Review Article

CMR in Heart Failure

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Heart Failure (HF) is a common syndrome with multiple causes. Cardiovascular magnetic resonance (CMR) is a medical imaging technique with significant advantages, allowing the understanding of aetiology and pathophysiology of HF in the individual patient, permitting specific therapy to be administered and predicting prognosis. This paper discusses the diverse role of CMR in HF.

1. Introduction

Heart failure (HF) is a syndrome arising from any cardiac disease severe enough to impair the heart’s ability to support the physiological circulation. Clinically, it is characterised by a number of symptoms and signs most of which relate to fluid accumulation; particularly fatigue, breathlessness, cough, and peripheral oedema. The incidence and mortality of HF is high with more than 24,000 deaths a year in the UK [1] and a 30% mortality within 1 year of first hospital admission [2]. In the United States, 1 in 8.6 death certificates (282,754 deaths/year) mention HF [3]. Management of HF involves prevention, diagnosis of the underlying aetiology, treatment of symptoms, and prevention of disease progression. This may include drug therapy aimed at maladaptive processes within the pathophysiology of heart failure—typically blockade of the renin-angiotensin system, drugs for symptoms (for example diuretics) and nonpharmacological treatment (particularly exercise and dietary changes). Device therapy may also be appropriate (ICD, biventricular pacemaker, ventricular assist devices) and at end stage, cardiac transplantation may be considered. Specific therapy tailored to the etiology of HF is also available. Common causes of HF are coronary artery disease (mainly infarction but with some role for myocardial ischaemia), hypertensive heart disease, valvular heart disease, diabetes, myocarditis, cardiomyopathies, tachycardia-induced cardiomyopathy, and systemic diseases with cardiac manifestations. Within an individual, often more than one etiology is present. Common contributory factors exacerbating symptoms within individuals may include lung disease, obesity, detraining, and simultaneous disease in virtually any other organ system.

The approach to the HF work-up is systematised. Diagnosis of the syndrome is based on history, clinical examination, electrocardiography, chest radiograph, and blood tests (including BNP and tests for heart failure mimics—hypoalbuminaemia, hypothyroidism, and anaemia). Imaging in HF is challenging and requires a multimodality approach. Echocardiography is the first-line investigation for assessment of cardiac anatomy, function, and dynamics. It is cheap, widely available and can be done at the bedside. However, it is limited by poor image quality in some patients and gives very little information about tissue characteristics and coronary anatomy. Cardiovascular magnetic resonance (CMR) provides additional incremental data in patients with HF, providing accurate quantification of systolic dysfunction, determining aetiology and prognosis and predicting response to and selecting therapy. The excellent image quality obtained by CMR allows a detailed examination of cardiac structure and function, without the need for ionising radiation. There are also the key advantages of tissue characterisation (unique to CMR) and perfusion imaging. Evaluation of HF is therefore one of the major current roles of CMR in clinical medicine with the pilot EuroCMR registry of 11,000 patients finding that the most common reason for referral for CMR was to evaluate for
potential myocarditis cardiomyopathy [4]. In this registry, CMR impacted on the management of two thirds of patients and in 16% a new cardiac diagnosis was made that had not been apparent previously.

Assessment of the coronary arteries is currently limited to assessment of origins and proximal courses using CMR. The small calibre of the vessels and movement during the cardiac cycle make flow limiting disease difficult to evaluate using this modality [5]. Cardiac CT or percutaneous angiography are therefore required for anatomical assessment of nearly all patients with potential coronary disease.

This paper provides an overview of the principles of CMR and then discusses its role in specific conditions that can cause HF.

