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Cationic Liposomes Target Angiogenic Endothelial Cells in Tumors and Chronic Inflammation in Mice

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Abstract

This study sought to determine whether angiogenic blood vessels in disease models preferentially bind and internalize cationic liposomes injected intravenously. Angiogenesis was examined in pancreatic islet cell tumors of RIP-Tag2 transgenic mice and chronic airway inflammation in Mycoplasma pulmonis–infected C3H/HeNCr mice. For comparison, physiological angiogenesis was examined in normal mouse ovaries. We found that endothelial cells in all models avidly bound and internalized fluorescently labeled cationic liposomes (1,2-dioleoyl-3-trimethylammonium-propane [DOTAP]/cholesterol or dimethyldioctadecyl ammonium bromide [DDAB]/cholesterol) or liposome–DNA complexes. Confocal microscopic measurements showed that angiogenic endothelial cells averaged 15–33-fold more uptake than corresponding normal endothelial cells. Cationic liposome–DNA complexes were also avidly taken up, but anionic, neutral, or sterically stabilized neutral liposomes were not. Electron microscopic analysis showed that 32% of gold-labeled liposomes associated with tumor endothelial cells were adherent to the luminal surface, 53% were internalized into endosomes and multivesicular bodies, and 15% were extravascular 20 min after injection. Our findings indicate that angiogenic endothelial cells in these models avidly bind and internalize cationic liposomes and liposome–DNA complexes but not other types of liposomes. This preferential uptake raises the possibility of using cationic liposomes to target diagnostic or therapeutic agents selectively to angiogenic blood vessels in tumors and sites of chronic inflammation. (J. Clin. Invest. 1998, 101:1401–1413.) Key words: angiogenesis • blood vessels • cancer • endosomes • gene therapy

Introduction

Angiogenesis is a consistent feature of tumors and chronic inflammation. Indeed, the growth of tumors and the perpetuation of chronic inflammation may depend upon angiogenesis (1, 2). Therefore, one strategy of treating these diseases is to inhibit new vessel growth (2–4). The identification of distinctive features of angiogenic endothelial cells could make it possible to target therapeutic agents selectively to angiogenic vessels. Studies showing that angiogenic endothelial cells have distinctive adhesion molecules (5), receptors (6), oligosaccharides (7), or other cell surface proteins (4) provide promise that selective delivery to these cells is feasible. Indeed, the utility of these features to target agents to angiogenic blood vessels has been demonstrated in specific disease models (8–10).

However, angiogenesis is not a singular process, and newly formed blood vessels may have very different properties. For example, at least two types of angiogenesis may contribute to vessel growth in tumors. One process involves sprouting and migration of endothelial cells from existing capillaries (sprouting angiogenesis [2]), while the other involves the creation of multiple new vessels from existing ones that are divided by intraluminal septae or pillars (intussusceptive angiogenesis [11]). The anatomic location of tumors or exogenous growth factors can influence the physiological properties of the resultant vessels (12, 13). In addition, angiogenic vessels at sites of inflammation can vary with different stimuli and in different animal strains or species (14–17).

Cationic liposomes have been used to deliver DNA for gene therapy in vivo (18–20). When cationic liposome–DNA complexes are injected intravenously into normal mice, transgene expression occurs in specific organs including the lung and heart (18, 20, 21). This organ-specific pattern of transgene expression is partly related to the pattern of endothelial cell uptake of cationic liposomes. We have found that cationic liposome–DNA complexes injected intravenously into normal mice are taken up by endothelial cells in an organ- and vessel-specific pattern (21). Endothelial cells in the lung and anterior pituitary avidly internalize cationic liposome–DNA complexes into endosomes, while endothelial cells in the brain and posterior pituitary show little or no uptake (21).

Liposome-mediated gene therapy is an attractive approach for diseases dependent on angiogenesis. Noncationic liposomes have been used to deliver therapeutic drugs such as doxorubicin to tumors (22). However, most studies have sought to exploit the increased macromolecular permeability of tumor vessels to deliver liposome-encapsulated agents preferentially to the tumor interstitium (23–26). This approach relies on gradual, passive accumulation of the liposome-encapsulated therapeutic agent in tumors, generally requiring 1–24 h (24, 25, 27). Sterically stabilized liposomes with prolonged circulation times have been developed for this purpose (23, 24). In contrast, cationic liposomes are rapidly cleared from the circulation by the liver, spleen, and lung (21, 28, 29) and have not been used with the intent of targeting agents to endothelial cells in tumors and other sites of angiogenesis.

In view of the finding that cationic liposomes are taken up by endothelial cells in an organ-specific and vessel-specific pat-
tern in normal mice (21), we sought to determine whether angiogenic blood vessels that form in solid tumors or chronic inflammatory diseases are sites of preferential uptake. Because of the differences in angiogenic blood vessels in different disease models, we examined two models of angiogenesis in tumors and another model involving chronic airway inflammation. These disease models in mice were compared with physiological angiogenesis in the normal mouse ovary.

The specific aims of our study were to: (a) compare the amount of cationic liposomes taken up by tumor vessels with that taken up by corresponding normal vessels; (b) identify the locations of cationic liposomes in the wall of tumor vessels, and determine the mechanism of entry into the endothelial cells; (c) compare the uptake by tumor vessels with that by vessels at sites of chronic airway inflammation and normal ovarian follicles and corpora lutea; (d) determine the specificity of uptake by comparing cationic, anionic, neutral, and sterically stabilized neutral liposomes as well as cationic liposome–DNA complexes; and (e) determine whether cationic liposomes can be used to deliver macromolecules, including DNA, preferentially to angiogenic endothelial cells.

