Technical Note

High-Resolution MRI of Deep-Seated Atherosclerotic Arteries Using Motexafin Gadolinium

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Purpose: To evaluate the potential of using motexafin gadolinium (MGd) to characterize atherosclerotic plaques of deep-seated arteries with MRI.

Materials and Methods: We exposed vascular endothelial cells (EC) and smooth muscle cells (SMC) in vitro to varying concentrations of MGd. The fluorescence properties of MGd were then exploited using confocal microscopy to image exposed cells. For an in vivo validation study, we performed surface coil–based and intravascular coil–based high-resolution MRI of the iliac arteries and the abdominal aorta of three atherosclerotic Yucatan pigs. Subsequently, MGd enhancement of the target vessel walls was quantitatively evaluated and MR images were correlated with histology of the target vessels.

Results: The in vitro study confirmed the intracellularization of MGd in both cell types and determined the optimum MGd dosage of 0.004 mmol/kg that produced the sufficiently high intracellular fluorescent intensity. The in vivo study showed a steady increase of MGd enhancement to approximately 25% at three hours postinjection of MGd. MRI showed areas of strong enhancement along the lumen boundary, which corresponded to fibrous tissue seen in histology.

Conclusion: This study provides initial evidence that MGd may enhance MR vessel wall imaging for the characterization of plaque in deep-seated arteries.

Key Words: atherosclerotic cardiovascular disease; motexafin gadolinium; contrast agent; vessel wall; intravascular MRI


Atherosclerotic cardiovascular disease is still the leading cause of death in developed countries. Atherosclerosis refers to the buildup of fatty deposits or “plaque” in the innermost lining of the arterial wall. Atherosclerotic plaque causes hardening of the artery and narrowing of the vessel lumen, which can result in the obstruction of blood flow and the reduction of oxygen supply to the important tissues and organs of the body. Current diagnostic techniques, such as conventional X-ray angiography, focus on determining the degree of stenosis in arteries, but it is now understood that the composition and shape of the plaque, and not the degree of stenosis, are the best indicators of future acute events (1–3). A large lipid core with a thin or ruptured fibrous cap is thought to indicate “vulnerable” plaque with an elevated risk of thrombosis. To diagnose a plaque as vulnerable, the plaque composition, rather than plaque burden, must be identified. Previous works using MR contrast agents to discriminate between plaque components in the surface-seated arteries, primarily the carotid arteries, have shown an increased contrast between plaque components, such as fibrous tissue and lipid (1–4).

However, it is a challenging task to generate high-resolution MRI of atherosclerotic plaques in the deep-seated arteries, such as the iliac arteries, the renal arteries, and the aorta, due to insufficient MR signal-to-noise ratios (SNRs) when using the body coil or surface coils. Motexafin gadolinium (MGd; Pharmacyclics Inc.), a gadolinium texaphyrin analog, is primarily designed as an intracellularly localized anticancer agent (5). MGd selectively accumulates in metabolically active tissues, such as tumors. In addition, MGd can produce red fluorescence when stimulated by blue light, and, thus, can be tracked under fluorescence microscopy (5,6). Because MGd contains gadolinium, it can also function as a T1 MR contrast agent. These characteristics of MGd encouraged us to evaluate MGd for high-
resolution MRI of atherosclerotic plaques, which are also metabolically active tissues, similar to tumors. Currently, no studies have been performed investigating the in vitro localization of MGd within the vascular cells involved in atherosclerosis. The potential of MGd to provide plaque-specific contrast enhancement for vulnerable plaque diagnosis of atherosclerotic disease is also unknown.

The objectives of the present study were to: 1) evaluate, in vitro, the intracellular properties of MGd, specific to atherosclerotic plaques, by confirming the internalization of MGd by vascular endothelial cells (EC) and smooth muscle cells (SMC); and 2) investigate, in vivo, the potential of using MGd as an MR T1 contrast agent to discriminate between tissues in the atherosclerotic plaques of the deep-seated iliac arteries in an atherosclerotic animal model.

**MATERIALS AND METHODS**

**MGd**

MGd is an expanded porphyrin (texaphyrin) that contains bound gadolinium within its macrocyclic core. Texaphyrin molecules are known to be attracted to metabolically active cells, such as tumor cells, and therefore have strong potential in tumor imaging (5,7). The mechanisms of cell uptake of MGd are not clear, but considered as the effects of microenvironmental factors present in cells such as low pH and hypoxia (8). The fluorescent properties of MGd allow it to be imaged under a confocal microscope, with fluorescence at 740 nm when excited at 470 nm.

