Magnetic Resonance and Computed Tomography Imaging of Arrhythmogenic Right Ventricular Dysplasia

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Arrhythmogenic right ventricular dysplasia (ARVD) is a familial cardiomyopathy that causes fibro-fatty replacement of the right ventricle (RV), leading to sudden death due to ventricular arrhythmias. The disease is an important cause of sudden death in individuals younger than 35 years of age. Structural and functional abnormalities of the RV constitute an important diagnostic criterion for the disease. Diagnosis of ARVD is often a challenge as conventional imaging modalities have significant limitations to visualize the RV. Recently, magnetic resonance imaging (MRI) and computed tomographic (CT) imaging have emerged as robust clinical tools for evaluation of myocardial pathology. In addition to providing morphologic and functional information, both imaging modalities have the ability to demonstrate intramyocardial fat, which is the pathological hallmark in ARVD. This article discusses the current status and role of MRI and CT imaging in the diagnosis of ARVD.

Key Words: MRI; CT; right ventricular dysplasia; cardiomyopathy; diagnosis

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strengths, and limitations of MRI and cardiac CT in evaluation of patients with suspected ARVD.

**MRI OF ARVD**

Among the current cardiac MR applications in cardiomyopathies, the greatest potential as well as biggest challenges are in the diagnosis of ARVD. Compared to conventional imaging modalities, MR is uniquely suited to evaluate ARVD. The three-dimensional depiction of anatomy by MRI allows both qualitative and quantitative analysis of RV function (14). MR has the ability to demonstrate intramyocardial fat, which is the pathological hallmark of ARVD (12). The last 10 years have seen significant improvements in MR hardware, with tremendous increases in acquisition speed and image quality. The development of ECG gating and the use of breath-hold imaging have significantly reduced motion artifacts. Improved tissue contrast is currently achieved by the use of inversion recovery black-blood imaging techniques for depiction of cardiac morphology (15). Improvements in cine imaging using ECG-gated steady-state free precession imaging (SSFP) pulse sequence have resulted in better delineation of endocardial borders, enabling accurate and reproducible volumetric measurements (14). For these reasons, MRI has been increasingly used in evaluation of the RV and has evolved as the noninvasive modality of choice in ARVD. Casolo et al (12) were the first to describe the use of MRI to assess ARVD in 1987. They demonstrated intramyocardial fat deposits in the RV on conventional spin echo (SE) imaging in a patient with advanced ARVD. Since that time, several authors, including our group, have reported MR abnormalities in ARVD (16–25). Broadly, MRI abnormalities in ARVD can be grouped into two major categories: 1) morphological abnormalities and 2) functional abnormalities. Morphologic abnormalities include intramyocardial fat deposits, focal wall thinning, wall hypertrophy, trabecular disarray, and RVOT enlargement. Functional abnormalities include regional contraction abnormalities, aneurysms, RV global dilation/dysfunction, and RV diastolic dysfunction. The sites of involvement of these abnormalities are observed in the “triangle of dysplasia,” which is constituted by the inferior subtricuspid area, RV apex, and RV infundibulum (26) (Fig. 1). The goal of MRI in ARVD is to accurately assess the RV for the presence or absence of these abnormalities, which aids not only in the diagnosis but also in follow-up of patients.

**MR Assessment of Cardiac Morphology in ARVD**

Accurate depiction of morphology is very important in most cardiac applications, and ARVD exemplifies this statement. Morphologic evaluation is generally performed by the use of black-blood techniques. Conventional SE pulse sequences were initially used to evaluate cardiac morphology, and the majority of studies in literature have used SE imaging for ARVD. Black blood in the cardiac chambers is obtained by the saturation of inflowing blood signal both above and below the acquired volume. Disadvantages include long acquisition times, precluding breath holding, and resulting motion artifacts, which degrades image quality. Blood suppression is often poor in the long-axis planes and is ineffective for slow flow in the atria and RV.