We examined the uptake of fluorescently labeled cationic liposomes and liposome–DNA complexes by vascular endothelial cells in transgenic mice with solid tumors, in mice with chronic airway infection due to Mycoplasma pulmonis, and in normal mice. At 20 min or 4 h after red fluorescent liposomes were injected intravenously, the vasculature was stained by perfusion of a green fluorescent lectin, and the location and amount of liposomes in tissue whole mounts were determined by fluorescence and confocal microscopy. The specific cell associations and the intracellular distribution of the liposomes were determined by transmission electron microscopy.

Methods

Murine models of angiogenesis

Tumor models in transgenic mice. We used two tumor models in transgenic mice in which angiogenesis has been well characterized (30, 31). In the first model, designated RIP-Tag2, expression of the SV-40 virus early antigen (Tag) oncogene is driven by the 5′ flanking region of the rat insulin gene including the promoter (RIP). When inserted into the murine genome, this construct induces expression of T antigen specifically in the β cells of pancreatic islets. One important attribute of this model is that various stages of tumor development, and therefore various stages of angiogenesis, are present concurrently in each RIP-Tag2 mouse. Although all of the 300–400 islets express T antigen, the islets initially develop normally. However, about half of the islets become hyperplastic by 6 wk of age, and a small proportion develop into tumors by 12 wk. Tumorogenesis appears to be closely linked with the onset of angiogenesis (32). RIP-Tag2 C57BL/6 mice were bred and raised under barrier conditions in the transgenic mouse facilities at University of California, San Francisco. Mice were genotyped at weaning (3 wk) by PCR on tail-tip DNA using primers for T antigen (33). Nontransgenic littermates were used as controls. Male and female mice were used at 9–11 wk of age.

In the other tumor model, designated K14-HPV16, the oncogene from the human papilloma virus (HPV) is driven by a region of the keratin 14 (K14) promoter (34, 35). In these transgenic mice, viral oncogene expression occurs in epidermal cells. All of the mice develop skin dysplasia accompanied by angiogenesis, and a small proportion develop skin tumors (35).

Chronic airway inflammation model. Angiogenesis associated with chronic airway inflammation was studied in C3H/HeNCR mice infected with M. pulmonis, which afforded the opportunity to examine a type of vascular remodeling different from that in RIP-Tag2 or K14-HPV16 tumors. Previous studies have shown that M. pulmonis infection in the airways of F344 rats causes chronic inflammation with angiogenesis, characterized by extensive proliferation of mucosal blood vessels (14, 16). Recently, we characterized the vascular remodeling in the airways of C3H/HeNCR mice, a strain known to be sensitive to M. pulmonis (36). Airway blood vessels of F344 rats and C3H/HeNCR mice infected with M. pulmonis are similar in that both have normal baseline permeability (with no intervention apart from the infection) but both have exaggerated leakiness after exposure to substance P (14, 16). However, one difference is that airway blood vessels in infected C3H/HeNCR mice become conspicuously enlarged without an increase in vessel number or length (17). Nonetheless, this enlargement is accompanied by a doubling of the number of endothelial cells, consistent with it being a form of angiogenesis (17). Although the terminology concerning angiogenesis is not consistent in the literature, vessel enlargement has been reported in another model of angiogenesis associated with inflammation (15) and may explain the presence of large, dilated, congested blood vessels in the airways of people with asthma (37–40).

Pathogen-free, 8 wk-old, male and female C3H/HeNCR mice (Charles River Laboratories, Hollister, CA) were inoculated intranasally with 3 × 10^7 cfu of M. pulmonis (strain 5782C-UBC C1; obtained from Dr. J. Russell Lindsey, University of Alabama, Birmingham, AL) in a volume of 50 μl after anesthesia (87 mg/kg ketamine and 13 mg/kg xylazine intraperitoneally). Pathogen-free mice were used as controls. Infected and control mice were caged separately under barrier conditions for 4 wk. Antibody titers to M. pulmonis were measured in serum obtained from control and infected mice at the time of the experiment (Microbiological Associates, Bethesda, MD). All of the infected mice had significant titers to M. pulmonis, but none of the pathogen-free mice had measurable titers.

Normal ovarian angiogenesis model. Ovaries were examined in normal female C57BL/6 mice at 9 wk of age with a focus on the angiogenesis associated with mature antral follicles and developing corpora lutea (41, 42).

Animals were housed in the University of California, San Francisco Animal Care Facility and handled in accordance with the procedures of the UCSF Committee on Animal Research.

Fluorescent liposomes and liposome–DNA complexes

Cationic liposomes were prepared with 55 mol% 1,2-dioleoyl-3-trimethylammonium-propane (DOTAP) and 45 mol% cholesterol (both Avanti Polar Lipids, Alabaster, AL). Dimyristoyladeyl amnomion bromide (DDAB; Sigma Chemical Co., St. Louis, MO) was used in place of DOTAP in some experiments. Neutral liposomes were prepared with 55 mol% egg yolk phosphatidylcholine (Avanti Polar Lipids) and 45 mol% cholesterol. Anionic liposomes were prepared with 33 mol% brain phosphatidylserine (Avanti Polar Lipids), 33 mol% egg yolk phosphatidylcholine, and 34 mol% cholesterol. Sterically stabilized liposomes were prepared with 50 mol% egg yolk phosphatidylcholine, 5 mol% PEG2000 DOPE (1,2-dioleoyl-sn-glycerol-3-phosphoethanolamine-N-(polyethylene glycol) 2000); Avanti Polar Lipids), and 45 mol% cholesterol.

1. Abbreviations used in this paper: DOTAP, 1,2-dioleoyl-3-trimethylammonium-propane; HPV, human papilloma virus; K14, keratin 14; RIP, rat insulin promoter; Tag, T antigen.