2. The CMR Process

CMR is the use of MRI to image the heart. Any modern MRI scanner can be used, once ECG gating and specific sequences for cardiac work are installed. Scans are typically performed during multiple 7–15 second breath holds, with standardized protocols according to the referral indication [6, 7]. The first 10 minutes of a CMR study is always to assess anatomy and cardiac structure and function—a volume and function study. This typically uses double-oblique imaging orientated to the long and short axes of the heart using steady-state free precession (SSFP) ECG gated cine imaging. The result is a highly reproducible, stereotyped set of images which are acquired in the same way in all CMR centres [8]. At the end of almost any CMR study is late gadolinium enhancement imaging (LGE, see below) looking for myocardial scar from infarction, cardiomyopathy, or myocarditis. In between, additional techniques may be used, including myocardial blood velocities/flow (e.g., for detecting shunts, valve disease), cardiac iron quantification (iron overload), oedema imaging (acute myocarditis, sarcoidosis, infarction), T1-weighted imaging (pericardial disease and masses), tagging (pericardial disease, diastology), real-time imaging during respiration (diaphragm paralysis, pericardial constriction), stress imaging (vasodilator or inotropic stress for ischaemia/hibernation), tumour tissue characterisation, and early postcontrast imaging (thrombus, microvascular obstruction). A typical HF protocol would take 40 minutes. For further details, see Table 1.

3. Gadolinium Enhancement Imaging

CMR is excellent for distinguishing different soft tissues based on their magnetic properties described by the relaxations times T1, T2, and T2*. Normal and abnormal myocardium may have different relaxation times, but the addition of a gadolinium contrast agent greatly magnifies these differences. Gadolinium is chelated and behaves as a passive, extravascular, extracellular contrast agent. After an intravenous bolus, three time phases are considered. The first pass can be used for perfusion imaging and is often given during vasodilator stress to detect ischaemia. Early after contrast, severely hypovascular regions will not enhance (thrombus or microvascular obstruction in acute myocardial infarction). In the late phase (5 minutes plus post bolus), contrast lingers in areas of infarction or focal fibrosis due to slower contrast kinetics and a greater volume of distribution in extracellular water associated with collagen. When imaged using a sensitive sequence (inversion recovery), focal fibrosis appears bright—a region of late gadolinium enhancement (LGE). This can be assessed either visually or quantitatively based on relative enhancement compared to the background [9]. The extent and pattern of LGE varies according to the underlying disease process and is frequently of prognostic significance and can be diagnostic of the underlying etiology.

4. CMR Safety

Sternal wires and all cardiac prosthetic valves and cardiac stents in use in the UK are CMR safe at 1.5 and 3 T. Many intravascular clips are not safe. All ICDs and most pacemakers are not safe with the risk of malfunction and significant heating of the pacemaker leads. There are now CMR conditional pacemakers (generator and leads) available (Figure 1). In the vast majority of patients, gadolinium contrast agents are safe—safer than iodine-based contrast. However, in patients with an estimated glomerular filtration rate of <30 mL/min, only cyclic gadolinium-based contrast agents should be used, and then only after risk-benefit analysis and at the minimum dose because of the possibility of the severe condition, nephrogenic systemic fibrosis (NSF) [10].

5. Practical Clinical Issues

The most common reason for the failure to complete a CMR examination is claustrophobia. In one study, this resulted in 4.2% of patients refusing to undergo CMR [11]. However, with scanning patients feet first, prone, the occasional use of low dose oral (or intra venous) sedation and sensitive treatment, nearly all patients who arrive in a department can be successfully scanned. Some patients with HF may be difficult to scan—either due to inability to lie flat, poor breath holding, or atrial fibrillation with uncontrolled ventricular rates. Real-time, ungated cine imaging, parallel imaging, highly efficient sequence, arrhythmia rejection sequences, and single shot late enhancement sequences help overcome some of these difficulties. On occasions, scanning may be deferred or repeated when patients clinically improve, particularly if the patient is not able to lie flat due to an acute decompensation of their HF.

6. CMR in Myocardial Infarction

6.1. Acute Myocardial Infarction. CMR can be safely carried out in patients following acute infarction and primary angioplasty and aids risk stratification [12–15]. T2-weighted imaging detects myocardial oedema, allows for early diagnosis of myocardial ischaemia, area at risk and salvage [16], and may differentiate between acute and chronic infarction. T2* changes occur in intramyocardial haemorrhage in acute infarction, an adverse prognostic marker [17–20].
Figure 1: A CMR conditional pacemaker. (a, b) show the pacemaker on a chest X-ray. On (b) the arrows point to MRI conditional markers on the header and leads. (c) shows a large artifact from the pacemaker box on a transverse white blood stack. (d) shows susceptibility artifact from the pacemaker leads in the heart.

