Regional Heterogeneity of Human Myocardial Infarcts Demonstrated by Contrast-Enhanced MRI

Potential Mechanisms

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Abstract
**Background** Myocardial reperfusion is pivotal to the prognosis of patients with acute myocardial infarction. In these patients, coronary flow is generally assessed by angiography and tissue perfusion by tracer scintigraphy. This study was designed to examine whether magnetic resonance imaging (MRI) provides information on myocardial perfusion and damage beyond that supplied by angiography and thallium scintigraphy after acute myocardial infarction.

**Methods and Results** Twenty-two patients with recent myocardial infarction had ECG, echocardiography, coronary angiography, and fast contrast-enhanced MRI. Twelve patients also had exercise thallium scintigraphy. Time-intensity curves obtained from infarcted and noninfarcted regions were correlated with coronary anatomy and left ventricular function. Two perfusion patterns were observed in infarcted regions by comparison with the normal myocardial pattern. All patients but 1 had persistent myocardial hyperenhancement within the infarcted region up to 10 minutes after contrast. In 10 patients, this hyperenhanced region surrounded a subendocardial area of decreased signal at the center of the infarcted region associated with coronary occlusion at angiography, Q waves on ECG, and greater regional dysfunction by echocardiography. Moreover, the extent and location of the MRI abnormalities correlated well with the extent and location of the fixed single-photon emission computed tomography thallium defects.

**Conclusions** Large human infarcts, associated with prolonged obstruction of the infarct-related artery, are characterized by central dark zones surrounded by hyperenhanced regions on MRI. Conversely, reperfused infarcts with less regional dysfunction have uniform signal hyperenhancement. The MRI hyperenhanced segment correlates well with the fixed scintigraphic defect in patients with acute myocardial infarction.

**Key Words:** reperfusion • edema • magnetic resonance imaging • myocardial infarction

**Introduction**

The timing and adequacy of myocardial reperfusion largely determine the prognosis of patients with acute myocardial infarction. When myocardial perfusion is quickly restored after coronary occlusion and maintained to the entire territory at risk, myocardial infarction may be aborted. In most cases, however, only partial restoration of coronary blood flow is achieved after significant injury to the threatened segment of the left ventricular wall has already occurred. Therefore, methods to evaluate the extent and adequacy of spontaneous or therapeutically induced reperfusion are necessary to assess the prognosis and management of patients with myocardial infarction.

Coronary angiography is currently the method of choice to evaluate the efficacy of reperfusion after myocardial infarction. Although it reflects flow only in the large epicardial vessels and thus provides limited information on the extent of myocardial tissue reperfusion, its impact on the treatment of patients after infarction is well established. Radionuclide scintigraphy, generally used to assess myocardial reperfusion, is limited in terms
of spatial resolution and its dependence on both myocardial blood flow and tracer cellular uptake. The limitations of the present methods are particularly relevant in view of recent reports of incomplete myocardial reperfusion after successful thrombolysis or angioplasty in patients with acute myocardial infarction.

The advent of fast magnetic resonance imaging (MRI) permits the acquisition of MR tomograms during a breath-hold. This creates the possibility of studying the influence of paramagnetic contrast agents on myocardial signal intensity with much greater temporal resolution than achieved before by spin-echo MRI. Previous experimental work using spin-echo MRI identified different contrast enhancement patterns in animals with occlusive versus reperfused infarcts. Our study was designed to examine whether, in patients with acute myocardial infarction, contrast-enhanced fast MRI provides information on myocardial perfusion and myocardial tissue damage additional to that supplied by coronary angiography and thallium scintigraphy.

In patients with acute myocardial infarction, we identified different contrast enhancement patterns that are correlated with epicardial vessel patency defined by coronary angiography and perfusion defects assessed by thallium scintigraphy. The interpretation of these enhancement patterns is based on the knowledge of the fundamental mechanisms of MR contrast enhancement defined in isolated myocardial tissue and experimental models. In addition, we demonstrate that these patterns relate to the severity of acute myocardial damage modulated by attempts to restore coronary blood flow to the infarcted region.

Methods

Study Patients
The study group consisted of 22 patients (16 men) with a mean age of 58 years (range, 40 to 76 years) (Table 1). The entry criteria included typical symptoms of acute myocardial infarction, accompanied by a creatine phosphokinase rise above two times the upper limits of normal, with an at least 5% MB band. All patients gave written informed consent according to the standards established by the Joint Committee for Clinical Investigation of the Johns Hopkins Hospital.