Currently, black-blood techniques using breath-hold imaging with double-inversion recovery fast SE (DIR-FSE) techniques are preferred to traditional SE imaging. Breathhold FSE sequences consistently provide end-diastolic images with minimal motion artifacts and improve resolution of myocardial detail (27–29). Black-blood inversion-prepared half-Fourier single-shot turbo SE (HASTE) imaging has not been systematically evaluated, but is currently not recommended due to blurring of detail with this sequence. A dedicated cardiac coil is recommended for best results, although we use only the anterior coil elements to prevent wraparound artifact when using a small field of view (FOV). An anterior saturation band (Fig. 2) is placed over the anterior subcutaneous fat for further suppression of motion artifacts.

**Morphologic Features of ARVD**

**Intramyocardial Fat**

Normal myocardium shows an intermediate signal similar to that of skeletal muscle, and fat appears as a hyperintense signal (bright) on black-blood images. Figure 3A shows a black-blood image from a normal volunteer. In normal individuals epicardial fat overlies the RV and is abundant toward the RV apex and in the atrioventricular groove. There is often a clear line of demarcation between the gray RV myocardium and the epicardial fat. Disruption of this line of demarcation (Fig. 3B) and extension of the hyperintense signals into the RV myocardium are frequently noted in ARVD, indicating infiltration of epicardial fat into the RV wall.
The prevalence of intramyocardial hyperintense signals in ARVD on T1-weighted SE imaging has ranged from 22% to 100% in different studies (16–25). The largest series is by Auffermann et al (20), who imaged 36 biopsy-proven ARVD patients and found intramyocardial hyperintense signals in only 22% of patients. The patients in this study had different stages of ARVD, and a significant number had localized ARVD. An interesting finding of this study was that fat infiltration on MRI, not on biopsy, predicted inducibility of ventricular tachycardia at electrophysiologic testing. Wichter (21) added 16 additional patients to the series by Auffermann et al (52 total patients) and concluded that patients with extensive ARVD had higher incidence of fatty replacement of the RV, compared to localized forms (96% vs. 58%). Menghetti et al (25) described SE MRI findings in 15 ARVD patients diagnosed using the Task Force criteria and reported intramyocardial hyperintense signals in 62% of patients. The differences in incidence of fat signal in ARVD are largely based on differences in patient selection and the definition of abnormal intramyocardial hyperintense signals.

We used the breath-hold DIR-FSE technique to evaluate intramyocardial fat in ARVD and found a high intramyocardial T1 signal (fat) in 9 of 12 patients (75%) who were prospectively diagnosed using the current Task Force criteria (16). The use of spectrally selective fat suppression with the DIR-FSE sequence provided additional evidence of fat infiltration due to high contrast between epicardial fat and the RV myocardium. A fat-suppressed sequence may demonstrate a signal void within the myocardium, in a location where a high signal was observed in the non-fat-suppressed image. This increases the confidence for the diagnosis of fat signal, in particular in exams where the results of non-fat-suppressed images are inconclusive (Fig. 4A and B). Fatty infiltration was more commonly noted in the basal regions (RV inflow and outflow) and less frequently at the RV apex (one of nine patients). Biopsy was abnormal in six of the nine patients and the rest had normal biopsies. Biopsies, however, may be inconclusive due to sampling error, since the biopsy site is often the RV septum, which may not be involved in the disease process. Biopsy of the thin anterior RV wall carries a risk of perforation and hemopericardium.

**Wall Thinning**

RV wall thinning is defined focal abrupt reduction in wall thickness of <2 mm, surrounded by regions of normal wall thickness. The exact pathogenesis of wall thinning is not known, but it is thought to be due to apoptotic loss of RV myocytes. Progressive loss of epi-
cardial and myocardial layers leaves behind a thin rim of endocardial cells, which are usually not involved until late in the disease process. This finding often observed in pathologic specimens was not revealed in vivo until the emergence of MRI. Compared to intramyocardial fat, fewer reports have addressed the issue of wall thinning in ARVD. Typically the in-plane resolution of DIR-FSE is approximately 1.5 mm. The resolution is further reduced by motion artifacts. Auffermann et al (20) reported wall thinning in 67% of biopsy-proven ARVD patients on SE MRI. In our series we found wall thinning in less than 25% of patients who met the Task Force criteria (Fig. 5). Wall thickness is often difficult to assess due to adjacent high epicardial fat signal and motion artifacts that are increased in patients with arrhythmias. The use of chemical shift fat suppression may reveal the true thickness of the RV wall, making its identification more reliable.

