Coronary artery disease still is the main cause of death worldwide in spite of recent improvements in therapeutic and interventional methods. With the introduction of revascularization therapy, using coronary artery bypass grafting and percutaneous coronary intervention (PCI), the need for identifying viable myocardium has arisen. Furthermore, by preoperatively assessing the transmural size of the infarct, the functional recovery of dysfunctional segments after revascularization therapy in patients with chronic LV dysfunction can be predicted (1-3).

A number of indirect techniques have been introduced to assess myocardial viability. These techniques are echocardiography (for assessing recovery of contractile function), fluoro-2-deoxyglucose (FDG) positron emission...
tomography (PET) (for assessing glucose metabolism) and $^{201}$Tl single-photon emission computed tomography (SPECT) in patients with ischemic heart disease. Unlike echocardiography, magnetic resonance imaging (MRI) does not present the limitations of the acoustic window. MRI also offers more spatial and temporal resolution than nuclear medicine modalities and better tissue characterization. In addition, MRI does not use ionizing radiation, which is another advantage in respect to other imaging modalities such as computed tomography (CT) or nuclear medicine (4).

MR images are acquired with a high contrast between blood pool and myocardium using a steady state free precession (SSFP) sequence. Similar to echocardiography, cine MRI allows dynamic imaging of cardiac wall motion, but with superior endocardial border definition, facilitating more accurate wall motion assessment. Left ventricular function and mass, measured with cine MRI, are used in clinical practice as an endpoint in clinical trials (5,6). End-diastolic wall thickness is easy to measure; however, this indicator does not measure the size of myocardial infarct (MI).

Another method for detecting myocardial viability and critical coronary stenosis is perfusion MRI. With this technique a bolus of contrast media is used to visualize the delay in perfusion in ischemic myocardium compared to healthy myocardium. Perfusion of ischemic myocardium can be assessed quantitatively by calculating perfusion indices (curve upslope, maximum signal intensity and time to the peak). However, clinical studies indicated that MI size is a stronger predictor of future events than LV functional or perfusion parameters, where significant areas of stunning and/or hibernating may be present (7-9).

Unlike echocardiography, PET and SPECT, MRI has the unique ability to provide quantitative information on cardiac function, perfusion and viability. Delayed enhancement MRI (DE-MRI) provides high contrast between viable and nonviable myocardium, thus it has been frequently used for detecting and measuring MI size. Furthermore, discrimination between viable and infarcted myocardium allows patients to avoid the risks associated with revascularization therapy when they are unlikely to benefit. MRI examination for evaluating suspected coronary artery disease consists of T2-weighted imaging for area at risk (AAR) demonstration and DE-MRI for infarct visualization. A recent study using T2 mapping, however, indicated that edematous reaction during the first week after ischemia/reperfusion is not stable and warranted the use of edema extent in evaluating therapies (10). Inversion-recovery prepared T1-weighted gradient-echo (GRE) sequence after intravenous administration of a gadolinium-chelate demonstrates MI, microvascular obstruction (MVO) zone, patchy microinfarct and peri-infarct zone (incomplete infarct at the infarct border). Evidence suggests that the likely mechanism is different volumes of distribution of gadolinium-based MR contrast media in viable and infarcted myocardium (11). It should be noted that an increased volume of distribution occurs in both acute and chronic (scar) MI, as there is an increase in the interstitial space. In the former, the loss of sarcomere membrane integrity increases the potential interstitial volume, whereas in the latter, the presence of fibrotic tissue increases the interstitial space. Additionally, dynamic first pass perfusion and cine SSFP sequences are used for detection of perfusion deficits and LV remodeling. The main disadvantages of the current cardiac MRI sequences are breath-holds, the exam time, contrast media reaction and relatively high cost. New real-time MRI sequences eliminated part of these disadvantages.

### Cardiac MRI

Cardiac MRI is a noninvasive, tomographic, nonionizing technique, thus it has been clinically used for assessing expansion of infarcted segments, late wall thinning of infarcted regions, LV volumes, distortion of LV shape and compensatory hypertrophy of non-infarcted myocardium (12). At UCSF, MRI is used for ischemic, non-ischemic heart diseases, hypertrophic cardiomyopathy as well as heart failure and congenital heart disease (13-16).

