The role of cardiac magnetic resonance imaging in the assessment of heart failure with preserved ejection fraction

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Heart failure (HF) is a major cause of morbidity and mortality worldwide. Current classifications of HF categorize patients with a left ventricular ejection fraction of 50% or greater as HF with preserved ejection fraction or HfPEF. Echocardiography is the first line imaging modality in assessing diastolic function given its practicality, low cost and the utilization of Doppler imaging. However, the last decade has seen cardiac magnetic resonance (CMR) emerge as a valuable test for the sometimes challenging diagnosis of HfPEF. The unique ability of CMR for myocardial tissue characterization coupled with high resolution imaging provides additional information to echocardiography that may help in phenotyping HfPEF and provide prognostication for patients with HF. The precision and accuracy of CMR underlies its use in clinical trials for the assessment of novel and repurposed drugs in HfPEF. Importantly, CMR has powerful diagnostic utility in differentiating acquired and inherited heart muscle diseases presenting as HfPEF such as Fabry disease and amyloidosis with specific treatment options to reverse or halt disease progression. This state of the art review will outline established CMR techniques such as transmitral velocities and strain imaging of the left ventricle and left atrium in assessing diastolic function and their clinical application to HfPEF. Furthermore, it will include a discussion on novel methods and future developments such as stress CMR and MR spectroscopy to assess myocardial energetics, which show promise in unraveling the mechanisms behind HfPEF that may provide targets for much needed therapeutic interventions.

KEYWORDS
HfPEF, CMR in HfPEF, diastolic function, diastolic dysfunction, diastolic heart failure
Introduction

Heart failure (HF) is a clinical syndrome caused by abnormalities in cardiac structure and function resulting in increased intracardiac pressures and/or reduced cardiac output (1). Patients with HF frequently present with dyspnea, fatigue and fluid retention, organ dysfunction due to hypoperfusion and have a higher risk of sudden cardiac death due to ventricular arrhythmia or pump failure. Although the range of pathologies resulting in impaired cardiac function is broad, current clinical guidance on the classification and management of HF places great emphasis upon left ventricular ejection fraction (LVEF). Thus, patients with HF and LVEF of 50% or greater are commonly categorized as having HF with preserved ejection fraction (HFpEF). The prevalence of HF is estimated at 64.3 million globally (2) with more than half of these people having HFpEF (3). In this review, we discuss the utility of cardiovascular magnetic resonance imaging (CMR) for the assessment of HFpEF.

In patients presenting with HFpEF, the pathological hallmarks are abnormal left ventricular (LV) filling and increased LV end diastolic pressures (LVEDP), collectively termed diastolic dysfunction. HFpEF has diverse causes associated with a multitude of co-morbidities. Established risk factors implicated in the development of HFpEF include hypertension, type 2 diabetes (T2D), chronic kidney disease and obesity. HFpEF also appears to be more prevalent in females and presents at an older age than heart failure with reduced ejection fraction (HFrEF). Importantly, atrial fibrillation (AF) has a complex association with HFpEF (4) and AF commonly coexists with HFpEF with a reported prevalence of up to 65% in older patients (5). The underlying pathological changes are thought to be related to chronic inflammation, neurohormonal activation, changes in intracellular signaling pathways, endothelial and microvascular dysfunction and myocardial fibrosis (6).

To date, a number of diagnostic algorithms have been proposed since accurate identification of HFpEF can be problematic (7). Importantly, a wide range of HFpEF clinical phenocopies also exist, each with underlying pathophysiologically distinct etiologies including hypertrophic cardiomyopathy (HCM), ATTR cardiac amyloidosis, Fabry disease iron overload cardiomyopathy and cardiac sarcoidosis. Moreover, unique disease-targeted therapies in these conditions might reverse, halt or slow disease progression.

Diagnosing HFpEF with CMR

Cardiac imaging plays a pivotal role in the diagnosis of HFpEF. Echocardiography is the initial modality of choice to assess LV diastolic function particularly because of its availability, cost-effectiveness and technical capabilities through Doppler imaging. However, CMR is increasingly being utilized for the further evaluation of patients with HFpEF (Figure 1). CMR is currently second line to echocardiography in the imaging assessment of diastolic function as it remains a limited resource, is less economical and is challenging in patients with claustrophobia. Practical limitations in HFpEF patients include difficulties in breath holding in older patients and in obese patients. Image quality may be impacted in patients with AF and in those with cardiac devices and post valvular intervention. In addition, there remains a larger clinical and research evidence base for echocardiographic assessment of HFpEF. Despite these drawbacks, the strengths of CMR include its higher spatial resolution and ability to quantify structural changes in the heart with greater precision and reproducibility compared to other imaging modalities, and its unique tissue characterization capability (8). The diagnostic utility of CMR is highlighted by a study comparing CMR and echocardiography. Kanagala et al. (9) found that CMR diagnosed new, significant clinical cardiac pathologies such as myocardial infarction, HCM and constrictive pericarditis in 27% of 154 patients with HFpEF and these patients were at higher risk of death and HF hospitalization at 6 months. Recent data further suggests integration of different CMR measures of diastolic function into a novel diagnostic algorithm (10) in a similar vein to echocardiography but further validation and consensus agreement is required before putting such algorithms into routine clinical practice.