**Wall Hypertrophy**

Wall hypertrophy is defined as RV wall thickness of >8 mm. This finding is seldom observed in pathologic specimens as the true RV myocardium is measured exclusive of the epicardial fat (30). In vivo this differentiation is sometimes not possible due to extensive fibro-fatty infiltration with loss of distinction between epicardial fat and the true myocardium. In such cases the RV wall appears hypertrophied with MR images showing islands of gray muscle surrounded by bright signals compatible with fat. This finding was observed in 5 of the 12 patients (42%) in our series. Use of fat suppression reveals multiple signal voids within the RV myocardium in locations that showed hyperintense signals in the non-fat-suppressed images (Fig. 6A and B).

**Trabecular Disarray**

Molinari et al (24) were the first to describe giant Y-shaped trabeculae and hypertrophy of the moderator band in patients with ARVD. This finding has been equated to the angiographic finding of deep fissures with a “pile d’assiettes” (stack of plates) appearance. We found a prevalence (40%) of trabecular hypertrophy and disarray similar to that of the above study in ARVD patients (Fig. 7). This finding is not specific for ARVD.
RVOT Enlargement

The RVOT is a common location for localized ARVD. The RV outflow is usually equal to or marginally smaller than the aortic outflow tract at the level of the aortic valve. An exception to this rule is pediatric patients in whom the RVOT may be larger than the left ventricular outflow. Presence of an enlarged RVOT beyond adolescence is uncommon. Ricci et al (22) reported enlarged RVOT in 15 patients with ARVD, compared to patients with dilated cardiomyopathy. In this study, ARVD patients also had increased RV end-diastolic diameters and volumes compatible with RV dilation. This suggests that RVOT enlargement may be a part of generalized dilation involving the right heart. More important than a simple enlargement is a dysmorphic appearance of the outflow tract (Fig. 8). Abnormal appearance of the RVOT, which is dyskinetic in systole, is highly suggestive of ARVD in the absence of pulmonary hypertension.

Figure 6. A: Axial black-blood image from a patient with RV dysplasia showing heterogeneously increased T1 signal in the RV anterior wall (arrow). B: Fat-suppressed image at the same level shows multiple signal voids in the same location of the hyperintense signals on the non-fat-suppressed image (arrow).

Figure 7. Axial black-blood image from a patient with RV dysplasia showing hypertrophied trabeculations (arrowheads).

Figure 8. Axial black-blood image from a patient with RV dysplasia showing enlarged and dysmorphic outflow tract with focal bulging anteriorly (arrow).

and may be present in any condition that results in RV hypertrophy or enlargement.
MR Assessment of Cardiac Function in ARVD

One of the major advances in MR cardiac imaging has been the ability to accurately assess global and regional ventricular function without using geometric assumptions. Until the availability of cine MRI, the complex geometry of the RV made quantification by conventional two-dimensional techniques impossible. The techniques to image cardiac function are called bright-blood techniques, derived from the appearance of intracavitary blood. These techniques yield both morphologic and functional data. Multiple consecutive images that are acquired with a high temporal resolution can be viewed dynamically to generate functional information. Ventricular volumes and masses using bright-blood imaging have been shown to be accurate and reproducible, and MRI is considered the standard of reference (31,32). Although a number of sequences exist for bright-blood imaging, SSFP imaging is the most preferred technique (FIESTA, true FISP, Balanced Fast Field Echo). SSFP sequences result in improved contrast between the blood pool and the myocardium with better delineation of papillary muscles and trabeculation compared to segmented k-space cine gradient echo (GRE) images (33). Blood signal is hyperintense on SSFP images, and this is especially important in imaging the RV. If SSFP is not available, segmented k-space cine gradient echo images (e.g., fast low-angle shot (FLASH), fast cardiac-gated gradient echo (FASTCARD)) can be used. Conventional cine k-space GRE images rely on flowing blood to generate bright blood. In the dysfunctional RV, blood velocities are reduced and signal intensity decreases with conventional GRE imaging. With SSFP, the signal intensity remains high since the signal intensity is proportional to T2 time.