2. Evans blue concentrations in the trachea of pathogen-free C3H/HeNCR mice were 17.4±1.8 ng/mg wet weight under baseline conditions and 16.2±1.7 ng/mg after substance P (5 μg/kg, i.v.), whereas the corresponding values in M. pulmonis–infected mice 4 wk after inoculation were 19.4±3.7 and 33.1±3.5 ng/mg (Baluk, P., G. Thurston, T.J. Murphy, and D.M. McDonald, unpublished observations).
Liposomes were prepared by rotary evaporation of lipid solutions in chloroform, at 42°C, under vacuum, in a stream of argon. The flask containing the thin lipid film was then placed under high vacuum for 1 h to remove traces of solvent. Sufficient 5% glucose was added to the flask to give a 10 mM total lipid concentration, and the mixture was incubated at 42°C for 1 h with occasional vortexing. The suspension was then sonicated by immersion of the flask in an ultrasonic bath (Laboratory Supplies Co., Hicksville, NY) for 20 min to produce small unilamellar vesicles. These were filtered through a sterile 0.2-μm Nalgene syringe filter and stored at 4°C. Liposomes were prepared and stored under argon.

The fluorescent phospholipid Texas red–DHP-E (1,2-dihexadecanoyl-sn-glycero-3-phosphoethanolamine; Molecular Probes, Eugene, OR) was incorporated during liposome synthesis to give a final fluorophore concentration of 0.25–1.0 mole%. For transmission electron microscopy, the electron-dense lipid Nanogold 1,2-dipalmitoyl-sn-glycero-3-phosphoethanolamine (Nanoprobes, Stony Brook, NY) was incorporated synthetically into DOTAP/cholesterol liposomes to give a final concentration of 0.2 mole%. In some liposomes, lysine-fixable fluorescent Texas red–dextran MW 3000 (10 mg/ml; Molecular Probes) was encapsulated by addition to the 5% glucose used for resuspension. After sonication, the liposomes were separated from free dextran by column chromatography on Sephadex G-75 superfine (Pharmacia, Biotech, Piscataway, NJ). The columns were preloaded with DOTAP/cholesterol liposomes and washed thoroughly with 5% glucose before loading the Texas red dextran liposomal mixture. Liposomes encapsulating Texas red dextran eluted in the column void volume when the column was flushed with 5% glucose.

Cationic liposome–DNA complexes were prepared by the rapid addition of 80 μg of plasmid DNA in 5% glucose to an equal volume of liposomes (1.920 nM total lipid) in 5% glucose. Plasmid DNA (5435 bp) was isolated from Escherichia coli SURE (Stratagene, La Jolla, CA) and purified by alkaline lysis and anion exchange chromatography (Qiagen, Chatsworth, CA). Fluorescently labeled DNA was prepared by the enzymatic addition of Cy3-labeled nucleotides to the termini of linearized DNA. In brief, plasmid DNA was digested with Bsp1280I (New England Biolabs, Beverly, MA) to give five fragments ranging in size from 85 to 1323 bp. Terminal deoxynucleotidyl transferase (Promega Corp., Madison, WI) was used to catalyze the addition of Cy3-dCTP residues (Amersham Life Science, Inc., Arlington Heights, IL) to the 3’ termini of the DNA fragments. Labeled DNA was purified by Qiagen column chromatography as above. Labeled and unlabeled DNA fragments migrated indistinguishably when analyzed by agarose gel electrophoresis.

Fluorescence and confocal microscopy

Liposomes (1,440 nM total lipid in 5% glucose) or liposome–DNA complexes (60 μg plasmid DNA in 1,440 nM total lipid in 5% glucose) were injected via a tail vein in a volume of 0.1–0.2 ml. At 20 min after the injection, the mice were anesthetized by intraperitoneal injection of pentobarbital sodium 50 mg/kg (Abbott Laboratory, North Chicago, IL) and supplemented as necessary. In some mice, Monastral blue (30 mg/kg in 0.1 ml), instead of the liposomes, was injected via a tail vein 5 min before fixation to assess endothelial permeability (14). Blood (0.2 ml) was drawn from the right jugular vein of infected mice and their pathogen-free controls into a heparin-containing syringe for fluorescence microscopy. At 20 min or 4 h after injection of liposomes or liposome–DNA complexes, the vasculature was fixed by perfusion of 1% paraformaldehyde in PBS, pH 7.4, for 3 min at 120 mmHg, and stained by perfusion of fluorescein Lycopersicon esculentum lectin (Vector, Burlingame, CA) (21). Tissues were embedded in paraffin, sectioned (3 mm thick) with a tissue chopper (Sorvall, Newtown, CT), and mounted. Tissues were kept refrigerated and examined within 24 h using a Zeiss Axioskop fluorescence microscope with Fluor objectives or a Zeiss Axioskop 410 confocal microscope equipped with a krypton-argon laser and optimized photomultiplier tubes. Extravasated Monastral blue was viewed in cleared whole mounts of the pancreas or trachea using bright-field optics (14). Images were recorded on Kodak Ektachrome film (ASA 400) or as digital confocal image files.

Transmission electron microscopy

Electron-dense lipid Nanogold 1,2-dipalmitoyl-sn-glycero-3-phosphoethanolamine-DOTAP/cholesterol liposomes (1,440 nM total lipid in 5% glucose) were injected via tail vein in a volume of 0.1–0.2 ml. At 20 min after the injection, tissues were fixed by vascular perfusion of 3% glutaraldehyde in cacodylate buffer, pH 7.1, for 10 min at 120 mmHg, removed, and fixed overnight at 4°C (21). Intercartilaginous and posterior membrane regions of tracheas from M. pulmonis–infected and pathogen-free mice were isolated with a razor blade. Pancreatic islets and tumors were removed and sectioned (80–100 μm thick) with a tissue chopper (Sorvall, Newtown, CT).