Two-dimensional echocardiography was performed by standard techniques (Hewlett Packard Sonos 1000) in all patients within the first week after infarction. Twelve patients performed a predischarge treadmill exercise test (Naughton protocol), and at peak exercise, a dose of 3 mCi (111 MBq) of \( ^{201} \text{Tl} \) thallium chloride was administered intravenously. Single-photon emission computed tomography (SPECT) thallium scintigraphy was performed according to our standard clinical protocol. Thirty 1-minute projection images over 180° (from 45° right anterior oblique to 225° left posterior oblique) were obtained in a 64x64 matrix by use of a low-energy, all-
purpose collimator and a 35x35-cm field of view. SPECT images were reconstructed by uniformity correction with a 150 million count flood, center-of-rotation correction and low-pass filtering, no attenuation correction, and reconstruction into 1-pixel-thick (5.3-mm) transaxial sections. Oblique angle reorientation and summation produced 3-pixel-thick short-axis left ventricular images. Correction algorithms were applied to the projection images to compensate for motion artifacts.

MRI Protocol
Images were acquired during multiple breath-holds on a 1.5-T whole-body magnet (Signa, General Electric). The pulse sequence used in this study\textsuperscript{19} is similar to an inversion-recovery turboFLASH sequence\textsuperscript{11} in that pixel intensity is heavily T1-weighted. It was specifically designed to minimize contamination of pixel intensity by both T2 effects, by spoiling magnetization in the $xy$ plane\textsuperscript{19} achieved by use of standard radiofrequency phasing algorithms,\textsuperscript{20} and T2* effects, by use of a very short TE (2.3 ms). Briefly, within each RR interval, 60 nonselective dummy radiofrequency pulses are transmitted before imaging to drive magnetization to steady state. These are followed immediately by 32 image phase-encoding steps acquired with TR=6.5 ms, TE=2.3 ms, and flip angle=45°. A total of 96 phase-encoding steps per image were acquired, such that each image was completed in three cardiac cycles. K-space lines 1, 4, 7 . . . etc; 2, 5, 8 . . . etc; and 3, 6, 9 . . . etc were acquired during the first, second, and third beats, respectively. Matrix size was 256x96, field of view was 36 cm, and voxel size was 0.9x3.7x10.0 mm.

After scout images were completed, four base-to-apex short-axis cross sections were acquired with prospective ECG gating during each 12-heartbeat breath-hold every 30 seconds for 5 minutes and then at each minute to complete a 10-minute interval begun immediately before contrast administration. The nonionic contrast agent gadoteridol (Squibb, 0.1 mmol/kg) was administered as a bolus by hand injection in a peripheral vein. The entire examination lasted 45 minutes on average, and there were no untoward reactions to gadoteridol.

MRI Data Analysis
Signal-intensity curves over time were generated with the aid of a commercially available software package (GPIX, General Electric). In brief, regions of interest were defined inside the infarcted region represented as a region of hyperenhanced or hypoenhanced signal in the territory perfused by the infarct-related artery determined by coronary angiography. Regions of interest were also defined inside the noninfarcted territory and left ventricular cavity. In patients with hypoenhanced subendocardial zones at the center of the infarcted region, the central region of interest was defined within the central dark zone, while the other regions of interest were placed within regions of increased signal intensity surrounding the central dark zone. The pulse sequence used in these studies produces dark and homogeneous precontrast cardiac images (Fig 1). Signal intensity from each image was quantified, and the time-intensity curves generated for each patient (Fig 2) were expressed as the percent increase in signal intensity (SI) over baseline precontrast signal intensity as shown below:

**Figure 1.** A. Left ventricular short-axis cross section from a patient with inferior myocardial infarction. Cardiac signal is suppressed to create a homogeneous image to be enhanced by contrast given...
immediately after baseline images were obtained. B, Both ventricular cavities are highlighted at 50 seconds after contrast, showing also delivery of contrast material to normal myocardium and the delineation of a dark region in the left ventricular inferior wall (arrow). C, Images obtained 160 seconds after contrast administration show hypoenhancement of the central portion of the infarcted zone (large arrow) with hyperenhancement of infarcted regions adjacent to the dark zone (small arrows). D, Late images, obtained 10 minutes after contrast injection, demonstrate hyperenhancement of the infarcted region, which appears as bright as the left ventricular cavity (small arrows). Note also a residual hypoenhanced zone in the center of the infarcted region (large arrow).