Global functional assessment is useful and can be accurately performed using cine imaging. But it is well known that ejection fraction reduction and ventricular dilation may not be sensitive indicators of myocardial pathology. This is especially true of ARVD where regional functional abnormalities often precede changes in global function. The multiplane nature of MRI, the high contrast between the blood pool and myocardium, and the high spatial resolution have made it possible to visually assess regional RV dysfunction, which has been the focus of several authors in ARVD assessment. There exists an excellent correlation between MRI and RV angiography (17), the later being the traditional gold standard for RV function. For the above reasons, MRI is a noninvasive alternative for RV functional assessment in screening of first-degree relatives for ARVD and also for follow-up.

Functional Abnormalities in ARVD

Global RV Dilation/Dysfunction

Fibro-fatty replacement of the RV in ARVD eventually leads to RV dilation and dysfunction. RV dysfunction is often asymptomatic (34), but is preceded by symptoms related to an associated arrhythmia. MRI does not offer any advantage in patients who have severe reduction in RV function (ejection fraction < 35%), as this can be detected even on echocardiography. The advantage of MRI is to be able to detect minor changes in ventricular volumes over time, which provides insight into an evolving disease process. MR has the capability to accurately and quantitatively measure ventricular volumes, and minor increases in RV end-diastolic volumes may be a subtle and early sign of ARVD. Auffermann et al (20) reported an increased RV end-diastolic volume index in 10 patients with ARVD who were inducible on electrophysiologic studies, compared to control subjects. The RV volume indices and the global function of ARVD patients, who were noninducible, did not differ from those of the control subjects, suggesting that the patients in this group had localized ARVD. Several other authors have reported RV enlargement and dysfunction using a variety of patient selection criteria. In our report we found that a majority of patients (75%) who meet the Task Force criteria have some degree of RV enlargement and dysfunction at presentation. Also, there was a linear correlation between the RV end-diastolic volumes and the duration of symptoms, suggesting the progressive nature of the disease (16). Serial quantitative volumetric assessment of RV may play an important role in assessing disease progression and may have an important role in evaluation of first-degree relatives.

Regional Dysfunction

Much remains to be understood about the pathophysiology of regional dysfunction in ARVD. but regional dysfunction is generally thought to be due to focal fibro-fatty infiltration. Areas of fibrosis result in wall thinning with focally reduced contraction and aneurysm formation. These are thought to precede changes in global ventricular function, and accurate identification may improve the sensitivity of diagnosis. RV angiography was traditionally used to evaluate this, but currently MRI has replaced RV angiography due to its noninvasive nature.

Regional functional abnormalities of the RV described in ARVD include focal hypokinesis (wall thickening of <40%), akinesis (systolic wall thickening of <10%), dyskinesis (myocardial segment, which moves outward in systole), and aneurysms (segments with persistent bulging in diastole and dyskinetic in systole). Studies have consistently reported high incidence of regional dysfunction in ARVD (16–25). One study, which compared MRI to angiography, showed 86% correlation between the two modalities (17). The areas of dysfunction corresponded to the areas of signal abnormality observed on black-blood MRI. In patients presenting with arrhythmias of RV origin, the finding of isolated regional functional abnormalities limited to the RV may not be specific for the diagnosis of ARVD. But the presence of signal abnormality associated with abnormal wall motion is more suggestive of ARVD than either of them alone. In our series, 67% of the patients had regional contraction abnormalities, which correlated to the area of adipose replacement on MRI (Fig. 9A and 9B). Of these patients, 50% had aneurysms localized to the region of adipose replacement. But overall, aneurysm formation was only observed in 25% of the patients with a final diagnosis of ARVD.
Diastolic Dysfunction

Few investigators have used MRI to assess diastolic function in ARVD. Auffermann et al (20) were the first to use time-volume curves obtained from cine GRE MRI to assess diastolic function in biopsy-proven ARVD patients. Comparison with control subjects showed that patients with ARVD who had arrhythmias induced during electrophysiologic testing had a significant delay in diastolic relaxation of the RV. The same patients also had increased RV volumes and reduced function, so that diastolic relaxation may not provide additional diagnostic information. More recently, Kayser et al (35) evaluated diastolic function in 14 patients with ARVD with preserved systolic function using MR velocity mapping of transtricuspid flow. Compared to controls, tricuspid flow patterns in ARVD patients showed a significant decrease in peak filling rate and in the slope of the descending part of the early filling phase. The ratio of peak early filling rate to peak atrial contraction and ratio of integrated early filling to integrated atrial contraction (i.e., volume) were significantly lower in patients than in healthy volunteers. These data are consistent with studies using echocardiography, suggesting that diastolic abnormalities may precede systolic dysfunction and may have a role in early diagnosis.

