Cardiovascular magnetic resonance imaging – a pictorial review

Introduction

Cardiovascular magnetic resonance imaging (CMR) is a powerful problem-solving tool and arguably offers the most comprehensive assessment of cardiac morphology and function, as well as the opportunity of rebuilding the bridge between cardiologists and radiologists. The role of CMR-trained imaging physicists is also valuable, and many CMR centres harmoniously incorporate these three sub-specialty fields.

This paper comprises an overview of several CMR techniques, outlining both the strengths and limitations of the modality.

Cardiovascular magnetic resonance imaging

The combinations of cardiac and respiratory motion make the heart a challenging organ to image. Electrocardiogram (ECG) gating is done almost without exception, and patients are often asked to hold their breath for each scanning sequence to minimise respiratory motion artefacts. In prospective gating, the acquisition is triggered by the QRS complex, and usually only 90% of the cardiac cycle is acquired. In retrospective gating, data are acquired continuously while the ECG trace is recorded, and processing is done retrospectively to provide images throughout the cardiac cycle.

MRI is a relatively slow modality, and samples need to be taken from a number of heartbeats. Image acquisition is made more time-efficient by capturing interleaved samples or slices during each heartbeat. The resulting samples from different phases of the cardiac cycle can be represented as a multi-slice collection of images, or interleaved into a cinematographical series of images.

Figures

- Fig. 1a. Four-chamber T2-weighted image showing evidence of active inflammation (arrow) in a patient with myocarditis. (1b.) Four-chamber bright-blood image showing a mediastinal neoplasm, post-operatively confirmed to be a benign thymoma (block arrow) impressing the right atrium (arrow) but not invading the heart. (1c.) Short axis post-contrast T1 image showing diffuse RV infiltration, confirmed on catheter directed myocardial biopsy to represent primary cardiac lymphoma, bulging into and severely narrowing the RV outflow tract (arrow).

- Fig. 2a. Bright-blood image showing thrombus (arrow) in a typical site with the left atrium. (2b.) Dark-blood image in the same patient as in (a) confirming thrombus (arrow) within the left atrium. CMR was used in this case following equivocal echocardiography for confirmation of thrombus and to differentiate this from an atrial myxoma. (2c.) Extensive mural thrombus (arrows) is shown in a patient with previous infero-septal myocardial infarction with aneurysm formation.
monitoring the liver-lung interface using navigator echoes, which are incorporated in some MRI pulse sequences. A number of fast imaging techniques have been developed, including steady state free precession (SSFP) and echo planar imaging (EPI). Details of these sequences can be found in many MRI texts.1 A specialised phased-array radiofrequency chest coil is used for cardiac imaging to maximise the MRI signal. If multiple phased-array coils are used, parallel imaging2,3 can be applied to significantly reduce the scan time, with only a marginal trade-off in signal.

Techniques and clinical applications

Morphology

Various image sequences can be used, either rendering ‘dark blood imaging’ where the blood pool is darker than the myocardium, or ‘bright blood imaging’ where the myocardium is darker than the blood pool. Cardiac imaging planes (short axis, long axis and four-chamber orientations) are traditionally used. Standard imaging planes (axial, coronal and sagittal) are usually also employed to ensure that the extracardiac structures are assessed.

CMR is used with excellent effect in patients with suspected cardiomyopathies, myocardial inflammation, tumours and infiltration (Fig. 1). Both the right ventricle (RV) and left ventricle (LV) can be viewed in their entirety.

Intracardiac thrombus can also definitively be identified and differentiated from neoplasms (e.g. atrial myxoma) with CMR as a problem-solving tool following equivocal echocardiography (Fig. 2).

Myocardial iron deposits associated with transfusion-dependent anaemias can be quantified using T2* mapping. This allows pre-clinical monitoring of iron deposition with immense therapeutic implication (iron deposition can be reversed), as the first clinical manifestation is often intractable heart failure.4

CMR also offers the most complete evaluation of the pericardium. This allows the distinction between constrictive pericarditis and restrictive cardiomyopathy, which can both be clinically and haemodynamically indistinguishable (Fig. 3).

General chamber orientation and morphology can be evaluated as well as abnormal communications (atrial septal defect, ventricular septal defect and patent ductus arteriosus), often applied to congenital cardiac disease.

Function

Both the right and left ventricular parameters can be evaluated with excellent reproducibility. A series of semi-automatically defined contours are used to obtain estimates of ventricular volume and myocardial mass. Evaluation of RV function is important in patients with suspected arrhythmogenic right ventricular cardiomyopathy (ARVC) and in therapeutic follow-up of patients after repair of certain congenital heart diseases, and cannot be reliably measured on any other non-invasive modality.

Stress tests can be performed where controlled doses of dobutamine can be administered to investigate wall motion abnormalities under...
stress conditions. This is, however, less accurate than perfusion stress CMR (see Perfusion below).