MRI pulse sequences are software programs that control the magnitude and timing of the radiofrequency (RF) pulses emitted by the RF transmitter, switching of the magnetic field gradient and data acquisition. The major components of a pulse sequence are the engines and modifiers. The engines include fast spin-echo (FSE), GRE, SSFP, echo-planar imaging (EPI), and single-shot versus segmented modes, while the modifiers include fat suppression (used to null the fat), inversion recovery pulse (used to null the blood and viable myocardium on DE-MRI, saturation-recover preparatory pulses (used with perfusion weighted imaging and cardiac tagging), Phase-contrast imaging with velocity-encoded imaging (for measurement of vascular blood flow, regurgitate fraction and strain) and parallel imaging (applicable to most forms of cardiac imaging). Parallel imaging is a robust method for accelerating the acquisition of MRI data, and has made possible many new applications, such as cardiac MR imaging. Parallel imaging works by acquiring a reduced amount of k-space data with an array of receiver coils. These
undersampled data can be acquired more quickly, but the undersampling leads to aliased images. One of several parallel imaging algorithms can then be used to reconstruct artifact-free images from either the aliased images (SENSE-type reconstruction) or from the undersampled data (GRAPPA-type reconstruction). The advantages of parallel imaging in a clinical setting include faster image acquisition, which can be used, for instance, to shorten breath-hold times resulting in fewer motion-corrupted examinations. Several current applications of parallel imaging are presented and recent advancements and promising research in parallel imaging are well described by Heidemann et al. (17).

**Bright blood imaging**

In general, cardiac MRI sequences can be classified into either black-blood (spin-echo-based acquisitions) or bright-blood (GRE-based acquisition) sequences (Figure 1). Bright blood imaging sequences include GRE and SSFP. GRE sequences are also named spoiled gradient recall (SPGR), turbo FLASH, turbo field echo and fast-field echo (FFE) depending on the manufacturer of the scanner. The SSFP is the sequence of choice at 1.5T because of its high signal-to-noise ratio (SNR), which relies on a steady state of magnetization between the longitudinal and transverse magnetizations (T2/T1 ratio). This imaging sequence produces information on blood flow and cardiac function and has a spatial resolution of approximately 1.5×1.8 mm² and a temporal resolution of 50 ms. The signal in this sequence depends on steady-state T1 signal of blood and myocardium. Bright blood imaging describes the high signal intensity of fast-flowing blood and is typically used to evaluate global and segmental LV function. It has been used for measurements of LV mass, myocardial perfusion, blood flow and coronary anatomy. Investigators recommended cine SSFP sequence as an alternative to conventional echocardiograph.

**Dark blood imaging**

Dark blood imaging sequences refers to the low-signal-intensity appearance of fast-flowing blood and is mainly used to delineate LV chambers. Spin-echo (SE) sequences have been used for dark blood imaging and have relatively little artifact from metal (stents or other devices), but longer acquisition times compared with GRE. FSE dark blood T2-weighted sequences are used (double inversion-recovery, triple inversion-recovery, or short-tau inversion recovery (STIR) in the ventricular short axis. Unlike, conventional SE sequence, FSE and turbo spin-echo (TSE) techniques have lower SNR, but are faster than SE, which minimizes the effects of respiratory and cardiac motion. A potential source of artifact is bright slow-flowing blood adjacent to dysfunctional LV segment (Figure 2). It has been accepted that in acute myocardial infarct (AMI) the hyperintense territory on T2-STIR sequences reflects myocardium at risk (18). T2-mapping sequences have been used for quantitatively assessing myocardial edema (T2 relaxation time) and not qualitatively (arbitrary signal intensity) like T2-STIR sequences (19).

**Dynamic first-pass perfusion imaging**

Myocardial perfusion gives new insight into major and minor coronary artery disease. It can also be used as alternative
surrogate endpoints in studies evaluating revascularization therapy and new therapeutic treatments, such as genes and stem cells. CT, ultrasound, PET and SPECT sequences are designed to obtain kinetic parameters that describe myocardial perfusion and microvascular permeability. Myocardial first pass perfusion sequences are T1-weighted GRE sequences, with a high temporal resolution acquired as gadolinium is being injected. With these sequences it is possible to monitor the contrast as it disseminates throughout the myocardium. Arterial spin-labeling MRI is also a sensitive and useful technique for evaluating myocardial perfusion and angiogenesis. Arterial spin labeling uses flow sensitive alternating inversion recovery MRI technique (20,21) with multiple inversion times and echo planar imaging readout and endogenous tracer (water). The arterial spin labeling technique allows the spin population in arterial water to be magnetically labeled by inversion pulse and used as an endogenous diffusible tracer (21-23). Unlike first-pass perfusion imaging, arterial spin labeling imaging technique has limited use in myocardial perfusion, because of confounding effects of cardiac motion and the relative modest signal changes achievable with spin labeling (24,25).