Role of MRI in Diagnosis of ARVD

The diagnostic role of MRI in ARVD remains somewhat controversial. The ability of MRI to characterize fat in the RV free wall has been brought to question. The sensitivity and specificity of intramyocardial fat on MRI in ARVD remain to be answered. In our experience, this finding alone is neither sensitive nor specific for the diagnosis. Our experience with MRI of autopsy hearts led us to conclude that the achievable spatial resolution in current state-of-the-art clinical protocols substantially limits the capability to detect subtle RV intramyocardial fatty changes. Isolated areas of fat replacement are not specific to ARVD and have been reported in elderly patients, patients receiving long-term steroids, and in other cardiomyopathies (36,37). Discrete areas of fat substitutions have also been reported in idiopathic ventricular tachycardia, which is an important differential diagnosis for ARVD (38–40).

Recently, the reliability of interpreting intramyocardial high T1 signals has been brought to question. Bluemke et al (41) reported poor interreader reproducibility for detection of intramyocardial fat signal on conventional SE MR images performed on patients evaluated for ARVD. The study raises an important issue in defining the role of MRI in ARVD using conventional (i.e., noncardiovascular) MR scanners. The poor agreement for identifying intramyocardial fat is not entirely surprising. The normal presence of epicardial and pericardial fat makes identification of true intramyocardial fat difficult. Some areas such as the subtricuspid region are not easily distinguished from the atrioventricular sulcus, which is rich in fat (42). The RV free wall is only 4–5 mm thick, and the spatial resolution is often unsatisfactory to reliably comment on wall thickness, let alone fat infiltration (43). To differentiate fatty infiltration from the normal epicardial fat requires considerable expertise, and the diagnostic sensitivity and specificity of detecting fat on MRI still need to be defined. Until these issues are resolved, the presence of intramyocardial fat on MRI should not be considered synonymous with ARVD and the diagnosis should not be made in the absence of other clinical criteria.

Since the disease is so rare, most imaging centers have little or no experience with diagnosis of ARVD. Technical problems in imaging patients with arrhythmias, lack of a standardized protocol for ARVD, and lack of experience by imaging physicians suggest that MRI should be only one part of a comprehensive evaluation for these patients. According to the Task Force
criteria (5). MRI provides information related to RV size, global and regional function, and aneurysm formation. Even using conventional SE imaging, the morphologic features appear to distinguish ARVD patients from normal individuals (41).

MRI, on the other hand, can identify patients requiring invasive testing. A completely normal MR study in a patient with no abnormalities on electrocardiography or echocardiography is reassuring, and such patients may not need invasive testing (angiography/biopsy) in the absence of other clinical criteria. If signal abnormalities and wall motion abnormalities coexist, invasive testing should be undertaken to confirm the findings. Minor structural abnormalities, i.e., signal abnormalities in the absence of wall motion changes, present a challenge as further evaluation of such patients is unclear. Adherence to Task Force criteria is recommended, and these minor criteria may not necessitate invasive testing. Further, it must be recognized that ARVD Task Force criteria do not currently recognize fat signal on MR (or CT) as a diagnostic criterion for the disease.

Future Directions

In spite of the current problems faced by MRI, it is undeniable that MR certainly has enormous potential in ARVD. If the significant strides that have been made in MR technology over the last 10 years continue, each of the above-mentioned problems will be systematically addressed and solutions will hopefully emerge.