Figure 2: CMR in a patient with ischaemic cardiomyopathy. The cine imaging had shown normal wall thickness throughout but septal akinesis. (a) First pass perfusion sequence following administration of vasodilator stress with adenosine. A defect is seen throughout the septum. (b) Almost full thickness LGE is seen in the septum highly suggestive that this area is nonviable.
Table 1: Sequences commonly employed during a CMR examination.

<table>
<thead>
<tr>
<th>Sequence</th>
<th>When commonly used</th>
<th>Information obtained</th>
</tr>
</thead>
<tbody>
<tr>
<td>Localisers</td>
<td>All studies</td>
<td>Allows cardiac piloting of all future images. Also allows extra cardiac structures to be assessed</td>
</tr>
<tr>
<td>Transverse Thoracic Stack</td>
<td>All studies</td>
<td>Assessment of extra cardiac thoracic structures</td>
</tr>
<tr>
<td>Cardiac long axis and LV short axis cines</td>
<td>All studies</td>
<td>Assessment of LV and RV volumes, systolic function and regional wall motion abnormality</td>
</tr>
<tr>
<td>Flow</td>
<td>Valve Disease/Shunts</td>
<td>Quantification of Qp: Qs shunt ratio. Quantification of valve regurgitant fraction and forward flow velocity.</td>
</tr>
<tr>
<td>T1 weighted turbo spin echo</td>
<td>ARVC assessment</td>
<td>Demonstration of myocardial fat</td>
</tr>
<tr>
<td>STIR</td>
<td>Acute Myocardial Infarction and Myocarditis</td>
<td>Demonstration of myocardial oedema</td>
</tr>
<tr>
<td>T2*</td>
<td>Iron overload syndromes (e.g., Beta Thalassaemia major)</td>
<td>Demonstration and quantification of myocardial and liver iron</td>
</tr>
<tr>
<td>Gadolinium First Pass Perfusion</td>
<td>Ischaemic Heart Disease, Cardiac mass assessment</td>
<td>Demonstration of perfusion defects in areas of ischaemic myocardium. Assessment of the vascularity of a cardiac mass</td>
</tr>
<tr>
<td>(with and without vasodilator stress)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Gadolinium Enhanced Aortogram</td>
<td>Aortic Disease</td>
<td>Demonstration of dissection/dilatation of the aorta.</td>
</tr>
<tr>
<td>Early Gadolinium Inversion Recovery</td>
<td>Assessment of intracardiac mass</td>
<td>Demonstration of LV or left atrial appendage thrombus</td>
</tr>
<tr>
<td>Late Gadolinium Inversion Recovery</td>
<td>Nearly All Studies</td>
<td>Assessment of LV infarction and viability in ischaemic heart disease. Assessment of fibrosis in Cardiomyopathies.</td>
</tr>
</tbody>
</table>

Figure 3: CMR cine sequences demonstrating the differing morphologies of inherited cardiomyopathies. (a) HCM: asymmetric basal anterospetral wall hypertrophy; (b) LVNC: left ventricular hypertrabeculation and dilatation; (c) FDCM: left ventricular wall thinning and dilatation; (d) ARVC: right ventricular dilatation.
Microvascular obstruction, the tissue equivalent of the no-reflow phenomenon, can be easily detected by CMR. It is associated with adverse ventricular remodelling and clinical outcome [21, 22]. Within areas of microvascular obstruction, myocardial haemorrhage may occur which carries an incremental adverse prognosis [23].