Gradient echo cine images are also useful for valvular imaging and have a niche application since slow moving or turbulent blood results in low signal intensity (Fig. 4). These can be quantified with phase contrast (PC) velocity encoding, which provides an instantaneous measure of fluid velocity, and can be used to provide velocity-time and flow-time curves through the valves and the great vessels, thus allowing calculation of pressure gradients. Flow can also be measured in structures not easily or accurately assessed by echocardiography e.g. post-valvar pulmonary stenosis and aortic coarctation (Fig. 5). This technique can also be applied to the myocardium, yielding regional measures of velocity and strain rate.

Three-dimensional (3D) phase contrast velocity encoding is also being used as an emerging technique for creating high-resolution angiograms without the need for contrast agents.

MR angiography

3-D contrast angiography can be performed in conjunction with CMR to fully evaluate the great vessels, and is particularly useful in patients
with congenital heart disease where intracardiac and extra-cardiac abnormalities often co-exist (e.g. tetralogy of Fallot with associated post-valvar pulmonary stenosis and aortic coarctation with associated VSD). Fig. 6 shows a post-contrast MR angiogram maximum intensity projection (MIP) image of a patient with Scimitar syndrome.

**Viability and delayed enhancement imaging**

MRI viability imaging is unparalleled by any other techniques. Gadolinium contrast agents settle in oedematous areas and take longer to be washed out of scar tissue. Delayed-enhancement MRI involves imaging the heart 10 minutes after administering the intravenous contrast using inversion recovery sequences that null normal myocardial tissue. Scar tissue appears unmistakeably bright in these images. This technique can map the severity and extent of myocardial infarction, with obvious prognostic implications (Fig. 7).

This technique is also increasingly being used to differentiate ischaemic from non-ischaemic cardiomyopathy. Typical patterns of delayed enhancement have been described for various non-ischaemic cardiomyopathies, allowing non-invasive diagnosis and prognostication (Fig. 8).

**Perfusion**

Myocardial perfusion can be quantified by rapidly measuring the uptake of gadolinium. A cine image series with manually introduced contours is
post-processed to yield regional measures of tissue perfusion. In contrast with the longer-term gadolinium uptake, ischaemic myocardium appears darker than normal myocardium in the first images owing to reduced perfusion. This can be incorporated into a stress test but requires the use of Adenosine, which is unfortunately not currently licensed for this application in South Africa.

Coronary angiography

Although MR coronary angiography has been steadily improving, this arguably remains the one shortcoming of CMR. Resolution lags behind multidetector CT, and techniques such as volume imaging with respiratory navigator tagging are not as robust.

Myocardial tagging and DENSE

In MRI tagging, the myocardium is modulated by a series of saturated dark bands or tags. These are a material property of the tissue and can be seen to deform as the heart moves. Tagged images provide insight into contractility and myocardial mechanics, and can be processed to yield meaningful measures of myocardial strain. Displacement encoding with stimulated echoes (DENSE) is a more recent technique that provides more accurate measures of displacement and hence myocardial strain. Both myocardial tagging and DENSE allow a user to track discrete portions of tissue as they traverse the cardiac cycle. Fig. 9 shows an example of this technique, where a number of points in the left ventricle are tracked through the cardiac cycle.

Myocardial dysynchrony can be mapped by using tagging or DENSE regional strain-time curves. Dysynchronous but healthy segments of myocardium can be identified using a variety of metrics such as the time to onset of strain or the time to peak strain. This technique can be used in combination with delayed-contrast MRI to improve the planning of cardiac resynchronisation therapy. Fig. 10 presents a patient with a right bundle branch block and an anteroseptal myocardial infarct. Fig. 10a is an end-systolic displacement and strain map obtained using cine DENSE imaging. The black lines represent motion relative to end-diastole, and the underlying colour represents tissue deformation corresponding to circumferential shortening. The infarct is clearly apparent as a region of zero or slightly positive strain. The positive strain occurs when infarcted tissue is stretched by the adjacent healthy tissue. Further information can be obtained by looking at regional strain as a function of time. The plots in Fig. 10b show strain for the anterior septum and regions in the right and left ventricular free walls. A delay in onset to contraction is clearly apparent in the RV segment, but the peak strain is normal; this implies that pacing the RV free wall would benefit this patient but pacing the anterior septum would yield no benefit.

Conclusion

Just as MRI has made and continues to make a major impact on neuroscience, sports medicine, orthopaedics and various other fields, it is currently also making a major impact in cardiology. CMR is established as the gold standard imaging technique for assessing myocardial and pericardial pathology. It is the most accurate and reproducible non-invasive investigation for ventricular function and viability assessment. CMR is also a valuable problem-solving adjunct to echocardiography in congenital and valvular heart disease. Inability to depict detailed coronary artery anatomy is the major shortcoming of CMR, but physicists continue to develop new and faster methods of imaging to overcome this factor and further improve established applications.