Figure 2. Time-intensity curves from the patient shown in Fig 1 with recent myocardial infarction. Percent increase in signal intensities in blood (▲) and noninfarcted regions (■) is sharp, with slow decay after contrast. Signal intensity in the periphery of the infarcted region (♦) also rises rapidly after contrast but persists at an elevated level, while blood signal intensity is falling. The center of the infarcted region exhibits reduced signal intensity in the first minutes after contrast administration (●), followed by a progressive rise secondary to delayed contrast penetration into the infarct core.

Normalized SI (%) = (SI−baseline SI/baseline SI)×100.

Normalized time-intensity curves were sampled at three time points (50, 160, and 600 seconds) to describe the time course of myocardial enhancement beyond contrast first pass. The time courses of myocardium-to-blood normalized signal-intensity ratios were analyzed at the same time points. The assessment of the extracellular volume index (ECVi, see "Appendix" for details) was performed in 10 patients, with large areas of signal hyperenhancement uncontaminated by central dark zones. The ECVi was calculated according to the following equation:

$$ECVi = \frac{\Delta SI_{tissue}}{\Delta SI_{blood}} (1 - Hct_{blood})$$
Study Outcomes

The circumferential extent of the sum of hyperenhanced plus hypoenhanced regions was compared with the circumferential extent of the fixed SPECT thallium defect assessed visually from MRI images and thallium scans by two independent investigators. Only the short-axis images obtained at the redistribution phase of the thallium studies, which therefore delineated the fixed thallium defects, were used in this analysis. Both MRI and thallium SPECT abnormalities were quantified at the mid left ventricular wall level and matched by location along the left ventricular long axis. The match was performed by calculating the relative distance of short-axis images from the left ventricular base represented by the mitral valve level. Because only four MRI cross sections were available for seven or eight thallium SPECT cross sections, the thallium short-axis scan that best approximated the MRI short-axis location along the left ventricular long axis was selected for comparison. The average time interval between thallium scintigraphy and MRI was 1 day (range, 0 to 7 days). Thallium scans were interpreted for the presence of fixed or redistribution defects by independent observers who were unaware of the MRI results.

Coronary angiograms were classified by two independent observers in relation to the extent of radiographic contrast penetration downstream from a coronary lesion according to the criteria proposed by the Thrombolysis in Myocardial Infarction (TIMI) study group: grade 0 (no flow), grade 1 (minimal penetration of contrast), grade 2 (delayed flow of contrast), or grade 3 (brisk flow of contrast). Vessels with TIMI grade 0 or 1 were considered occluded and those with grade 2 or 3 were considered patent for statistical analysis purposes. Since 16 of our 22 patients had coronary angioplasty, TIMI flow in the infarct-related artery was assessed both before and after coronary angioplasty. The average time interval between coronary angiography and MRI was 4 days (range, 0 to 10 days).

Echocardiograms were quantified by an independent observer blinded in relation to the MRI and thallium scintigraphic results according to methodology previously described. Global parameters included left ventricular volumes and ejection fraction. The left ventricular wall, defined by endocardial and epicardial contours, was divided into 16 segments by equiangular radial lines placed around the left ventricular cross section. Wall thickness was calculated as the ratio of the segment area to the average of the endocardial and epicardial arc lengths in the short-axis images. Systolic wall thickening was calculated as \[ \frac{(end-\text{systolic} - end-\text{diastolic} \text{ wall thickness})}{end-\text{diastolic} \text{ wall thickness}} \times 100 \]. Regional dysfunction was characterized as the percent circumferential extent of the sum of segments with systolic wall thickening <5%.

ECGs were analyzed by two observers for the presence of Q waves and for infarct location, defined by ECG leads demonstrating Q waves and/or ST-segment changes.