Following the onset of coronary occlusion, irreversible myocardial injury/myocyte necrosis develops as a transmural wavefront which begins in the subendocardium and moves progressively towards the epicardium [24]. LGE CMR can detect even small volumes of focal injury (in the order of grams) and identifies subendocardial scar not otherwise captured by regional contractile function or SPECT imaging [25]. The volume of LGE in acute infarction has been shown to be a strong independent predictor of both heart failure and poor outcomes [26].

The excellent spatial resolution and wide field of view in CMR also allows for identification of the complications of acute infarction including thrombus, true or false aneurysm, papillary muscle involvement, and ventricular septal defect.

6.2. Chronic Myocardial Infarction (Figure 2). The use of CMR in chronic myocardial infarction helps to guide therapeutic and revascularisation decisions. As well as looking at rest perfusion images, imaging can be performed during vasodilator stress to derive flow reserve. Perfusion defects can therefore be observed and the cause of these can be clarified by comparing the LGE images and regional wall motion abnormalities on cine imaging. Low dose dobutamine stress CMR can also be used to assess myocardial contractile reserve.

Left ventricular dysfunction secondary to coronary artery disease typically is either reversible as in states of ischaemia, or potentially irreversible, as in transmural myocardial infarction. As ischaemic cardiomyopathy is potentially reversible, the assessment of myocardial viability is extremely valuable. CMR can assess multiple markers of myocardial viability in a single examination [27].

Myocardial infarction has a characteristic LGE pattern due to the wavefront of myocardial necrosis. This always involves the subendocardium at the core of the infarct, although it may be transmural. Transmurality predicts potential functional recovery but with a grey-zone of 50% transmural infarction recovery with revascularisation it is more difficult to predict [28]. Low dose dobutamine can provide incremental information.

The identification of viable (dark on late gadolinium imaging) myocardium in a patient with regional or global left ventricular systolic dysfunction in the setting of ischaemic heart disease can represent hibernation, stunning, dilated cardiomyopathy, or dyssynchrony [28–30].
myocardium may recover functionally with time (stunning), revascularisation (hibernation), or resynchronisation (see below). In addition, medical therapy (beta-blockers in particular) are more likely to improve cardiac function in the presence of living but dysfunctional myocardium.

7. CMR in Inherited Cardiomyopathy (Figures 3 and 4)

7.1. Hypertrophic Cardiomyopathy (HCM). This autosomal dominant disease resulting in otherwise unexplained left ventricular hypertrophy can present with a number of different clinical phenotypes and in some patients will result in the development of HF and sudden death [31]. CMR is an important investigation in its work-up. High resolution cine images allow accurate hypertrophy assessment, particularly useful in apical hypertrophy which is difficult to visualise using echocardiography [32]. Complex left or right ventricular outflow tract obstruction is well visualised to plan intervention. First pass gadolinium imaging can show perfusion defects whilst LGE sequences show the quantity and distribution of fibrosis, which predicts future HF [33]. In contrast to ischaemic cardiomyopathy, the late enhancement pattern will not follow a coronary distribution and rarely involves the subendocardium. CMR can also identify phenocopies (or genocopies) of HCM, with characteristic findings in Anderson Fabry disease (such as basal inferolateral mid wall late enhancement) and amyloidosis (different gadolinium kinetics with global subendocardial enhancement), both of which may have specific therapeutic options [34].

7.2. Familial Dilated Cardiomyopathy (FDCM). FDCM is characterised by left ventricular (LV) dilatation and impairment on the background of a family history [35]. CMR provides the accurate assessment of LV and right ventricular (RV) volumes, mass, and systolic function. Around 35% of patients with FDCM will have mid myocardial wall LGE, with some evidence that this is a risk factor for sudden cardiac death and ventricular arrhythmia [36]. The pattern of LGE can sometimes be useful in suggesting the underlying genetic basis of FDCM, with dystrophinopathies often showing enhancement of the basal inferolateral segment [37]. It can also help to rule out phenocopies of FDCM, particularly ischaemic cardiomyopathy in older patients.