**In Vitro Study**

The in vitro study included two cell groups: 1) the first group to confirm intracellularization of MGd in vascular ECs and SMCs; and 2) the second group to determine the optimum concentration of MGd for subsequent in vivo animal study.

In the first cell group, human ECs and SMCs (Cambrex Bio Science Walkersville, Inc., Walkersville, MD, USA) were cultured in a six-well plate. 0.1 mL of 2 mM MGd and the cell culture medium (provided by the manufacturer) in a 37°C incubator for 24 hours. The cells were then allowed to culture in cell culture medium for an additional 24 hours. After detaching the cells from each other by 0.5 mL trypsin (25%) per well, the cells from each well were collected in a centrifuge tube. Nonexposed control cells were prepared in a similar manner. Subsequently, cells were centrifuged, scraped from each of the four tubes, and then fixed in formalin on microscope slides for confocal microscopy imaging.

The cells were imaged with a Zeiss LSM 410 confocal microscope using a 470 ± 10 nm excitation band-pass filter with a 700-nm emission long-pass (far-red) filter, and a power of 63 mW/cm². Because MGd has an emission wavelength (>700 nm) at which mammalian cells do not autofluoresce, all fluorescent signal detected on the confocal microscope with the 700-nm long-pass filter comes from the presence of MGd. A standard black and white microscope image was also taken and fluorescence images were displayed in pseudocolor over the black and white images to show the cellular location of the MGd.

The second cell group was used to determine the optimum MGd dosage for subsequent in vivo animal study. Additional ECs and SMCs were subcultured and transferred to other six-well plates. The cells in each well were exposed to solutions of 0, 5, 10, 25, 50, and 100 μM MGd, respectively, for 24 hours. They were washed, incubated, and extracted with the same method described above. In each of cell groups, five cells were randomly selected and imaged with the confocal microscope. Signal intensity was averaged over the volume of the cells to obtain a measure of fluorescence, which was subsequently plotted against MGd concentration. The MGd concentration at which the increase fluorescent intensity became negligible was considered the optimum MGd concentration. This concentration was used to determine dosages for the subsequent in vivo study with atherosclerotic animal models.

**In Vivo Study**

**Animals**

In vivo MRI studies were performed on three atherogenic Yucatan pigs (approximately 65 kg). The pigs were subjected to injury of the iliac arteries by balloon denudation followed by a high-cholesterol diet (Research Diets, Inc., New Brunswick, NJ, USA) for 18–24 months. In addition to essential nutrients, vitamins, and salts, the atherogenic diet contains 4% cholesterol, 14% beef tallow, and 1% hog bile extract in daily amounts of 1000 g (9). All animals were treated according to the “Principles of Laboratory Animal Care” of the National Society for Medical Research and the “Guide for the Care and Use of Laboratory Animals” (10). The Animal Care and Use Committee at our institution approved the experimental protocol.

**Surface Coil-Based MRI**

Under general anesthesia, the first and second Yucatan pigs were placed in the 1.5T GE MR scanner. Dual 3- and 5-inch surface coils were placed on the abdomen and lower back, respectively. Two-dimensional time-of-flight (TOF) MR angiography was performed to obtain scout images to localize stenotic areas in the iliac arteries distal to the aortic bifurcation and in the lower abdominal aorta. Then, transverse images covering 3 cm of the iliac artery starting at the aortic bifurcation and in the lower abdominal aorta were obtained using high-resolution MRI. MR images were acquired with a double inversion-recovery, fast spin-echo (FSE), black blood, electrocardiograph (ECG)-gated pulse sequence using breathholds in several locations along the 3-cm-long iliac arteries. The imaging parameters were as follows: TI = 280 or 400 msec, TE = 4.7 msec, heart rate (HR) = 80 bpm, TR = 1 heartbeat (~750 msec), field of view.
In Vitro Study

Confocal imaging showed internalization of MGd in both ECs and SMCs. MGd was shown to accumulate in the cytoplasm and was retained there for longer than 24 hours. Figure 2 shows fluorescence signal intensity plotted against concentration for confocal microscopy imaging. Near-maximal intensity was achieved with a 50 μM concentration of MGd. This concentration cor-