A current limitation for black-blood imaging as mentioned above is poor spatial resolution. Increases in spatial resolution results in longer and prohibitive breath-hold duration, which is often counterproductive. One approach would be the use of navigator echo-gated techniques, which allow free breathing. This technique was initially described by Ehman and Flemlee (44) in 1989 and is currently used in coronary imaging. Navigator echoes are positioned over the right diaphragm and imaging is triggered for a 4- to 5-mm window. In conjunction with three-dimensional black-blood techniques, this approach could improve spatial resolution for detection of intramyocardial fat. Limitations to this technique include increased scanning times (by a factor of two), increased edge blurring, and susceptibility to motion artifact due to arrhythmias and in patients with irregular breathing patterns.

Another aspect of ARVD evaluation that needs improvement is regional function assessment. Visual analysis is often inadequate due to the complex contraction pattern of the RV. Myocardial tissue tagging provides accurate and quantitative data regarding regional function and may potentially improve the sensitivity of MRI to detect regional dysfunction in ARVD. Fayad et al (45) have previously described this approach in the RV. However, tissue tagging is not easily applied due to the very thin RV free wall and resultant poor signal to noise ratio.

Currently, two large clinical trials in ARVD patients are under way. The European ARVD registry (46) attempts to prospectively validate criteria for clinical diagnosis of ARVD/cardiomypathy (ARVD/C), evaluate the accuracy of clinical diagnosis, and assess the natural course of the disease. The Multidisciplinary Study of Right Ventricular Dysplasia (U.S. ARVD study) (47), which aims to prospectively enroll 100 patients with ARVD and 500 first-degree relatives, attempts to develop quantitative methods to assess RV function to enhance the specificity and sensitivity of the diagnosis of ARVD/C. These studies may define the diagnostic role of MRI in ARVD as well as in family members.

CT OF ARVD

CT imaging utilizing x-ray technology was developed in the early 1970s, and ever since has been used extensively for its ability to provide cross-sectional images of the body (48). There has been a continuous and accelerated development of CT technology, particularly in the last 10 years, focused toward cardiac imaging. The latest-generation CT scanners, namely, electron beam CT (EBCT) and multidetector CT (MDCT), provide excellent spatial resolution and allow accurate high-resolution assessment of morphological detail of both cardiac chambers (49–51). The use of a nonionic contrast agent provides excellent contrast resolution with clear delineation of the ventricular endocardium. Multiple cardiac phases can be extracted, with animated movies of the beating heart made available for visual assessment of global and regional function (49). Quantitative determinations of ventricular mass, right and left ventricular volumes, and global ventricular function can be performed in a variety of cardiac pathologic states (50). The widespread availability of MDCT scanners and less dependence on technical factors have made CT imaging popular compared to MRI in cardiovascular evaluation. In addition, CT is fast, easy to perform, and has more reliable image quality. Although images are acquired only in the axial plane, the acquisition of a three-dimensional data set allows reformatting in any desirable plane. For the above reasons, CT is a clinically valuable, noninvasive tool for assessment of myocardial pathology.

Similar to MRI, CT imaging also has the capability to provide tissue characterization of the myocardium. The ability to depict fatty tissue along with cardiac morphologies makes CT imaging an option for evaluation of ARVD. Dery et al (52) were the first to demonstrate a dilated hypokinetic RV with a markedly thin anterior wall and normal left-sided chambers using CT in an elderly patient with ARVD who presented with RV failure. However, the ability of conventional CT to detect intramyocardial fat in ARVD was first reported by Villa et al (53) in a series of seven patients with ARVD, and subsequently Sotozono et al (54) provided biopsy confirmation of CT findings. They also demonstrated the ability of CT imaging to provide excellent anatomic details of the RV and left ventricle in a patient with advanced ARVD. Since that time, there have been only a few investigators who have used CT to image ARVD, utilizing EBCT and MDCT.