Unlike arterial spin labeling technique, first pass MRI perfusion uses exogenous tracers (gadolinium-based intravascular and extracellular MR contrast media). Fast perfusion images provide (I) strong T1 contrast; (II) large coverage of myocardial segments; and (III) adequate spatial resolution. Balanced SSFP, GRE and gradient echo-planar (GRE-EPI) are the fastest sequences for perfusion (26).

In multicenter trials, first-pass MRI perfusion demonstrated higher accuracy than SPECT in detecting ischemic myocardium (27). Perfusion sequences are typically more effective at 3T than 1.5T. The 109% higher SNR at 3T (28) can be used to improve spatial or temporal resolution. Peak enhancement is also better at 3T than 1.5T (29). The higher spatial resolutions achieved at 3T may result in decreased size and intensity of the dark rim artifact (Gibbs artifact) at the interface of the LV chamber and myocardium (30), a finding that often is confused with pathologic subendocardial ischemia and is believed to be secondary to low spatial resolution, susceptibility artifact or motion. At 3T, the diagnostic accuracy of myocardial perfusion deficit in coronary artery stenosis is 86% higher than at 1.5T, with good correlation with microspheres (31).

**Figure 3** shows the correspondence between the perfusion deficits on first pass perfusion MRI, wall thinning and dysfunction in the infarcted wall on cine MRI and compensatory hypertrophy in remote myocardium on histochemical triphenyltetrazolium chloride (TTC) stain.

### Delayed contrast enhanced MRI

Detection of myocardial viability is an important issue that needs to be addressed when patients with dysfunctional myocardium are considered for coronary revascularization. Viable myocardium can be detected with nuclear
Techniques (SPECT, PET), low dose dobutamine stress echocardiography and MRI. On MRI, detection of viable myocardium can be accomplished on T1 mapping, DE-MRI, dobutamine stress and stress/rest perfusion imaging.

T1-weighted 2D/3D GRE sequences with an inversion-recovery pulse were used for assessing myocardial viability (32). The inversion pulse is intended to cancel the signal of viable myocardium. New scanners often have a “look-locker (LL)” sequence that provides precise inversion time that nulls myocardial signal intensity at each time point post contrast injection. In infarcted myocardium, the inversion time (200 and 300 ms at 1.5T scanners and slightly higher at 3T) must be adjusted based on the imaging time after contrast media injection due to the clearance of gadolinium from the blood. DE-MRI is usually performed 10-15 min after injecting gadolinium-based MR contrast media. Because of the longer T1 of viable myocardium, the inversion time on 1.5T is shorter (260±30 ms) than 3T (330±48 ms). Several studies have demonstrated SNR and contrast-to-noise ratios (CNR) of up to 3.9 and 3.3 times higher at 3T than 1.5T, respectively (33,34) (Figure 4).

The sensitivity of DE-MRI for detection of AMI and scar is relatively high (99% and 94%, respectively) compared with SPECT (35,36). DE-MRI has the potential to demonstrate MI as small as 1 cm³, which is substantially less than other imaging modalities (26,37-41). Some investigators also suggested that hyperenhanced myocardium on DE-MRI is a reliable parameter to predict true MI size. Early MRI studies in our lab, however, demonstrated that the hyper-enhanced region does not optimally delineate MI (11,42,43), because DE-MRI overestimates AMI by including the edematous border surrounding the infarct (peri-infarct zone). Since the

Figure 3 First pass perfusion (A), cine MR images of diastole (B) and systole (C) and stained LV section (triphenyltetrazolium chloride, TTC, D) show perfusion deficits (arrows, A), lack of wall thickening at systole (arrows, B,C), wall thinning at the site of infarction (arrows, D) and compensatory hypertrophy, respectively, at 5 weeks after infarction. TTC, triphenyltetrazolium chloride.
extent of peri-infarct zone is time dependent, investigators considered it the target of gene and stem cell therapies. Others investigators consider it a source of arrhythmia and poor prognosis (44,45).

The principle of differential AMI on DE-MRI is that the cellular membrane is ruptured and an extracellular contrast media diffuses passively into the intracellular space resulting in increased signal intensity. In chronic myocardial infarctions, scar tissue mainly contains collagenous material; thus, there is an absence of the viable myocardium with increased extracellular space that results in increased contrast concentration and enhancement on DE-MRI (Figures 5, 6). The detection of diffuse myocardial fibrosis is limited on DE-MRI, therefore investigators used myocardial T1 relaxivity for assessing the degree of fibrosis. This technique involves the estimation of the distribution volume of gadolinium in the myocardium based on T1 relaxivity of myocardium relative to that of the blood pool over time. Diffuse myocardial fibrosis has been identified in cases of heart failure, dilated cardiomyopathy and congenital heart disease.