To silver-enhance the gold-labeled liposomes, tissue sections were treated with glycine (50 mM in PBS, pH 7.4), rinsed with Heps buffer (50 mM), pH 5.8, containing sucrose (200 mM), stained with a LI Silver enhancement kit (Nanoprobes) (43) using a development time of 20 min, fixed with sodium thiosulfate (250 mM) in Heps buffer (20 mM), pH 7.4, and rinsed with Heps buffer (50 mM), pH 5.8, containing sucrose (200 mM).

Tissue sections were postfixed in osmium tetroxide in one of two ways: silver-enhanced tissues were treated with 0.1% osmium tetroxide in 100 mM cacodylate buffer, pH 7.4, for 30 min at 4°C, while non-enhanced tissues were treated with 2% osmium tetroxide in the same buffer for 14–18 h at 4°C. After treatment with uranyl acetate (2% for 48 h at 37°C), tissues were embedded in epoxy resin (LXI12; Ladd Research, Burlington, VT), thin-sectioned (Ultracut; Leica, Deerfield, IL), and stained with lead citrate. The specificity of the silver enhancement step was tested by omitting the silver enhancement, osmium, or uranyl acetate staining. Sections 80 nm in thickness were examined with a Zeiss EM-10 electron microscope.

Quantitation of uptake of fluorescent and gold-labeled liposomes

To quantitate liposome uptake, a series of 10–12 confocal images separated by 2.5 μm in the focal (z) axis was collected in both the fluorescein and Texas red channels using a ×20 NA 0.6 lens (Zeiss) and standardized settings of the confocal pinhole size, photomultiplier tube gain, and laser power. Images were recorded of vessels in tumors >1 mm in diameter from RIP-Tag2 mice and of mucosal vessels across the cartilage rings in tracheas of M. pulmonis–infected mice. Corresponding images were collected from normal islets of wild-type mice and tracheas of pathogen-free mice. Projections, showing the vessel images (fluorescein-labeled) and liposome images (Texas red-lipid) separately, were generated from the image series. The confocal software was used to outline regions ~200 μm² on the vessel images, then the average fluorescence value of the pixels within the corresponding regions of the liposome images was measured. The pixel fluorescence values, ranging from 0 (dark) to 255 (saturated), represent the digitized output from the confocal photomultiplier tubes. Background intensity was determined by measuring fluorescence in selected regions adjacent to the vessels and was subtracted from the measured intensities. Measurements were made on 25 vessels per trachea and 4 tracheae per group (n = 4), or on 3 vessels per islet or tumor, 10 islets or tumors per mouse, and 4 mice per group (n = 4).

Silver-enhanced, gold-labeled liposomes associated with endothelial cells in RIP-Tag2 tumors were photographed by electron microscopy and printed at ×56,000. The locations of 400 sequentially photographed liposomes having two to four silver grains were classified as located on the luminal endothelial cell surface, inside an endothelial cell, or near the abluminal surface of the endothelium. Liposomes on the luminal endothelial cell surface were further classified as assci-
ated with fenestrae, membrane invaginations (caveolae or coated pits), or other regions of membrane. Liposomes inside an endothelial cell were further classified as in small vesicles, larger organelles resembling multivesicular bodies, or complex structures composed of multiple interconnected vesicles. Liposomes near the abluminal surface of the endothelium were further classified as associated with (within 100 nm of) fenestrae, associated with coated membrane invaginations or caveolae, or not associated with endothelial cells. The total length of luminal endothelial cell surface and the proportion of membrane composed of fenestrae were measured on electron micrographs at \( \times 56,000 \) using a digitizing tablet (44).

**Statistical analysis**

Measurements are presented as mean±SE (n = 4 mice per group) unless otherwise indicated. Significance of differences between groups was assessed by unpaired Student’s t test. Differences with \( P < 0.05 \) were considered significant.

**Results**

*Avoid association of cationic liposomes with tumor vessels.* Pancreatic islets and tumors were readily seen by fluorescence microscopy after the vasculature was stained by perfusion of the green fluorescent lectin (Fig. 1, A and B). In these preparations, the vasculature of normal nontransgenic islets had little or no Texas red fluorescence at 20 min or 4 h after injection of fluorescein-labeled DOTA/cholesterol cationic liposomes (Fig. 1 A). By comparison, the vasculature of tumors in RIP-Tag2 mice had intense Texas red fluorescence at both time points (Fig. 1 B). In addition to the avid uptake by tumor vessels, there was a normal pattern of uptake of the cationic liposomes by macrophages in the liver and spleen and by endothelial cells in the lung and certain other organs, as described previously (21).

Because tumorigenesis proceeds asynchronously among the 300–400 islets in the pancreas of RIP-Tag2 mice, various stages of tumor development could be compared in the same pancreas (31, 32). The intensity of Texas red fluorescence corresponded to the size of the islets or tumors (Fig. 1 C). Normal-sized islets in RIP-Tag2 mice had little or no Texas red fluorescence, as in wild-type mice. However, vessels in enlarged (hyperplastic) islets had focal regions of red fluorescence (Fig. 1, C and D), and those in tumors had uniformly intense red fluorescence (Fig. 1 E). Many focal regions of red fluorescence in hyperplastic islets were associated with vessel protrusions that appeared to end blindly (Fig. 1 F). The amount of Texas red fluorescence due to cationic liposomes in tumor vessels averaged 33 times that associated with corresponding vessels in wild-type mice (Fig. 2). Similarly, angiogenic blood vessels in regions of dysplastic skin of K14-HPV16 mice showed greater Texas red fluorescence than did vessels in normal skin.

*Uptake and intracellular localization versus extravasation of liposomes by tumor vessels.* Fluorescent liposomes in RIP-Tag2 tumors examined by confocal microscopy appeared to be closely associated with lectin-stained blood vessels. No Texas red fluorescence was seen in the surrounding tissue (Fig. 1, E and F), suggesting that the liposomes were associated with the endothelial cells or other components of the vessel wall. Little or no extravasated Monastral blue was found within tumors, but focal regions of extravasated Monastral blue were found on some vessels at the interface of tumors and the surrounding pancreatic tissue.