Statistical Analysis

The changes over time of signal intensity obtained from infarcted regions in the MRI scans were compared with the patterns obtained from noninfarcted regions by repeated-measures ANOVA. Differences between specific regions at specific time points were isolated by Bonferroni t tests. Moreover, differences in the myocardium-to-blood signal-intensity ratio between patients with open and closed infarct-related arteries were analyzed by profile analysis with repeated-measures ANOVA. Fisher's exact probability tests, Student's t tests, and linear regression analyses were also used, as indicated in the text. The data are presented as mean±SEM.
Results

Myocardial Infarction and Reperfusion
Therapeutic myocardial reperfusion was attempted in 18 of the 22 patients (see Table 1). Thrombolytics were used in 17 patients, combined with rescue angioplasty in 4 patients. Direct angioplasty was performed in 1 patient. Four patients had contraindications to therapeutic reperfusion and were treated conservatively. Except for the 5 patients who had either rescue or direct angioplasty, all patients had catheterization at least 48 hours after the onset of myocardial infarction. Eleven of these 17 patients had percutaneous coronary angioplasty at that time. The average time interval between coronary angiography and myocardial infarction was 4 days (range, 0 to 11 days).

Myocardial MRI Enhancement Patterns
Three patterns of myocardial signal enhancement on MRI were observed in patients with acute myocardial infarction (see Figs 1 through 3). In noninfarcted regions, a rapid increase in signal intensity reflecting adequate tissue contrast delivery is followed by a slower decay (110±17%, 85±12%, and 57±9% at 50, 160, and 600 seconds after contrast, respectively, Fig 3) caused by the combination of a slowly decaying blood contrast concentration and the extravasation of contrast material into the interstitial space.17 23 24

Figure 3. Graph showing percent increase in myocardial signal intensity (pooled data from all patients) at 50, 160, and 600 seconds after contrast administration. Signal intensity in noninfarcted regions (○) increases rapidly in the first minute, before decaying slowly over the next 9 minutes. Infarcted regions (■) appear hyperenhanced due to rapid signal increase with persistently high signal intensity. In 10 patients, the center of the infarcted region (▲) showed low levels of signal intensity in the first 2 to 3 minutes after the contrast bolus.

The second enhancement pattern is characterized by a similarly sharp rise in myocardial signal intensity, followed, however, by a continued rise in signal intensity over the first 2 minutes and a slower decay than that observed in normal noninfarcted regions (128±21%, 151±26%, and 116±20% at 50, 160, and 600 seconds, ANOVA P<.001 versus noninfarcted, Fig 3). This enhancement pattern is seen as a "bright area" occupying the infarcted region and was found in 21 of the 22 patients. One patient had a normal and uniform enhancement pattern despite a clinical diagnosis of myocardial infarction.

The third pattern, characterized by a slower rate of signal-intensity increase after contrast administration, was present in 10 patients (37±9%, 41±10%, and 39±9% at 50, 160, and 600 seconds, ANOVA P<.02 versus...
noninfarcted, Fig 3). It was seen as a "dark zone" involving the subendocardial half of the left ventricular wall at the center of the infarcted region surrounded by regions of hyperenhanced signal intensity. Eventually, protracted contrast penetration into the infarct core produced signal enhancement in the central dark zones, observed in images obtained 5 to 10 minutes after contrast injection (Figs 1 and 2).

Precontrast absolute signal-intensity levels were not different in noninfarcted versus infarcted regions (59.3±8.3 versus 53.2±7.5 for all patients, \( P=NS \)). In patients with central dark zones, precontrast signal intensity was similar in noninfarcted (56.4±12.6) and infarcted regions, which later appeared hyperenhanced (52.9±12.6) and hypoenhanced (57.8±15.3, \( P=NS \), ANOVA).

Infarct-Related Artery Patency and Myocardial Enhancement Patterns

Infarct-related artery obstruction at the time of cardiac catheterization was related to the presence of subendocardial dark zones by MRI. When the first angiogram obtained after myocardial infarction was analyzed, coronary arteries with TIMI flow 0 or 1 were associated with a high prevalence (75%) of central dark zones by MRI. The prevalence was lower (20%) in patients with patent coronary arteries but with slow antegrade flow (TIMI 2) and was nil in patients with brisk antegrade flow (TIMI 3, \( P<.003 \), Table 2).