DE-MRI identifies AMI/chronic MI, peri-infarct zone, as well as MVO with high spatial resolution independently from LV function. The transmural extent of hyperenhancement on DE-MRI has been used to predict functional recovery after revascularization in AMI. DE-MRI differentiates non-viable myocardium on the bases of contrast media distribution in infarcted and fibrotic tissues (Figures 5-7). Wu et al. (46) studied 122 patients with STEMI following acute percutaneous revascularization using DE-MRI. The investigators found that AMI size correlated linearly with the initial end systolic volume index, end-diastolic volume index and ejection fraction. Furthermore, Roes et al. (47) found that the size of hyperenhanced infarct on DE-MRI is superior to LVEF and LV volumes for predicting long-term mortality in patients. Masci et al. (48) have demonstrated that the size of hyperenhanced infarct on DE-MRI determines the pattern of infarct healing and regional and
Figure 5 The three fluid compartments in healthy myocardium, namely intravascular (~10% of tissue volume), interstitial (~15%) and intracellular (~75%) compartments.

Figure 6 Schematic presentation of diffuse myocardial fibrosis in non-ischemic heart diseases (A) and contiguous chronic infarct (B) in ischemic heart disease.
global LV remodeling. Currently, DE-MRI is recognized as the standard of reference for assessment of myocardial viability and the size and pattern of hyperenhanced infarct provides important prognostic information on ischemic and nonischemic cardiomyopathy (49).

Coronary revascularization for treating AMI is associated with MVO in the core and peri-infarct zone at the border. Ahmed et al. observed MVO zone in 20-50% of the patients after revascularization of AMI (50). MVO zones can be detected on MRI using three different strategies: first pass perfusion, early enhancement and delayed enhancement (8,51). The last two methods are more specific, because the first method shows the perfusion deficit in infarct with and without MVO zones. Imaging early (1-2 min) after gadolinium-based MR contrast media administration provides valuable information on MVO zone. The dark appearance of MVO zones on T1-weighted imaging is related to lack of delivery of MR contrast media to the infarct core. MVO zones subside on day 9 after infarction.

In an experimental study, it has been demonstrated that hemorrhage occurs within MVO zones (52). Myocardial hemorrhage can be detected on T2 and T2* sequences. On T2 sequences it is identified as a hypointense area within the hyperintense edematous AAR. On T2* sequence the size of the hemorrhagic zone is smaller than MVO zone and is identified as a hypointense area (52). Multiple MR studies have shown that patients with large MVO zone have poor prognosis (51,53). Wu et al. found that the presence of MVO zone in AMI is a marker of poor prognosis (54).

We were the first to demonstrate the peri-infarct zone on DE-MRI in experimental animals (55). The major part of differentially enhanced myocardium on DE-MRI represents MI and the other small portion represent peri-infarct zone (45). Investigators found strong correlation between the extent of peri-infarct zone and ventricular arrhythmias and sudden death (44,45). Both MVO and
peri-infarct zones are visible and can be monitored on DE-MRI. In an electrophysiological study of 40 patients before implantation of pacemakers or implantable automatic defibrillators, the size of the peri-infarct zone on DE-MRI predicted ventricular tachycardia (56). Additionally, DE-MRI has the potential of measuring infarct resorption in animal models (57,58) and humans (59-61).

**T1 mapping**

T1 and T2 relaxation times with and without MR contrast media can be used for mapping myocardium (*Figure 8*) (19,62-64). LL, modified Look-Locker inversion-recovery (MOLLI) and shortened modified Look-Locker inversion-recovery (ShMOLLI) MRI sequences are the sequences of choice for T1 mapping.

However, previous T1 mapping methods, such as MOLLI (65), are based on LL approaches and measure apparent T1 (T1*), which underestimates the true T1 values (65). SMART1 map is a new single-point technique for cardiac T1 mapping. Recently, we compared SMART1 map with MOLLI in an animal model with AMI. SMART1 map yielded higher myocardial T1 values in both normal and infarcted myocardium than MOLLI.