7.3. Arrhythmogenic Right Ventricular Cardiomyopathy (ARVC). This disease is characterised by fibrofatty infiltration of the right or both ventricles [38]. It may present with sudden cardiac death, although in latter stages can cause biventricular systolic impairment and hence HF. It is a challenging disease to diagnose as assessment of the RV by echocardiography is difficult and limited due to its complex and variable geometry. The multiple image planes that can be performed during CMR make RV assessment less unreliable. Recently published international task force guidelines emphasise assessment of regional wall motion abnormality and RV volumes and ejection fraction [39]. Although not mentioned in the task force criteria, CMR can also be used to identify areas of myocardial fibrofatty infiltration [40].

7.4. Left Ventricular Noncompaction (LVNC). This disease is characterised by the presence of a thick endocardium with prominent trabeculation and deep recesses communicating with the LV cavity [41]. In some cases it can result in HF via LV systolic impairment. Noncontrast-enhanced echocardiography is limited in the evaluation of LVNC as the LV apex is often poorly visualised. Cine CMR...
Figure 6: Gadolinium-enhanced imaging in specific cardiac diseases. (a) Subendocardial LGE in a patient with AL amyloidosis. (b) Mid anterior wall focal LGE lesions in a patient with sarcoidosis. (c) Epicardial and mid wall anterior LGE in a patient with myocarditis. (d) Subendocardial enhancement in a patient with Churg-Strauss syndrome.

imaging will provide excellent views of the LV apical endocardium, with early and late gadolinium contrast imaging allowing evaluation for thrombus. However at present LVNC is a complicated and often debateable diagnosis particularly due to wide normal variation and there is no consensus on diagnostic criteria [42].

8. CMR in Intracardiac Shunting (Figure 5) and Other Congenital Heart Disease

Heart failure is the final common pathway for many types of congenital heart disease. CMR can assist in the morphological diagnosis, functional assessment, angiography, flow and tissue characterisation, monitoring ventricular function, and tracking change over time. Flow analysis allows shunt detection—a ratio of >1.5 is considered significant and likely to result in patient symptoms (including those of heart failure) at some stage [43]. Cine imaging can then be used to demonstrate the shunt anatomy including ASDs, VSDs, anomalous pulmonary venous drainage, and more complex disease [44]. CMR is the gold standard for investigating complex disease, and its use has dramatically reduced the need for invasive angiographic assessment [45]. Its primary limitation is that direct measurement of pressures cannot be made and so pulmonary vascular resistance cannot be directly assessed.

9. CMR in Pericardial Disease

Pericardial disease, particularly pericardial restriction can cause HF in the presence of preserved ejection fraction but impairing cardiac filling [46]. CMR can detect the anatomy of pericardial thickening; the local and systemic functional consequences (e.g., tethering of the pericardium to myocardium, effusions), and the nonquantitative haemodynamics of pericardial constriction with ventricular-ventricular interaction either at rest or on real-time cine imaging during inspiration (where preferential filling of one ventricle due to respiration restricts the other, with a clear septal deviation associated) [47–49]. The assessment of pericardial calcification is better performed using CT scanning.

10. CMR in Valvular Heart Disease

Echocardiography is the first-line test for valvular heart disease, providing excellent assessment of haemodynamics and anatomy in most patients. CMR has incremental value, particularly where echocardiographic windows are poor [50].

Cine imaging will allow precise evaluation of ventricular size and function which can be important in deciding when to surgically intervene in valvular disease [51]. It will also provide high-quality imaging of the valve anatomy allowing
accurate planimetry of the aortic valve to be performed. In aortic valve disease, cine imaging and gadolinium enhanced aortography will show the anatomy of the aorta in three dimensions. CMR is also very useful for assessment of pulmonary valve disease which can be difficult with echocardiography.

Flow sequences will allow the calculation of the regurgitant fraction in regurgitant valve disease and the forward flow maximal velocity in stenotic disease [52, 53].