To determine whether the liposomes were taken up by endothelial cells, extravasated, or both, we obtained more detailed information on the location of the liposomes in RIP-Tag2 tumors by examining silver-enhanced, gold-labeled liposomes by electron microscopy. At 20 min after injection, liposomes were found on the luminal surface of the endothelium (Fig. 3, A, C, and D), inside endothelial cells (Fig. 3, B and D), and on the abluminal side of the endothelium (Fig. 3 E). Many liposomes on the luminal surface were associated with membrane invaginations (Fig. 3 A) or fenestrae (Fig. 3 C). Some liposomes were in coated invaginations of the plasma membrane or near caveolae. Most liposomes inside endothelial cells were in small vesicles or larger organelles resembling multivesicular bodies (Fig. 3, B and D). Liposomes on the abluminal side of the en-

**Figure 1.** Fluorescence and confocal micrographs showing the vasculature of mouse pancreas after intravenous injection of Texas red–labeled cationic liposomes and vascular perfusion of fluorescein-labeled lectin. (A) Little fluorescence associated with cationic liposomes in islet (arrow) and acini of normal mouse pancreas, and (B) abundant fluorescence in large tumor (arrow) in transgenic RIP-Tag2 mouse. Yellow color in tumor is due to colocalization of red and green fluorescence. (C) Different stages of tumor development in pancreas of RIP-Tag2 mouse showing different amounts of cationic liposome fluorescence: weak red fluorescence in normal-sized islets (arrowheads), focal bright fluorescence in hyperplastic islets (short arrows), and more uniform bright fluorescence in small tumor (long arrow). (D) Hyperplastic islet with foci of bright fluorescence of cationic liposomes in some vessels (arrows) but not others (arrowheads). (E) Uniform bright fluorescence in vessels of large tumor similar to one in B; none is evident outside vessels. (F) Higher magnification view of hyperplastic islet showing focal sites of liposome fluorescence associated with vessel protrusions that end blindly (arrows). No Texas red fluorescence visible outside vessels. Scale bars: A and B, 200 \( \mu \)m; C, 200 \( \mu \)m; D and E, 50 \( \mu \)m; F, 20 \( \mu \)m.

Cationic Liposomes Target Angiogenic Endothelial Cells
Figure 3. Electron micrographs of silver-enhanced, gold-labeled cationic liposomes in tumor vessels 20 min after injection into RIP-Tag2 mice. (A) Clusters of silver grains (arrowhead) identifying liposomes near invagination of plasma membrane of endothelial cell. (B) Silver grains on liposomes (arrow) in multivesicular body in endothelial cell. (C) Silver grains on liposomes (arrowheads) on luminal surface of fenestrae. (D) Silver grains on liposomes near invaginations (arrowheads) on luminal plasma membrane of endothelial cell and in multivesicular bodies (arrows). Endothelial junction is closed. (E) Silver grains on liposomes (arrowheads) near endothelial fenestrae (arrows). Two liposomes are near abluminal surface of endothelium and one is on luminal surface near fenestrae. (F) Silver grains (arrow) associated with intracellular organelle in endothelial cell of tumor processed without uranyl acetate in block or lead citrate staining. Scale bar, 200 nm.
dothelium were confined to the basement membrane; many of these were near fenestrae (Fig. 3 E). Silver particles on liposomes in specimens in which the uranyl acetate or lead citrate was omitted appeared identical to those in other preparations, but cellular morphology was indistinct (Fig. 3 F).

The proportions of luminal, intracellular, and extravascular liposomes were determined from the distribution of silver-enhanced gold-labeled liposomes in 49 sequential transmission electron micrographs of vessels in RIP-Tag2 tumors. Of the 400 tumor vessel-associated liposomes examined, 32% were on the luminal surface of the endothelium, 53% were inside endothelial cells, and 15% were on the abluminal surface of the endothelium (Table I). Of the liposomes on the luminal surface, the largest proportion (51%) was associated with endothelial fenestrae, even though the fenestrae constituted only 4% of the endothelial surface (137 fenestrae, with a total length of 7.4 μm, were found in a 184-μm length of endothelium). The remainder was associated with caveolae or coated invaginations (17%) or with other regions of the plasma membrane (32%). Of the liposomes inside endothelial cells, 89% were in multivesicular bodies, 10% were in small vesicles, and ~1% were in complex structures composed of multiple interconnecting vesicles. Of the liposomes on the abluminal side of the endothelium, 47% were within 100 nm of fenestrae, 33% were near caveolae or coated invaginations of the abluminal plasma membrane, and 20% were located elsewhere in the endothelial basement membrane. No gaps were evident in the 11 endothelial cell junctions present in this sample.

Uptake of cationic liposomes by blood vessels in chronic inflammation. The vasculature of the tracheal mucosa of normal C3H/HeNCr mice had little Texas red fluorescence at 20 min or 4 h after injection of the cationic liposomes (Fig. 4 A). In contrast, these vessels in mice infected with M. pulmonis for 4 wk had conspicuous Texas red fluorescence (Fig. 4 B and C). At 20 min after injection, patches of Texas red fluorescence coated the luminal surface of some vessels (Fig. 4 B), whereas at 4 h the fluorescence was in the form of discrete dots (Fig. 4 C). Some of the red dots were slightly beneath the luminal surface, consistent with an intracellular location (Fig. 4 D), but extravasation could not be excluded. However, electron microscopic examination showed that liposomes were located on the luminal plasma membrane (Fig. 5, A and B), in coated (Fig. 5 B) and uncoated membrane invaginations, and in intracellular vesicles and multivesicular bodies similar to those in the tumor vessels. No endothelial fenestrations were found in these vessels in a total of 121 μm of endothelial cell surface examined in 29 electron micrographs.