*Figure 9* demonstrates the difference in T1 between MI and viable myocardium on both MOLLI and SMART1. At 1.5T magnetic field strength, the inherent T1 value of healthy myocardium is 1,007±30 ms and longer by 30% at 3T 1,220±70 ms, which are substantially shorter than the blood 1,580±130 ms and 1,660±60 ms, respectively (66). Administration of Gd-DTPA substantially reduces the T1 values of infarcted > blood > normal myocardium, but Gd-DTPA relaxivity is not affected by field strength between 1T and 5T. Longer T1 is better for perfusion, T1 mapping and DE-MRI, because it increase the contrast between normal, hypoperfused and MI.

After injection of gadolinium-based MR contrast media (0.1-0.2 mmol/kg), T1 value of the healthy and
infarcted myocardium differentially decreases followed by continuous increase due to gadolinium washout, with normal myocardium consistently shows a higher T1 than infarcted myocardium or LV cavity (67). Myocardial T1 values that are similar to or lower than the LV cavity blood are indicative of abnormal accumulation of gadolinium in the myocardium (68). It should be noted that contrast enhanced T1 map values are highly dependent on (I) dose of MR contrast media; (II) the time elapsed after contrast media administration; (III) renal clearance of the contrast agent; and (IV) displacement of contrast material by the hematocrit (69). The dose of MR contrast media determines the magnitude of shortening of the T1 relaxation time in myocardium, thus clinical studies should standardize the optimum dose for T1 mapping. It is important to recognize the important of renal clearance of MR contrast media, particularly in patients with chronic renal disease.

**T2 mapping**

The assessment of myocardial salvage in AMI is clinically important because large salvageable tissue is associated with improved short- and long-term survival after AMI. Furthermore, detection of myocardial edema and salvage can allow for the identification of the infarct-related artery, guide management and act as surrogate end points in clinical trials (70). Detection of myocardial edema, using a dark blood TSE technique, has previously been shown to allow early detection of acute coronary syndromes and may identify both the AAR and myocardial salvage post-reperfusion (71,72). T2-weighted MRI identifies myocardial edema before the onset of irreversible ischemic injury and has shown value in risk-stratifying patients with chest pain. However, several problems inherent to T2-weighted MRI including respiratory/cardiac motion, variability in myocardial signal related to surface coil intensity inhomogeneity and sub-endocardial bright signal derived from slow flowing blood (73,74).

A quantitative T2 relaxation map (T2 mapping) has been introduced in 2007 for quantifying myocardial edema (19,26) and patients with MI (75). Kellman et al. found that T2-mapping sequences are superior to T2-weighted sequences

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**Figure 9** T1 maps (3T) showing the difference in T1 of AMI on MOLLI and SMART1 (arrow). The site of infarct corresponds to DE-MRI and histochemical TTC strain. MOLLI, modified Look-Locker inversion-recovery. AMI, acute myocardial infarct; DE-MRI, Delayed enhancement MRI; TTC, triphenyltetrazolium chloride.
in identifying tissues with interstitial edema (26), because T2-weighted sequences do not offer quantitative T2 measurements that would allow for comparisons between different studies (74). Figure 10 shows a T2 map of AMI acquire on a 3T scanner.

T2 mapping leverages the increased myocardial free water in the setting of AMI, taking advantage of the positive relationship between T2 signal intensity and tissue water content. After coronary revascularization, there is plasma accumulation in the interstitial and/or intracellular spaces of ischemic myocardium (76,77). It has been reported that T2 signal hyperintensity may identify myocardial ischemia even before detectable injury by troponin or DE-MRI (72). Recent studies showed myocardial T2 relaxation time to increase not only with AMI, but also with severe transient ischemia (18,72). Others found that myocardial T2 is not only determined by an absolute increase in myocardial water, but also by the movement of water molecules from the extracellular to the intracellular spaces and by the dissociation of water molecules from proteins leading to free instead of bound water (78). Hyperintense myocardial regions on T2-weighted images are thought to present edematous AAR of infarction (79-82). These observations have important clinical implications because patients with AAR of infarction are generally expected to benefit from early diagnosis and prompt intervention to arrest the process of cell death.

Recently, Fernández-Jiménez et al. (10) described in-depth the time course of myocardial edema in a porcine model of 40 min of coronary artery occlusion followed by reperfusion using T2-map and direct measurements of water content. They demonstrated marked fluctuation in myocardial water: increase in myocardial water content at 2 h of revascularization and substantial drop at 24 h followed by gradual increase after 4 to 7 days. The investigators concluded that the edema is not static, but fluctuates. Thus, this study suggests that investigators testing adjunctive therapies to reduce MI size should be careful about assuming that the zone of edema observed on T2-weighted MRI captures AAR of infarction (72). The edematous region fluctuates over time and may be influenced by therapy. Any therapy that reduces ischemic necrosis may reduce the edematous region in the early hours of coronary revascularization, thus falsely reducing the size of the AAR. Measurement of the edematous region at 24 h, when the early edema has resolved, will also falsely lower the AAR, thus, investigators recommended that anti-inflammatory agents should be tested between 4 to 7 days to determine the benefit of therapy.