Intravascular MRI

In the third Yucatan pig, we attempted to validate the feasibility of performing intravascular high-resolution MRI of the aortic plaques with MGd enhancement. Through a surgical cutdown, a 7-F introducer was inserted into the left carotid artery, through which a custom 0.032-inch MRI-guidewire (MRIG) (11,12) was positioned into the lower abdominal aorta under X-ray guidance. The pig was then transferred to the 1.5T GE scanner and high-resolution imaging was performed using the intravascular MRIG and a 5-inch surface coil placed on the abdomen. FSE, black-blood, ECG-gated scans were prescribed in cross-section to the lower abdominal aorta. The following MRI parameters were used: TI = 260 msec, TE = 11.5 msec, HR = 90 bpm, TR = 1 RR (~670 msec), FOV = 10 cm, matrix size = 256 × 192, ETL = 10, NEX = 10, slice thickness = 2 mm, and scan time = 130 seconds. The MRIG was operated in receive-mode only. After baseline imaging, 68 mL of 5 mM MGd (0.004 mmol/kg) was injected at a flow rate of 5 mL/second. Images were taken up to three hours postcontrast.

Image Analysis

Using the surface coil-based MR images, MGd enhancement of the iliac artery wall was measured. Three imaging slices of each artery were examined before and up to two to three hours postinjection. For each image, a series of 30 points were chosen randomly within the vessel wall. These were averaged and normalized for all three slices to achieve an average MR amplitude. Subsequently, the amplitude was plotted for each agent and for each time point to show how contrast enhancement developed.

Histology

Once MRI was completed, the pig was euthanized with a dose of 100 mg/kg of pentobarbital, and the imaged tissues from the bilateral iliac arteries and the lower abdominal aorta were harvested. The specimens were embedded in paraffin, cut into 5-μm slices on a cross-sectional view, and stained with hematoxylin and eosin (H&E) stain for microscopy examination.

RESULTS

In Vitro Study

Fluorescence was detected in exposed cells and was contained within the cellular membrane, as shown in Fig. 1. Confocal imaging showed internalization of MGd in both ECs and SMCs. MGd was shown to accumulate in the cytoplasm and was retained there for longer than 24 hours. Figure 2 shows fluorescence signal intensity plotted against concentration for confocal microscopy imaging. Near-maximal intensity was achieved with a 50 μM concentration of MGd. This concentration cor-

Figure 2. Average fluorescence intensity plotted against MGd concentration for EC and SMC. The paired histogram for each concentration was similar in shape and showed nearly maximal intensity at 50 μM concentration. au = arbitrary unit.
responded to the manufacturer-recommended maximum dose of 0.004 mmol/kg MGd, which was selected as our optimum concentration of MGd for the subsequent in vivo animal experiments. We achieved the optimal concentration of 0.004 mmol/kg MGd by estimating an amount of 5 liters of blood in the ~65-kg pigs we used. From there we converted the 0.004 mmol/kg dose into a blood concentration of about 50 μM ([0.004 mmol/kg] × [65 kg]/[5 liters] = 0.052 mmol/liter = 50 μM).

In Vivo Study

Since the vessels are small and deep inside the body, the surface coil MR images suffer from low SNR. Although this is the case, the effect of the contrast enhancements to the images could be visualized and quantified. Contrast enhancement with Magnevist developed rapidly to a maximum of 37% by five minutes, and then quickly began to decrease due to washout from the bloodstream. MGd exhibited a different behavior: enhancement increased steadily over the entire three-hour period of imaging. An increased average signal enhancement of approximately 25% was seen at three hours postinjection of MGd (Fig. 3). Wall thickening with nonuniform enhancement was visualized with the enhancement being greatest close to the lumen (Fig. 4).

With intravascular MR vessel wall imaging using the MRIG, image SNR increased significantly and thus we were able to reduce the FOV to 10 cm from 18 cm. Contrast enhancement in the atherosclerotic aortic wall was seen to occur again in a very nonuniform manner. Vessel wall thickening became more clear two hours after intravenous injection of MGd compared to the preinjection image. Areas of strong enhancement were seen along the lumen boundary, which corresponded to the fibrous tissue seen in histology (Fig. 5). Histological examination confirmed the presence of extensive wall thickening due to complicated plaques.