View this table: Table 2. Presence of Central Hypoenhanced Regions and Patency of the Infarct-Related Artery

The analysis was repeated taking into consideration the last angiogram obtained before the MRI study, because 16 patients had coronary angioplasty before MRI (Table 2). Four patients who initially had occluded infarct-related arteries (TIMI 1 or 0) and 3 patients previously in the TIMI 2 group crossed over to the group of patients with patent arteries and brisk antegrade flow (TIMI 3) after angioplasty. The prevalence of patients with central dark zones increased in both groups (83% and 36% in patients with TIMI flow 0 or 1 and TIMI flow 3, respectively), thereby reducing the statistical strength of the association between infarct-related artery obstruction and the presence of hypoenhanced regions by MRI (\( P<.056 \), Table 2).

Conversely, the presence of hyperenhanced regions on MRI was not related to patency of the culprit artery by angiography. All patients had either normal or hyperenhanced infarcted regions, indicating that at least the peripheral areas of most human infarcts are perfused by antegrade or collateral blood flow a few days after infarction.

Creatine Phosphokinase, ECG, and Left Ventricular Function

Peak creatine phosphokinase rise was not associated with myocardial perfusion patterns by MRI (1411±309 IU/L for patients with hypoenhanced regions versus 1109±366 IU/L for those without hypoenhanced regions, \( P=NS \)). However, the pattern of myocardial perfusion by contrast-enhanced MRI was related to the presence of Q waves on serial ECGs obtained at the time of infarction. Patients with Q-wave infarcts were more likely
(60%) to have hypoenhanced zones than those with non–Q-wave infarcts (14%, \( P < .07 \), Fisher's exact probability). Ten infarcts were anterior, 10 inferior, and 2 lateral by ECG criteria.

Patients with and without hypoenhanced subendocardial zones within the infarcted regions had similar left ventricular end-diastolic volume (145.3±23.4 versus 130.4±21.6 mL), end-systolic volume (96.1±19.9 versus 68.7±12.2 mL), and ejection fraction (39.7±14.9% versus 46.1±14.9%, respectively, \( P = \text{NS} \)), despite a trend for greater volumes and lower ejection fractions in those with MRI hypoenhanced zones. However, regional dysfunction characterized as a reduction in percent systolic wall thickening <5% was greater in patients with hypoenhanced zones (45.9±8.1%) than those without (22.3±5.6%, \( P < .03 \), Fig 4\textcircled{a}). The relation was still present, although statistically weaker, when the threshold for dysfunction was set at 0% systolic wall thickening (36.8±10.6% with versus 16.0±2.3% without hypoenhanced zones, \( P < .07 \), Fig 4\textcircled{b}).

**Figure 4.** Graph showing percent systolic wall thickening (mean ±SEM) in patients with (△) and without (●) central hypoenhanced zones for 16 segments distributed around the left ventricular short-axis echocardiograms. Wall-thickening maps from individual patients were averaged by superimposing the center of the segments containing the most severe wall thickening abnormalities. Comparisons between patients with and without central hypoenhanced zones were performed by arbitrary selection of the 5% and 0% thresholds (see "Methods").

**Thallium Scintigraphy**

Thallium scintigraphy was obtained in 12 patients during exercise and after tracer redistribution 4 hours later. Ten patients had fixed defects corresponding to the acute infarction, but 2 patients had no scintigraphic evidence of infarction despite a rise in creatine phosphokinase. Eight of the 10 patients with fixed thallium defects also had adjacent redistribution defects, suggesting the presence of myocardial ischemia.

The topographic location of fixed thallium defects correlated well with the location of hyperenhanced regions defined by contrast-enhanced MR images (Fig 5\textcircled{a}). In addition, the circumferential extent of the fixed thallium defect in the redistribution scan correlated well with the total extent of the MRI abnormality (the sum of the hypoenhanced plus hyperenhanced regions) in patients with acute myocardial infarction (Fig 6\textcircled{a}).

**Figure 5.** Top, Left ventricular short-axis single-photon emission computed tomographic thallium scan obtained during tracer redistribution from a patient with left circumflex occlusion who failed thrombolysis but was treated with rescue
percutaneous transluminal coronary angioplasty within 6 hours of the onset of chest pain. Inferolateral fixed thallium defect is seen 5 days after infarction. Bottom, Left ventricular magnetic resonance imaging (MRI) short-axis image matched by location to the thallium scan shown at top. The central zone of reduced signal enhancement seen in the subendocardial half of the left ventricular wall is surrounded by a region of hyperenhanced signal that corresponds in location and extent to the fixed thallium defect obtained the day before the MRI study.