**Diffusion imaging**

Diffusion MRI provides information regarding microscopic tissue structure through encoding random motion (diffusive) of water molecules. Diffusion tensor imaging (DTI) helps to characterize the myocardium and monitoring the process of LV remodeling after AMI (83). An example of microstructure information of an *ex vivo* healthy pig heart obtained with a DTI experiment is given in Figure 11. The information includes apparent diffusion coefficient (ADC), fractional anisotropy (FA), and primary diffusion direction weighted by FA.

The earliest applications of cardiac DTI were in characterizing healthy myocardium architecture (84-91).
In these studies, the anisotropy property of the diffusion in myocardium was confirmed and the organization of myocardial fiber was observed and verified with histology. Helix angle (HA) maps and fiber tracking of the LV wall showed the transition of the myocardial fibers from right-handed helices to left-handed helices that started in epicardium and ended in endocardium (85-91). Others found during contraction a decrease in subepicardial left-handed helical fibers (LHF) and an increase in subendocardial right-handed helical fibers (RHF) (87). It has also been established that the LV is organized in branching sheet-lets of approximately four-five cardiomyocyte thickness (91-94). The three orthogonal myocytes orientations in the LV are along the local myocyte axis, perpendicular to the local myocyte axis in the sheet-let plane and normal to the sheet-let plane, a structural arrangement known as orthotropy (95). Changes in local myocyte orientation and myolaminar sliding (shearing of adjacent myolaminae over each other) are thought to be the principal mechanisms of ventricular wall thickening in systole (96). The integrity and mobility of these sheet-lets are considered to be fundamental for myocardium function (97). Thus, knowledge of local myocytes and laminar architectures are important in the understanding of normal cardiac function and interpretation of electrical and mechanical studies of cardiac diseases. Investigators also found that myocardial mechanical properties and electrophysiological conductance are different along each orientation (96,98,99). Regarding clinical applications, most of the clinical application studies of DTI are on MI.

Chen et al. observed in an ex vivo rat hearts the increase in mean diffusivity, decrease in diffusion anisotropy and increase in angular deviation of the primary diffusion direction in scar infarct (100). The increase in mean diffusivity was related to increased extracellular space due to loss of membrane integrity and collagen. Another study showed that the decrease in anisotropy and increase in angular deviation were correlated with microscopic fiber disarray, which has also been found to correlate with viability map (83).

In an in vivo swine study, we demonstrated the structural changes in revascularized AMI using DTI. Figure 12 shows hyperintensity area on T2-weighted images and ADC maps. The area mirrors the increase in regional water content related to MI. The hypointensity areas on FA maps corresponded with hyperenhanced infarct on DE-MRI, but smaller than hyperintensity area on T2-weighted images or ADC maps. The areas with increase ADC, but without a decrease in FA, may reflect salvageable myocardium, while the areas with increase ADC and decrease in FA may represent regions with loss of cellular integrity, which are the cause for enhancement in DE-MRI. Histochemical and histopathological stains show the area risk and infarction.

Wu et al. reported infarct zone-dependent changes in ADC, diffusion FA and fiber orientation in patients with 25 days old infarct. ADC data show significant increase, while diffusion anisotropy showed progressive decrease from the infarct and peri-infarct to remote myocardium (83). MI exhibited a substantial loss in the percentage of right-handed helical fibers (% RHF) and an increase in the percentage of left-handed helical fibers (% LHF), while the remote myocardium showed an increase in % RHF and a decrease in % LHF. Furthermore, it was shown that the infarct region with low % LHF correlated well with infarct size.

These aforementioned studies used voxel-by-voxel...
information of the diffusion tensor such as ADC, diffusion anisotropy and HA maps. However, Mekkaoui et al. indicated that microstructural reorganization of remodeled LV after AMI can only be fully understood by looking at myofibers as continuous 3D entities (101). Therefore, they introduced statistical tractography-based HA metrics to yield robust measure of 3D myofiber architecture in normal hearts, where they demonstrated in normal hearts positive and negative distribution of HA in subendocardium and subepicardium, respectively. In remote myocardium, a significant shift toward positive HA was found (83,102).