Because the vasculature of the tracheal mucosa of infected C3H/HeNCr mice had a simple architecture, it was possible to identify the vessel components and determine that most of the uptake was in enlarged vessels in the position of postcapillary venules and collecting venules. These remodeled vessels are very conspicuous after M. pulmonis infection (17). The average amount of liposome fluorescence associated with the remodeled vessels in infected tracheas was 15-fold that of corresponding normal vessels (Fig. 6). A difference of 100-fold was found in some vessels.

Uptake of cationic liposomes by angiogenic vessels in the ovary. Intense Texas red fluorescence was found in blood vessels around large ovarian follicles and in newly formed corpora lutea (Fig. 4 E). The Texas red fluorescence was confined to the plexus of vessels surrounding large antral follicles (Fig. 4 F), while none was observed inside follicles. Vessels in large, rapidly growing corpora lutea also had intense punctate Texas red fluorescence (Fig. 4 F), but those in smaller, regressing corpora lutea had little or none.

Specificity of uptake of cationic liposomes by angiogenic endothelial cells. The specificity of the uptake of cationic liposomes by angiogenic endothelial cells in tumors and ovaries was assessed by comparing Texas red-labeled DOTAP/cholesterol.
terol cationic liposomes with other lipidosome formulations. We found no evidence of uptake of Texas red–labeled anionic, neutral, or sterically stabilized neutral liposomes by blood vessels in RIP-Tag2 tumors, ovaries, or other organs at 20 min after injection.

However, the uptake of Texas red–labeled DOTAP/cholesterol–DNA complexes by angiogenic blood vessels in RIP-Tag2 tumors and ovaries resembled that of the cationic liposomes. Similarly, liposome–DNA complexes formulated with dimethyl dioctadecyl ammonium bromide (DDAB), another cationic lipid, resembled that of free DOTAP/cholesterol liposomes. To determine whether the bright fluorescence of tumor vessels after injection of liposome–DNA complexes indeed reflected the uptake of liposome–DNA complexes and not just the uptake of free liposomes, we injected Cy3-labeled DNA complexed with unlabeled liposomes and examined the distribution of Cy3 fluorescence in normal islets and tumors of RIP-Tag2 mice. The tumor vessels had much brighter fluorescence than corresponding normal vessels (Fig. 4, G and H) and resembled those found after injection of labeled liposomes.

**Delivery of macromolecules to angiogenic endothelial cells.** The feasibility of using cationic liposomes to deliver macromolecules other than DNA to angiogenic endothelial cells was determined by examining the uptake of liposome-encapsulated Texas red–labeled dextran by tracheal blood vessels in *M. pulmonis*–infected mice. The uptake of the encapsulated dextran matched that of Texas red–labeled cationic liposomes. At 4 h after the injection, the encapsulated dextran appeared as abundant fluorescent red dots in the endothelial cells of the angiogenic blood vessels (Fig. 4 I). No red fluorescence was associated with tracheal vessels 4 h after the injection of free Texas red–labeled dextran without cationic liposomes.

**Discussion**

These experiments show that angiogenic blood vessels in tumors of RIP-Tag2 mice and K14-HPV16 mice bound and took up cationic liposomes much more than did the corresponding normal vessels. At 20 min after injection, most liposomes were on the luminal surface of angiogenic endothelial cells or in multivesicular bodies or vesicles inside the cells. Angiogenic endothelial cells at sites of chronic inflammation in *M. pulmonis*–infected mice and in normal ovaries also took up cationic liposomes. The magnitude of uptake assessed by confocal microscopy in vessels of tumors and chronically inflamed tracheas was 15–33-fold that of corresponding normal vessels. Angiogenic endothelial cells in these models of angiogenesis also took up cationic liposome–DNA complexes, but not anionic, neutral, or sterically stabilized neutral liposomes. Finally, cationic liposomes were used to deliver fluorescent dextrans to angiogenic endothelial cells as a model of macromolecular intracellular delivery.

**Endothelial cell uptake versus leakage of cationic liposomes.** Plasma leakage is a feature of some angiogenic blood vessels (45–49). Therefore, we sought to determine whether cationic liposomes associate with angiogenic vessels by virtue of binding and uptake by endothelial cells or by virtue of extravasation. Several lines of evidence suggest that the liposome accumulation observed in the models used in this study was primarily by endothelial cell uptake, not by leakage. First, confocal microscopic observations of fluorescent liposomes and lectin-stained vessels showed that the liposomes remained closely associated with the luminal surface of endothelial cells, both in the tumor models and in the infection model. Second, in all models the distribution of the liposomes changed from a diffuse coating of the luminal endothelial cell surface at 20 min to discrete dots at 4 h, without evidence of appreciable extravasation over time. Third, electron microscopic observations showed that 85% of the liposomes associated with tumor vessels in RIP-Tag2 mice were attached to the luminal surface of the endothelium or were inside endothelial cells. Only 15% of the liposomes associated with tumor vessels had crossed the endothelium at 20 min. This evidence of the internalization of liposomes into endosomes is consistent with the change in distribution over time observed by confocal microscopy and fits our previous observations of liposome internalization by certain types of endothelial cells in normal mice (21). Fourth, the avid uptake of liposomes was specific to cationic liposomes and was not observed with anionic, neutral, or sterically stabilized neutral liposomes, whereas sterically stabilized liposomes can extravasate from leaky tumor vessels (25). Addition of DNA to form liposome–DNA complexes did not change this property of cationic liposomes, and the pattern of uptake of liposome–DNA complexes was the same regardless of whether the lipid or the DNA was labeled. Finally, tumor vessels in RIP-Tag2 mice were not abnormally leaky as judged by the limited extravasation of the particulate tracer Monastral blue.

