were unable to do so, including Legionella pneumophila, Salmonella enterica, and Bacillus subtilis (10). Growth and multiplication in the cytoplasmic compartment require specific adaptation to that compartment. It is not known whether M. tuberculosis is adapted to or capable of growth within the extraphagosomal, cytoplasmic compartment.

While M. tuberculosis clearly resides within a membrane-bound phagosome in human macrophages, it remains possible that the phagosomal membrane contains small pores that allow the passage of molecules less than 50,000 Da. In this regard, contact-dependent hemolytic activity has been described for M. tuberculosis (14) and Mycobacterium haemophilum (8), but the mechanism underlying this activity and whether or not it is associated with formation of discrete pores has not been reported. It also is not clear whether this lytic capacity is confined to red blood cells or can also be exerted on macrophages. Contact-dependent hemolytic activity was not observed for M. africanum (14), the organism studied by Teitelbaum et al. (24), and it is unclear whether there is any relation between mycobacterial hemolytic activity and any putative pore-forming capacity. Many microbial pathogens induce pores of discrete sizes in the membranes of mammalian cells (2). The sizes and mechanisms of the pores that are formed by microbial pathogens are quite variable. Whereas streptolysin O creates pores that are permeable to large proteins, such as lactate dehydrogenase (140,000 Da), the pores that are formed by Legionella pneumophila are smaller than 3 nm and impermeable to dextrans of 3,000 Da or larger (15). While it is certainly possible that M. tuberculosis forms pores in its phagosome, our data indicate that any such pores are not permeable to proteins of 50,000 Da or larger.
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