Beside MI, diffusion-weighted imaging (DWI) was applied in hypertrophic cardiomyopathy (97,103,104). One important feature of this disease is myocardial disarray (103). A study by Tseng et al. showed significant differences in the mean FA and % HA that are less than $-45^\circ$ between normal (0.75, 6%) and hypertrophied myocardium (0.56, 24%) in patients with hypertrophic cardiomyopathy as evidence of myocardial disarray (104). Additionally, the correlation between diffusion anisotropy and myocardial hypokinesis, measured with strain-rate MRI, was stronger in the cross-fiber direction than fiber direction, implying a connection between fiber disarray and myocardial passive compliance. The angle between principal shortening direction and

Figure 12 T2-weighted and ADC maps show the correspondence in delineating myocardial infarct (top block), while the hypointensity area on FA maps corresponds with hyperintensity area on DE-MRI (bottom block), but smaller than the hyperintensity areas on T2-weighted or ADC maps. Histochemical methylene blue and TTC stains delineate the area at risk and infarct, respectively. ADC, apparent diffusion coefficient; FA, fractional anisotropy; DE-MRI, Delayed enhancement MRI; TTC, triphenyltetrazolium chloride.
fiber orientation was reported to significantly vary across hypertrophied myocardium, indicating an abnormal transmural coupling.

In a recent study, Ferreira et al. reported a larger E2A, which is the angle of the second eigenvector of diffusion relative to the local wall tangent plane, in systole than in diastole in control group (97). This finding is consistent with previously reported changes of transmural angulation of the sheet-let structure (97). The same study reported that E2A of hypertrophied hearts retaining similar value during systolic as during diastolic, which implies impaired sheet-let mobility. The abnormal retaining of E2A values was shown to be associate with increased wall thickness but not with the presence of scar infarct on DE-MRI. Evidence was shown to be associate with increased wall thickness but not with the presence of scar infarct on DE-MRI. Evidence of significantly greater contraction in hypertrophied hearts was shown through the increased E2A values in both systole (63.9° vs. 56.4° healthy volunteers) and diastole (46.8° vs. 24° healthy volunteers).

With the increased maturity of acquisition and post-processing techniques, cardiac DTI is gaining momentum and providing more insights to diagnosis and prognosis of myocardial diseases. Cardiac DTI has the potential to characterize structural changes extending beyond infarct areas such as myocyte death (apoptosis), scar formation, myocyte hypertrophy, fiber disarray, angiogenesis, and diffuse fibrosis (100). Thus, with DTI of myocardium, we cross the barrier of pure morphological imaging and move on to microstructural and functional imaging. Furthermore, a clinical study showed that DWI has the potential to detect MI in 78% of patients in less than 1 minute acquisition time, with a similar extent as on DE-MRI (105). Thus, investigators suggested the use DWI in emergency patients with atypical chest pain for confirming/denying the presence of AMI in a few minutes and for differentiating acute from scar infarct. Laissy et al. (105) recently investigated the clinical feasibility of DWI in detecting AMI and to differentiate it from subacute and chronic MI, with delayed contrast enhancement MRI sequence as reference. Furthermore, they measured the variation of the myocardial ADC according to the age of MI. They found that qualitative assessment of DWI compared with delayed contrast enhancement images yielded a sensitivity of 97% and a specificity of 61%/14% to differentiate acute from chronic/subacute MI, respectively. The absolute ADCs (acute =0.00632 ± 0.00037 mm²/s, subacute =0.00639 ± 0.00035 mm²/s, chronic =0.00743 ±0.00056 mm²/s, remote or normal =0.00895 ± 0.00019 mm²/s) and relative ADCs were significantly different between groups (P<0.001) except between AMI and subacute. This study shows that DWI is a sensitive technique to diagnose recent MI. DWI MR Sequences could help differentiate recent from chronic MI.