**Figure 4.** Confocal micrographs showing vessels in mouse trachea (A–D, I), ovary (E and F), pancreatic islet (G), and tumor (H) after intravenous injection of cationic liposomes or liposome–DNA complexes. Vessels stained green with fluorescein *L. esculentum* lectin. Weak fluorescence from Texas red cationic liposomes (red) in lectin-stained (green) tracheal vessels of pathogen-free mouse (A) compared with bright red-orange fluorescence (arrows) in tracheal vessels of *M. pulmonis*–infected mice at 20 min (B) or 4 h (C) after injection. (D) Higher magnification of tracheal vessels of *M. pulmonis*–infected mouse 4 h after injection. Texas red fluorescence in discrete red dots just beneath the luminal surface of endothelium (green), suggestive of intracellular location. Some adherent leukocytes (arrowhead) are also brightly fluorescent. (E) Low magnification fluorescence micrograph of ovary showing bright Texas red fluorescence of cationic liposomes in vessels of follicle (arrow) and corpus luteum (orange structure beneath follicle). (F) Confocal micrograph of ovary showing bright Texas red fluorescence of cationic liposomes in vessel plexus around follicle. Interior of follicle has no fluorescence. Also shown is bright Texas red fluorescence in part of a large corpus luteum. (G and H) Confocal micrographs showing Cy3-labeled DNA (red) complexed to nonfluorescent cationic liposomes in lectin-stained pancreatic vessels (green) of RIP-Tag2 mouse 4 h after intravenous injection. Weak Cy3 fluorescence in vessels of normal-sized islet (G) compared with bright Cy3 fluorescence in tumor vessels (H, arrows). (I) Texas red–labeled dextran encapsulated in unlabeled cationic liposomes in tracheal venules of *M. pulmonis*–infected mouse at 4 h after injection. Texas red dextran (red, arrows) in discrete dots just beneath luminal surface of lectin stained venule (green), suggestive of intracellular location. Scale bars: A–C, 50 μm; D, 12 μm; E, 200 μm; F, 50 μm; G and H, 20 μm; I, 20 μm.
Similarly, the tracheal vasculature of *M. pulmonis*-infected C3H/HeNCr mice has about the same amount of plasma leakage as that of pathogen-free rats.

The mechanism by which 15% of the liposomes in tumors of RIP-Tag2 mice extravasated was not apparent in the electron micrographs. Some liposomes were near the abluminal surface of fenestrae. Although fenestrae are not believed to allow passage of particulate or protein tracers (50, 51), interactions between the liposomes and the fenestral diaphragms may facilitate such passage. Other mechanisms of leakage, such as endothelial cell gaps or vesicular transport (49, 52), could contribute to liposome leakage in tumors of RIP-Tag2 mice, but no intercellular gaps were found in the sections examined by electron microscopy in this study. In other models of angiogenesis, the number of extravasated liposomes would depend on the particular permeability and transcellular transport characteristics of the endothelium and the driving forces in the vasculature.

**Mechanism of liposome uptake by endothelial cells.** In blood vessels in tumors of RIP-Tag2 mice and tracheas of *M. pulmonis*-infected mice, liposomes were found on the luminal surface of endothelial cells and inside endothelial cells in small vesicles and multivesicular bodies. Fluorescence was measured on confocal micrographs of 25 vessels per trachea, 4 mice per group (*n* = 4). Fluorescence values represent digitized signal (0 to 255) from red channel photomultiplier tube. Values are mean±SE. *Significantly different from control value, P < 0.05, unpaired Student’s *t* test.

**Figure 5.** Electron micrographs showing clusters of liposomes (not labeled with gold) adherent on the luminal surface (*A* and *B*, arrows) and in a coated invagination (*B*, arrowheads) of endothelial cells in tracheal vessels of *M. pulmonis*-infected mouse 20 min after injection. Scale bar, 200 nm.

**Figure 6.** Amount of Texas red fluorescence in tracheal vessels of pathogen-free C3H/HeNCr mice and *M. pulmonis*-infected (4 wk) mice 4 h after injection of Texas red-labeled cationic liposomes. Fluorescence was measured on confocal micrographs of 25 vessels per trachea, 4 mice per group (*n* = 4). Fluorescence values represent digitized signal (0 to 255) from red channel photomultiplier tube. Values are mean±SE. *Significantly different from control value, P < 0.05, unpaired Student’s *t* test.
issue of whether other mechanisms of uptake, such as uptake by noncoated vesicles or liposome-membrane fusion, can also occur.

At 20 min after injection, 51% of the liposomes on the endothelial cell surface in tumor vessels were associated with fenestrae, although fenestrae constituted only 4% of the luminal endothelial cell surface. The positive charge of the liposomes may explain their avidity for fenestrae. Cationic ferritin, but not native ferritin, binds to fenestrae, indicating that fenestrated diaphragms have a negative charge (55, 56).

Despite the preferential adherence of cationic liposomes to endothelial fenestrae in RIP-Tag2 tumors, the avid uptake of cationic liposomes by angiogenic endothelial cells cannot be fully explained by the abundance of fenestrae. First, islet vessels in normal mice had little uptake, despite the presence of fenestrated endothelial cells. Second, avid uptake was also found in tracheal vessels of infected mice, even though these vessels were not fenestrated. Third, intracellular liposomes were found in nonfenestrated regions of endothelial cells, suggesting that fenestrae are not necessary for vesicular uptake. Fourth, large differences in the amount of uptake are present among fenestrated vessels in different organs, such as the thyroid, anterior pituitary, posterior pituitary, and pancreatic islets (21).