Figure 6. Graph showing circumferential extent of the fixed thallium defect in the short-axis redistribution single-photon emission computed tomographic scan correlates well with the total extent of the magnetic resonance imaging (MRI) perfusion abnormality (the sum of hyperenhanced plus hypoenhanced segments) in patients with myocardial infarction.

Potential Mechanisms of Myocardial Hyperenhancement After Acute Myocardial Infarction

The mechanisms of signal hyperenhancement that characterize infarcted myocardium were explored by study of the time course of signal intensity in myocardium relative to blood after contrast administration. The ratio was not a function of time in noninfarcted regions (Fig 7A). However, in the infarcted regions of patients with occluded infarct-related arteries (TIMI 0 or 1), the ratio increased progressively during the first 10 minutes that followed contrast administration, reflecting the contribution of impaired contrast wash-in/washout kinetics as a mechanism of local myocardial hyperenhancement (Fig 7A). Conversely, in patients with open arteries (TIMI 3), the immediate delivery of contrast material to infarcted territory was followed by a further increase in the ratio during the following 2 minutes, reaching a plateau beyond that time point (Fig 7B).

Figure 7. Graphs. A, The myocardium-to-blood signal-intensity ratio in noninfarcted regions of patients with closed infarct-related arteries...
The upward horizontal displacement of the myocardium-to-blood signal-intensity ratio relative to normal regions suggests the presence of an increased contrast volume of distribution as a mechanism of myocardial hyperenhancement in infarcted territory. This possibility was further explored by plotting myocardial against blood signal intensity for individual images acquired after 2 minutes following contrast administration in 10 patients. Fig 8 shows that all data points obtained from noninfarcted regions lie essentially on the same straight line (myocardial SI = -0.8 + 0.32 blood SI, r = .96). The slope of this relation, which characterizes the volume of distribution for the contrast agent in normal myocardium, principally reflects the extracellular compartment volume. The index of extracellular volume, derived from similar relations for each individual patient (see "Appendix"), was greater in infarcted than in noninfarcted myocardium (45.1 ± 2.9% versus 28.4 ± 1.9%, respectively, P < .001). These results suggest that the contrast volume of distribution is increased in infarcted regions and may contribute to local myocardial hyperenhancement in infarcted but perfused myocardium.

Discussion

Figure 8. Graph showing pooled data from 10 patients with acute myocardial infarction. Δ SI indicates signal-intensity increase in postcontrast images relative to precontrast images but without normalization for baseline signal intensity (see "Methods"). The direct relation between blood (ΔSIblood) and myocardial signal enhancement (ΔSI tissue) in noninfarcted regions is shown (y = -0.8 + 0.32x, r = .96, P < .001). Hct indicates hematocrit.
Contrast-enhanced MRI allows the identification of two perfusion patterns within human infarcts by comparison with the perfusion pattern of normal, noninfarcted regions. Virtually all patients with clinical myocardial infarction had an area of increased signal intensity within the infarcted region in images obtained 2 to 10 minutes after contrast bolus administration. The only patient who did not have hyperenhancement of the infarcted region had a normal perfusion pattern. In addition, the extent of this area of hyperenhanced signal intensity correlated well with the size of the fixed thallium defect, and its presence was not related to the patency of the infarct-related artery. Therefore, our study, by contrast-enhanced MRI, demonstrates the presence of myocardial blood perfusion within all human infarcts a few days after coronary occlusion. This observation is in agreement with prior studies that used pyrophosphate scintigraphy to demonstrate tracer uptake within infarcted territory after permanent coronary occlusion.

The second pattern consisted of a dark hypoenhanced area located in the subendocardium of the infarcted region within 2 minutes of intravenous contrast bolus administration. These regions were surrounded by areas of increased signal intensity and slowly diminished in size over the course of 5 to 8 minutes after contrast injection. They were associated with closed or nearly closed infarct-related arteries at the time of cardiac catheterization, the presence of Q waves on the ECG, and greater segmental dysfunction by echocardiography. They probably result from protracted contrast penetration with or without hemorrhage, probably caused by severe capillary damage and obstruction at the center of the infarcted region.