CT Imaging Findings in ARVD

There has been some speculation that CT might be superior to MRI in ARVD due to factors inherent to the
disease process. The arrhythmic nature of the disease often leads to image degradation on MR images, and ARVD patients receive defibrillator hardware, which presently precludes MRI. Hamada et al (55) imaged four ARVD patients with RV arrhythmias who had abnormalities on electrocardiography and angiography using EBCT. With contrast-enhanced volume mode scanning, they were able to demonstrate morphologic abnormalities in ARVD: 1) abundant epicardial fat, 2) low attenuation trabeculations, 3) scalloping of RV free wall, and 4) intramyocardial fat deposits. Quantification of ventricular volumes was performed on cine mode scanning, which showed regional dysfunction and depressed global RV function, respectively. Tada et al (56) added 10 more ARVD patients to the above series and compared EBCT findings to 16 age-matched non-ARVD patients with RV dilation/dysfunction and 13 control subjects. Intramyocardial fat was defined based on tissue attenuation values. The attenuation value for epicardial adipose tissue is around –65 ± 10 Hounsfield units (HU), and 5 to –17 HU for intramyocardial fat, which is far less than that of myocardium. Using the above values, none of the control subjects and no patient without ARVD showed any evidence of intramyocardial fat or any other qualitative features of ARVD, as described by Tada et al (56). The frequencies of abundant epicardial fat, low-attenuation trabeculae, scalloping, and intramyocardial fat in this study were 86%, 71%, 79%, and 50%, respectively. An important finding of this study was that the abnormal area on EBCT corresponded to the areas of abnormality on electroanatomic mapping and was frequently larger than the electroanatomic maps.

Kimura et al (57) studied 32 ARVD patients using contrast-enhanced, nongated, single-row detector helical CT. Similar to the findings of EBCT, they found intramyocardial fat, RV enlargement, hypertrophied trabeculations, and abundant epicardial fat in patients with ARVD. This study also provided radiologic and pathologic correlation in one autopsy heart with ARVD, illustrating the applicability of widely available helical CT in ARVD evaluation. There have been no reports on the use of MDCT in ARVD yet in the literature. Our group studied 13 prospectively diagnosed ARVD patients with MDCT (four detectors), and the results were recently presented in abstract form (58). We found good correlation between RV inflow diameter (measured 1 cm below tricuspid valve) and RV end-diastolic volume, suggesting that this might be a crude surrogate to RV enlargement. Enlargement of the RVOT was also noted in the majority of patients (61%), and this too correlated with increased RV volumes (Fig. 10). Left ventricular involvement with focal wall thinning and fat infiltration of the inferior wall was observed in only one of the 13 patients (Fig. 11).

Current Role of CT in ARVD

The majority of patients who are diagnosed with ARVD receive implantable defibrillators for prevention of sudden death. Defibrillator artifacts do degrade image quality on MRI, and quantification of RV function may be compromised. In this cohort CT remains the only imaging technique to assess RV structure and function for serial morphologic evaluation. CT is also useful in the occasional patient who has frequent premature beats resulting in arrhythmic artifacts on MRI and in patients who are claustrophobic. An additional use of CT imaging is to assess the mediastinum for presence of lymphadenopathy, a marker of sarcoidosis, which occasionally mimics ARVD (59).
Most centers currently rely on MRI instead of CT imaging for evaluating patients with suspected ARVD, mainly because the former is devoid of radiation. MDCT radiation can be quite high, exceeding conventional angiography by a factor of two when performing retrospective gating (60). Thus, MDCT may not be optional for screening of ARVD in first-degree relatives. This would be of particular importance especially in young females in whom breast radiation should be considered. Current temporal resolution of CT is of the order of 105 msec (at a heart rate of 81 beats/minute), but is still suboptimal compared to MRI (61). Despite the above limitations, CT does provide certain advantages over MRI in terms of consistency in image quality, scan time, operator dependency, etc. With increase in familiarity of radiologists with the use of helical CT for ARVD and with the advances in both the temporal and spatial resolution, CT imaging may play an important role both in the diagnosis and in the follow-up of patients with ARVD.

Future Directions
Current multiple-row detector scanners can scan 40-cm-volume lengths in less than 30 seconds with submillimeter near-isotropic resolution and excellent image quality. Higher spatial resolution may allow accurate and reproducible evaluation of intramyocardial fat and RV wall thickness, thereby increasing the sensitivity in ARVD diagnosis. Increasing gantry speeds will improve temporal resolution approaching that of MRI. Accurate calculation of radiation dose with customized adjustment individualized to a patient’s body habitus may increase the utilization of CT in ARVD. Studies comparing MRI and MDCT in ARVD would further our knowledge of the relative utility of each of these technologies in evaluation of ARVD.

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