DISCUSSION

MR vessel wall imaging of deep-seated arteries has been a difficult task due to the low SNR when performing MRI of these vessels with a surface or body coil. To overcome this problem, we attempted to apply two different techniques, including: 1) an intracellular MR contrast agent, MGd, to specifically characterize different components of the atherosclerotic plaques, and 2) an intravascular MRIG to generate high-SNR MR vessel wall imaging.

This study has confirmed the in vitro internalization of MGd by both vascular endothelial and SMC using confocal microscopy. MGd was shown to accumulate in the cytoplasm of EC and SMC, and was retained there for longer than 24 hours. These findings are similar to...
those found by Woodburn (6) with tumor cells (7), which indicate that MgD can be taken up by not only tumor cells but also vascular cells. These results may indicate that under in vivo conditions, MgD should localize into fibrous tissue and healthy vessel wall tissue, which contain endothelial and SMC. Lipid pools have few of these cells while necrotic cores are noncellular; therefore, both should have low MgD concentrations at equilibrium. Thus, MgD may allow differentiation between fibrous caps and lipid cores under MRI. In addition, we also determined a suitable dosage (0.004 mmol/kg) of MgD for near-optimal internalization and enhancement of the vascular cells.

Surface coil–based MRI generated during this study demonstrate a potential for MgD to discriminate between atherosclerotic plaque components, which can be further enhanced by using the intravascular MRI technique for better plaque characterization in the deep-seated vessels. In addition, MRI showed that MgD enhancement of the target vessel wall increased slowly over at least a three-hour period. This may be reflective of the slow cellular uptake of MgD by the vascular cells. These results are particularly encouraging, suggesting that MgD is steadily accumulating in the vessel wall and being retained in the vascular cells over time. These manifestations further support that MgD may function as a useful MR contrast agent for plaque characterization of deep-seated vessels using high-resolution MR vessel wall imaging.

Magnevist, a commonly used contrast agent with well-documented effects, was used here as a control. Magnevist is a small molecule that diffuses freely from the bloodstream into the vascular wall. It remains in the extracellular fluid and does not chemically target specific cell types. It is also cleared from the bloodstream rapidly. In this study, Magnevist exhibited characteristic behavior: reaching maximal enhancement very quickly and then diminished as it was cleared from the blood. The administered dose of MgD was 100-fold less than that of Magnevist, which may explain the lack of immediate enhancement of the vessel walls. The slow build-up of enhancement is believed to be due to the necessary time period for vascular cells to uptake and internalize the agent. The limitation of this study is that it was unclear whether MgD enhancement continued to increase beyond this point of three-hours postinjection. A longer time period after injection would be needed to fully determine the enhancement behavior.

MgD is a texaphyrin molecule that covalently binds the paramagnetic metal gadolinium. Texaphyrins have several useful qualities. They accumulate in highly metabolic tissues, not only tumors but also plaques as demonstrated in this study. Some texaphyrins function as radiation enhancers, photosensitizers, or chemotherapy enhancers. In the present study, we demonstrated the added benefit of MgD in functioning as a T1 MR contrast agent. This function is due to the bound gadolinium ion within the MgD molecule. In addition, texaphyrins are fully aromatic with a dark green color and emit fluorescent light when excited by a laser. This allows for accurate molecular optical imaging. Apparently, the further combination of the two MgD functions, therapy and imaging, should enable us to open new avenues, using MgD-enhanced MRI technology to simultaneously diagnose atherosclerotic plaques and guide interventional therapies of atherosclerotic cardiovascular disease.

The limitation of this study is that the initial nominal concentration of MgD for our in vivo experiments was calculated assuming distribution in blood plasma only. Its actual free concentration may depend on binding with albumin, lipoproteins, and blood cells. Furthermore it is possible that MgD is able to diffuse through the vascular wall into the interstitium and some amount of MgD is cleared by the kidneys and liver over two to three hours of experiments. These factors warrant further studies to understand the working mechanisms of MgD as an intracellular MR contrast agent.

In conclusion, the present study provides initial evidence that MgD can generate contrast between various vascular tissue types, such as fibrous tissues involved in atherosclerotic plaques, and may be useful for plaque characterization of deep-seated arteries using high-resolution, contrast-enhanced MR vessel wall imaging.

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