The presence of flow inhomogeneity inside infarcted territory is well known from experimental studies using radioactive microspheres after coronary occlusion, which have shown a greater reduction in myocardial blood flow in the subendocardial half of the infarct core. In addition, previous clinical studies have shown flow inhomogeneity inside the infarcted region by contrast echocardiography immediately after direct angioplasty. Our study extends the findings of previous work to demonstrate the persistence of flow inhomogeneity several days after coronary occlusion in association with failure to achieve adequate patency of the infarct-related artery after myocardial infarction and with larger regions of myocardial damage by ECG and echocardiographic criteria. Hypoenhanced zones within the infarcted region were also documented in 2 patients with widely patent infarct-related arteries who failed thrombolysis and underwent successful rescue angioplasty (Fig 5). These central dark regions correlate well with "no-reflow" regions, characterized in previous experimental and clinical studies by the failure to achieve complete myocardial reperfusion after reestablishment of blood supply to the infarcted territory.

Prior experimental studies using spin-echo MRI have demonstrated dark zones surrounded by regions of hyperenhanced signal in occlusive infarcts and basically hyperenhanced signal in infarcted but reperfused myocardium. Our findings correlate well with the results of those animal studies in which patency of the infarct-related artery could be accurately controlled. Spin-echo contrast-enhanced MRI techniques have also shown heterogeneous enhancement patterns within infarcted territory in patients with acute infarction. However, those patterns were difficult to interpret because of the limited temporal resolution of the spin-echo imaging sequence used in those studies. Given the time course of regional enhancement patterns after contrast...
bolus administration (Figs 2 and 3), the temporal and topographic features of hypoenhanced zones within the injured territory are best characterized by fast imaging techniques. However, it is theoretically possible to assess the different MRI perfusion patterns documented in our study by conventional or fast spin-echo MRI.

Several previous studies using spin-echo imaging have documented a good correlation between infarct sizes estimated by MRI and by other techniques. Our results are in agreement with those studies that also used the planimetric area encompassed by the MRI signal-intensity abnormality to estimate the extent of ischemic and/or infarcted myocardium. However, because of the trade-off between temporal resolution and number of image planes obtained during a given breath-hold, infarct size was not measured in our study. In addition, in our study, images represent the average of three cardiac cycles with loss of beat-to-beat variations of signal intensity during the contrast bolus first pass through the heart. However, since we did not attempt to extract information on myocardial perfusion from the ascending limb of the time-intensity curves, it is unlikely that the latter factor influenced our results.

Finally, we explored the mechanisms of signal hyperenhancement in infarcted myocardial tissue to gain insight into the pathophysiology of myocardial damage within infarcted regions. Potential mechanisms to explain differences in the time course of the myocardium-to-blood signal-intensity ratio (Fig 7) include (1) impaired myocardial contrast wash-in/washout kinetics, (2) increased contrast volume of distribution, and (3) binding of contrast material to proteins released within injured myocardial tissue.

Our study documents a progressive increase in the infarcted myocardium-to-blood signal-intensity ratio in patients with a totally occluded or nearly occluded infarct-related artery. This finding suggests impairment of contrast wash-in/washout kinetics as a mechanism of myocardial hyperenhancement observed in those patients. Moreover, we estimated the magnitude of extracellular volume expansion in the injured territory of patients with acute myocardial infarction. The almost doubled extracellular space that characterizes human infarcted tissue could represent extracellular edema formation and/or membrane rupture, with consequent diffusion of gadoteridol into the intracellular space. However, although gadoteridol protein binding in normal myocardium is minimal, the possibility of contrast protein binding within infarcted territory cannot be excluded and could potentially contribute to myocardial hyperenhancement in damaged areas after coronary occlusion. Future studies should be directed toward gaining further insight into these mechanisms, which could provide the means to assess myocardial viability after coronary occlusion by MRI.

In conclusion, human myocardial infarcts containing central regions of impaired myocardial blood flow are associated with persistent occlusion of the infarct-related artery and greater regional left ventricular dysfunction. Conversely, infarcts showing uniform contrast hyperenhancement on fast MRI are associated with patent infarct-related arteries and less damage by ECG and echocardiography. Myocardial hyperenhancement by fast MRI results at least in part from impaired contrast kinetics and extracellular space expansion, which may reflect edema formation and/or ruptured cell membranes. Therefore, contrast-enhanced MRI provides unique information on regional myocardial blood perfusion and tissue damage within acute human infarcts.
Figure 9. Hypothetical representation of contrast distribution in blood (left) and myocardial tissue (right) under equilibrium conditions. The extracellular space in this text refers to the sum of the interstitial plus the intravascular volumes minus the volume occupied by red blood cells within the myocardial intravascular space.