Thus, mechanisms of binding other than the affinity of cationic liposomes for fenestrae must be important. One possibility is that liposome binding and internalization by nonfenestrated regions of the plasma membrane is receptor mediated, while binding to fenestrae is by nonspecific charge interaction and does not result in internalization. Because of the rapidity of the internalization, an examination of the binding distribution at earlier times after injection could help to determine the relative contributions of binding to fenestrated and nonfenestrated regions of endothelial cells.

Identification of sites of uptake of cationic liposomes. Cationic liposomes and liposome–DNA complexes are rapidly cleared from the bloodstream with a half-life of < 5 min (29). This clearance results predominantly from the rapid uptake by macrophages in the liver and spleen and by endothelial cells in an organ- and vessel-specific pattern (21). Endothelial cells in capillaries of the lung and anterior pituitary avidly bind and take up cationic liposomes, as do those in high endothelial venules of lymph nodes, while endothelial cells in many other sites, including pancreatic islets and trachea, take up few liposomes.

In this study, the avid uptake of cationic liposomes by angiogenic endothelial cells in mice with tumors or chronic airway inflammation was superimposed on the normal pattern of uptake (21). The liver, spleen, and lung were still the principal sites of uptake, because of their large size in comparison to that of the tumors or inflamed airways. Modification of the surface properties of cationic liposomes or addition of specific targeting molecules may make it possible to increase the uptake by angiogenic endothelial cells or decrease the uptake by normal endothelial cells and macrophages.

Most of the blood vessels in RIP-Tag2 tumors form during an ~ 3–4-wk period of rapid tumor growth (31), during which tumor volume increases by several orders of magnitude. Similarly, most of the vessels of enlarging ovarian follicles and corpora lutea form while these structures are growing rapidly. Because newly formed blood vessels predominate in these systems, most of the avid uptake of cationic liposomes is by new endothelial cells. This issue is not as straightforward for the model of M. pulmonis infection in C3H/HeNCr mice, where the blood vessels of the tracheal mucosa enlarge but do not become more numerous. In this case, the uptake is evidently by endothelial cells within existing vessels that enlarge (17). It is not yet known whether the avid uptake coincides temporally with the proliferation of the endothelial cells.

Heterogeneity of endothelial cells in angiogenic vessels. The avid uptake of liposomes was not uniformly distributed. In RIP-Tag2 mice, focal regions of vessels in hyperplastic islets (31) had an unusually high uptake. Furthermore, some of these foci were located in vessel sprout-like protrusions that ended blindly. In contrast, uptake was uniformly high in the vasculature of large tumors. In M. pulmonis–infected mice, the avid uptake was confined to particular segments of the microvasculature, in the position of postcapillary venules and collecting venules, but even here, the uptake was much higher in some vessels than others.

The presence of focal sites of avid liposome uptake raises the possibility of a novel heterogeneity among endothelial cells of newly formed or remodeled vessels. Because the uptake by vessels in hyperplastic islets and small tumors of RIP-Tag2 mice tended to be focal, whereas that in large tumors was uniform, the switch in uptake may occur at an early stage in the angiogenic process. It is unknown how these sites of uptake relate to the presumptive processes that occur during angiogenesis, for example, endothelial cell proliferation, migration, basement membrane remodeling, lumen formation, and vessel maturation (2).

Various phenotypic markers of angiogenic endothelial cells, including the integrin α,β (5), the profile of cell surface oligosaccharides (7), vascular endothelial growth factor receptors (6), and other cell surface proteins (4), have been reported in particular systems of angiogenesis. An important next step will be to determine whether these phenotypic markers define the same population of endothelial cells as does the avid uptake of cationic liposomes. If not, multiple endothelial cell phenotypes may participate in angiogenesis, and the concept of an angiogenic endothelial cell may require expansion (57). A precedent for such cell heterogeneity is the report that migrating endothelial cells are primarily at the leading tip of a vascular sprout while proliferating cells are at the base (58).

Although liposomes were avidly taken up by endothelial cells in all of the models of angiogenesis we examined, these observations need not apply to all other models of angiogenesis. We intentionally used models of diseases in which angiogenesis occurs, with the long-term aim of altering the disease progression by reversing the angiogenesis. However, the signals that induce angiogenesis in these systems are likely to be complex, and the process of vessel growth has not been fully characterized. It may be necessary to use simpler models of angiogenesis to elucidate the endothelial cell mechanisms, relevant signaling molecules, and particular stages of vessel growth involved in the uptake of cationic liposomes by angiogenic endothelial cells.

Using cationic liposomes to target agents to angiogenic endothelial cells. The preferential uptake of cationic liposomes raises the possibility of delivering diagnostic or therapeutic agents to growing or remodeling blood vessels. This approach differs conceptually from previous approaches of liposome-mediated delivery that targeted tumor cells directly by exploiting the increased permeability of tumor vessels to achieve...
higher drug concentrations near tumor cells (25, 27). One of the problems of using endothelial cell leakiness to deliver agents to tumors is that a high interstitial pressure may limit the convective movement of molecules from the vessel lumen into the tumor interstitium (52). In comparison, cationic liposomes would deliver substances to the endothelium as long as there is blood flow, and delivery would not depend on convective movement across the vessel wall. Thus, cationic liposomes could be used to deliver agents to angiogenic endothelial cells rather than to tumor or interstitial cells.

Strategies of delivering substances selectively to angiogenic endothelial cells include targeting specific growth factor receptors (4), upregulated coagulant factors (59, 60), and key cell adhesion proteins (61). The avid uptake of cationic liposomes by angiogenic endothelial cells described here may provide a new tool to deliver diagnostic or therapeutic agents to these cells. For example, substances that initiate coagulation and stabilize neutral liposomes are not taken up by these cells. This work was supported in part by National Institutes of Health Grants PO1 CA69094 and CA21765.

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