11. CMR in Specific Acquired Cardiomyopathies (Figure 6)

11.1. Sarcoidosis. In the thorax, sarcoidosis usually manifests with disease in the lungs, lymph nodes, or heart. Autopsy evidence has shown that cardiac involvement can occur in up to 30% of cases [54]. CMR assessment can show extracardiac disease. If cardiac involvement is present, T2-weighted sequences can highlight areas of acute inflammation. LGE sequences will show enhancement in areas of disease, which is usually multifocal and in the mid or subepicardial layers [55]. In patients who require endomyocardial biopsy to make the diagnosis, there is some evidence that targeting areas with LGE increases the diagnostic yield of the procedure [56].

11.2. Myocarditis. This condition is caused by inflammation of the myocardium, most commonly as result of infection [57]. It can progress and result in a dilated cardiomyopathy and hence HF [58]. T2-weighted CMR sequences will show areas of acute inflammation. LGE will identify areas of necrosis or severe oedema. This is most commonly seen in the septal or basal inferolateral wall in the mid or subepicardial layers [59]. Myocarditis is a common finding on CMR in patients presenting with chest pain and an ECG/biomarkers suggesting myocardial infarction, but a normal coronary angiogram. One recent study found that 60% of patients presenting in this manner had myocarditis [60]. Such patients may not have any regional wall motion abnormality and so tissue characterisation using CMR is the only noninvasive method for making the diagnosis.

11.3. Iron Overload. The most common cause of iron overload cardiomyopathy is repeated blood transfusions in patients with beta thalassemia major. This results in a 50% mortality in patients under the age of 35 years [61]. CMR allows the accurate quantification of cardiac iron levels using the T2 star method [17]. This allows identification of patients who are at risk of developing HF, allowing more aggressive iron chelation therapy to be administered.

11.4. Cardiac Amyloidosis. Cardiac involvement is common in patients with AL or TTR amyloidosis. The presence of amyloid protein results in accelerated removal of gadolinium from the blood and increased myocardial uptake, such that more of it is present in the myocardium than the blood pool [62]. This change in gadolinium kinetic behaviour is almost unique to amyloidosis. The LGE pattern in amyloidosis is often subendocardial and seen in a circular pattern throughout the LV. This is in keeping with histological evidence demonstrating where cardiac amyloid deposition occurs [63].

11.5. Eosinophilic Diseases. Cardiac hypereosinophilia (e.g., malignant, Loefflers; Churgg-Strauss) causes endomyocardial fibrosis, valve disease and papillary muscle dysfunction, diastolic dysfunction, intracardiac thrombus formation, and heart failure. These are highly characteristic and well diagnosed and followed by CMR using T2-weighted imaging, early and late gadolinium techniques [64, 65].

11.6. Chagas Disease. Endemic in South America, the chronic phase may result in heart failure [66] characterised by myocardial fibrosis, sometimes mimicking infarction. CMR LGE is common and the extent of LGE correlates with disease severity [67].

12. CMR and Biventricular Pacing

Cardiac resynchronisation therapy (CRT) using biventricular pacing is a commonly used treatment for patients with HF and limiting symptom [68]. Upto 1/3 of patients fail to respond despite stratification by the ECG QRS duration. Results using echocardiographic markers of dyssynchrony have been mixed [69]. CMR cine and tagging sequences are being investigated [70], but one promising marker is ventricular scar burden and distribution. A high scar burden, predicts nonresponse, and even more robustly, posterior scar (the location of the LV lead) virtually precludes response [71]. It may also be possible by CMR to determine coronary sinus LV branch anatomy to target the LV lead away from scar areas [72].

13. Conclusions

The incidence of HF remains high throughout the world. CMR using the methods described above, is establishing itself as the gold standard non invasive method for phenotyping this condition, providing important insight into both the aetiology and pathophysiology of this syndrome in individual patients. In future years we expect that CMR will provide us with continuing improvements in image quality. Many new sequences are likely to become available which for example will improve LGE imaging [73] and allow assessment of diffuse myocardial fibrosis [74]; a variable that at present can only be assessed by invasive biopsy. Most importantly, we envisage CMR becoming more widely available, allowing a greater number of patients with HF to undergo this important investigation.

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