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Increased Contrast Volume of Distribution as a Potential Mechanism of Signal Hyperenhancement in Infarcted Myocardium

The interpretation of the first-pass events that follow bolus administration of contrast are complex. The results of this study, however, in which at best only a single breath-hold occurred under first-pass conditions, are somewhat easier to interpret. By 1 minute after contrast administration, which comprises the vast majority of our data, the contrast agent would be expected to be mixed within the body blood pool. In addition, within myocardial tissue, contrast concentrations between the blood and interstitium would be equilibrated on the basis of known extravasation of this agent and molecules of similar size. Furthermore, the rate of contrast clearance from the blood pool by renal elimination and interstitial space diffusion is slow relative to the time required for equilibration between the myocardial blood pool and the interstitial space. These approximations for noninfarcted regions are supported by our results showing myocardial enhancement as a fixed percentage of blood enhancement (Figs 7 and 8). Thus, our data were acquired at a time when myocardial interstitial contrast concentration equals that of the blood.

Fig 9 shows the hypothetical myocardial distribution of the contrast agent under these conditions. Given the above assumptions, the change in pixel intensity in the tissue (ΔSI_{tissue}) due to the contrast agent can be obtained by

\[ \Delta SI_{tissue} = k R_{plasma} [IVV (1 - Hct_{tissue}) + ISV] \]

where IVV is intravascular volume, ISV is interstitial volume, k is the slope of pixel intensity versus 1/T1 for the specific pulse sequence used in this study, R is the relaxivity of the contrast agent as further discussed below, [plasma] is plasma contrast concentration, and Hct_{tissue} is tissue hematocrit. Similarly, for blood:
If we take the ratio of tissue to blood pixel intensity changes, the constants k, R, and \([\text{plasma}]\) cancel:

$$\Delta S I_{\text{tissue}} = \frac{IVV (1 - Hct_{\text{tissue}}) + ISV}{1 - Hct_{\text{blood}}}$$

From this equation, an ECVi can be defined that differs from true extracellular volume in that the volume of red cells within the myocardium is included as cell volume:

$$ECVi = \frac{\Delta S I_{\text{tissue}}}{\Delta S I_{\text{blood}}} (1 - Hct_{\text{blood}})$$

This equation implicitly assumes that (1) myocardial water exchange rates are fast and (2) the imaging pulse sequence results in pixel intensities that are linearly related to $1/T1$. These points are individually addressed below.

First, under these nontransient conditions, it is valid to assume that myocardial water exchange rates are fast and that the change in myocardial $1/T1$ is linearly related to tissue contrast concentration. Thus, the ratio of the change in $1/T1$ of tissue to that of blood is proportional to extracellular volume.

Second, pixel intensities for the pulse sequence used in this study are linearly related to $1/T1$ over the range of 0 to 10 s$^{-1}$. After the first pass of a 0.1 mmol/kg gadoteridol IV bolus, blood and myocardial voxel contrast concentrations would be approximately 0.3 and 0.1 mmol/L, respectively, assuming a myocardial extracellular volume of 30%. For a gadoteridol relaxivity of 4.3 (mmol/L)$^{-1} \cdot $ s$^{-1}$ and assuming fast myocardial water exchange, the change in $1/T1$ of blood and tissue would be about 1.3 and 0.43 s$^{-1}$, respectively. These values are well within the range in which image pixel intensity is linearly related to contrast concentration for our pulse sequence, such that the ratio of the change in tissue to blood pixel intensity is proportional to extracellular volume.

Thus, as shown in Fig 8, all patients have a similar extracellular volume index for normal myocardium and, within the bounds of the above assumptions, the slope of this line reflects the average extracellular space. In infarcted regions, the above arguments equally apply, and the increased index documented in these hyperenhanced regions (see "Results") reflects, at least in part, a local augmentation in the contrast volume of distribution secondary to an increased extracellular space.

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