### Hybrid imaging

Novel trends in hybrid imaging, such as cardiac PET/MRI and optical imaging/MRI, are recently under intensive investigation. For example, it has been shown in patients with chronic ischemic heart disease that FDG PET can differentiate viable myocardium from scar. However, in patients with AMI, FDG-PET is unreliable for the assessment of myocardial viability because reduced glucose uptake has also been observed in reperfused ischemic (stunned or hibernating) myocardium, resulting in overestimation of infarct size (106-109). On PET/MRI, the area of reduced FDG uptake was shown to correspond to the area of myocardial edema in a patient with reperfused AMI, which, in turn, has been demonstrated to delineate the AAR (72,108). Therefore, we hypothesized that reduced myocardial FDG uptake allows for the delineation of the AAR in patients with reperfused AMI. Nensa et al. (110) found in patients with infarcted myocardium and reduced FDG uptake, a good correlation between the area of reduced FDG uptake and the AAR (r=0.70, P=0.001). The area of reduced FDG uptake (31%±11% LV mass) was larger than the infarct size (10%±10%, P<0.0001) and the AAR (17%±13%, P<0.0001). In six out of 18 patients, no late contrast enhancement was seen, whereas all patients had an area of reduced FDG uptake (29%±8%) in the perfusion territory of the culprit artery. They concluded that in patients with reperfused AMI, the area of reduced FDG uptake correlates with the AAR and is localized in the perfusion territory of the culprit artery in the absence of necrosis, although the area of reduced FDG uptake largely overestimates the infarct size and AAR.

### MRI-guided procedures

In the last decade MRI has been extended from a diagnostic to dynamic modality by tracking intravascular guide-wires and catheters in real-time (111). MRI-guided interventions became possible because of major advancements in the speed of data acquisition, data transfer, interactive control-display, highly uniform magnetic fields, rapidly changeable magnetic field gradients, multi-channel receivers and computing systems. Open and closed bore MR scanners have been used for cardiovascular interventions (112).
Open scanners were designed to ease patient access/observation and increase comfort for the interventionists. Multiple vendors have recently introduced hybrid systems (combination of X-ray fluoroscopy or PET with MRI). These systems can be used for detecting and characterizing the disease prior to intervention, provide guidance during intervention and monitor the progress of treatment.

MRI-guidance has been frequently used for stenting of the aorta, pulmonary, coronary, renal, iliac and femoral arteries (113-118). MRI-guided coronary artery stent placement, however, is a challenging interventional procedure because of the small size of the coronary arteries combined with incessant motion during the respiratory and cardiac cycles. These obstacles necessitate higher temporal and spatial resolution for real-time MRI when compared with interventional peripheral MR angiography (118). Bock et al. described in detail the technical prerequisites for MRI-guided endovascular interventions and addressed the safety aspects of this technique (119).

Pre-clinical studies indicated that cell transplantation, delivered under MRI-guidance, is safe and feasible (120-122). Stem cell tracking on MRI is based on labeling injected cells with FDA approved super paramagnetic iron oxide particles. Weeks later, MRI examination showed the increase in collateral blood flow of infarcted myocardium after delivering vascular endothelial growth gene (121). Intramyocardial delivery of gene therapy has been also performed under MRI-guidance (123-125). The advancement of molecular and metabolic imaging will improve diagnostic accuracy and allow patient-specific targeted therapy, designed to maximize disease control and minimize side effects. The advancement of molecular and cellular therapies with MRI-guided targeted procedure will open new avenue in patient-specific targeted therapy in the future.

Future perspectives

MRI guided interventions have also been performed in patients with congenital heart diseases including for the placement of transjugular catheters and cardiac catheterization, as well as repair of aortic coarctation and atrio-septal defects. This minimally invasive technique has been used for closure of atrio-septal defects and local drug delivery (126-129). Currently, there is great interest to develop a catheter based MRI-guided high-intensity focused ultrasound (HIFU) for ablating the sites causing ventricular arrhythmia. The catheter tip creates an ablative energy source (heat) that can precisely penetrate into the target without affecting surrounding tissues. It also provides real-time anatomic and thermal mapping of targeted tissue (130). Clinical studies showed atrial scar on DE-MRI after HIFU (131-133).

Conclusions

It can be concluded that MRI has revolutionized cardiac imaging. MRI gives complementary information on LV function, regional perfusion, angiogenesis, myocardial viability and local orientations of myocytes and their arrays. Recent advances in cardiac imaging include T1 mapping, T2 mapping (for myocardial viability), diffusion imaging (for myocardial architecture) and MR-guided interventions (for delivery of local therapies and ablation). Diffusion MRI provides information regarding microscopic tissue structure, while DTI helps characterizing the myocardium and monitoring the process of LV remodeling after AMI. MR-guided intervention is another young and rapidly evolving field. With the promise of higher spatial-temporal resolution and 3D coverage, in the near future, cardiac MRI will be an indispensable tool in the diagnosis of cardiac diseases, coronary intervention and myocardial therapeutic delivery.

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None.

Footnote

Conflicts of Interest: The authors have no conflicts of interest to declare.

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