LV functional assessment

CMR is the current gold standard imaging technique to assess LV volumes and therefore LVEF, and LV mass, as it avoids the geometric assumptions made by echocardiography and provides superior reproducibility (11, 12). Advances in CMR machine learning further serve to increase repeatability for measuring LV volumes and mass (13). To provide the highest spatial and temporal resolution, ECG-gated bright-blood balanced steady state free precession (SSFP) sequences of contiguous short-axis slices from the LV base to apex are acquired (14, 15). In patients with HF, cardiac remodeling and left ventricular hypertrophy (LVH) are associated with impairment of myocardial contractility and diastolic dysfunction even with preserved LV ejection fraction (16). LVH is an established risk factor for adverse cardiovascular events (17–19) and is associated with incident HF events (20).

Phase contrast CMR is routinely used to determine flow for valvular assessment using through-plane velocity encoding. These sequences can also be utilized to measure transmural inflow, generating E (passive diastolic inflow) and A (active diastolic inflow) waves and pulmonary vein inflow, which correlate well with echocardiographic Doppler indices (21–23). However, CMR-derived values tend to be lower and may underestimate velocities compared to echocardiography, which
FIGURE 1
Approaches to the assessment of HFrEF by CMR. CMR, cardiovascular magnetic resonance; HFrEF, heart failure with preserved ejection fraction; LA, left atrial; LV, left ventricular; LVH, left ventricular hypertrophy; RV, right ventricular.

may relate to the lower temporal resolution with CMR (30–40 ms) vs. echocardiography (< 5 ms) (24, 25). Additional pitfalls of CMR include time consuming data acquisition and analysis in addition to positive or negative phase offset errors with through-plane flow imaging due to local non-compensated eddy currents (26). The recently developed CMR golden-angle method permits acquisition of 150 to 250 frames per cardiac cycle to match that of echocardiography (27). This novel method involves k-space lines acquired continuously determined by the golden-angle of each sector, together with alternating velocity-encoding signs (27). Advances in CMR sequences have led to the development of 3D and 4D flow sequences which may prove to be useful for a more advanced assessment of diastolic filling than measurement of mitral inflow and pulmonary vein inflow by phase contrast CMR alone. For example, a 3D velocity-encoded MRI with retrospective mitral valve annular plane tracking sequence had better agreement with Doppler echocardiography for LV diastolic filling patterns compared to a 2D one-directional velocity-encoded sequence (28). Furthermore, in 53 healthy volunteers, a comparison of 4D flow CMR kinetic energy to mitral inflow E/A ratio showed a stronger independent association with age than standard 2D metrics (29).

Tissue Doppler imaging (TDI) for the quantification of myocardial velocities is a fundamental, clinically validated method for the assessment of diastolic function by echocardiography (30, 31). Similarly, phase contrast CMR can be used to determine myocardial e’ velocities (peak modal velocity in early diastole by pulsed TDI waveform at the mitral valve annulus). CMR-derived mean e’ and E/e’ have consistently shown excellent correlation with echocardiographic values in patients with diastolic dysfunction (23, 32). In a small observational study of patients with hypertensive heart disease (n = 18), CMR E/e’ strongly correlated with invasively measured mean pulmonary capillary wedge pressure (PCWP) (r = 0.8, p < 0.0001) and had a 100% positive predictive value for E/e’ < 8 and PCWP ≤ 15 mmHg, and similarly so for E/e’ > 15 and PCWP > 15 mmHg (32). Measurement of CMR LV fractional area change during the first 30% of diastole (termed diastolic-index) in the short axis view correlated well with e’ on echocardiography (33).

LV strain

Although the use of LVEF is clinically the dominant imaging method for defining severity, subtype and progression of HF, it has limitations (34). Other imaging parameters, particularly LV strain may detect changes in myocardial architecture and
function before changes in LVEF occur (35). Strain is a measure of myocardial mechanics that describes the deformation of LV myocardial fibers which are orientated in the longitudinal, circumferential and radial directions. Changes in longitudinal function occur early on in HF (36). Accordingly, global longitudinal strain (GLS) has been identified as an important marker of early myocardial dysfunction (37). Indeed, GLS has proven to have powerful prognostic value superior to LVEF (38). In a systematic review involving 5,721 patients with cardiovascular disease (CVD), there was stronger independent association with mortality with each SD change in the absolute value of baseline GLS (HR 0.50, 95% CI 0.36–0.69; p < 0.002) compared to LVEF (HR 0.81, 95% CI 0.72–0.92; p = 0.572) (38). Different methods to measure strain by CMR have evolved from myocardial tagging and phase contrast velocity-encoding to feature tracking, the latter now being the more commonly employed sequence.

Strain measurement by myocardial tagging using Spatial Modulation of Magnetization (SPAMM) involves tagging orthogonally intersecting sets of lines marking rectangular grids in 2D image (39). In 1,500 participants from the MESA (Multi-Ethnic Study of Atherosclerosis) cohort, diastolic function from circumferential strain curves independently predicted incident cardiovascular disease (CVD), there was stronger independent association with mortality with each SD change in the absolute value of baseline GLS (HR 0.50, 95% CI 0.36–0.69; p < 0.002) compared to LVEF (HR 0.81, 95% CI 0.72–0.92; p = 0.572) (38). Different methods to measure strain by CMR have evolved from myocardial tagging and phase contrast velocity-encoding to feature tracking, the latter now being the more commonly employed sequence.

An alternative method to assess myocardial deformation is by phase contrast velocity-encoding which offers better spatial resolution but lower temporal resolution than myocardial tagging (45). This allows measures of instantaneous velocity over a short time period. Displacement encoding with stimulated echoes (DENSE) is a free-breathing, phase-velocity based method which permits measurement of displacement during most but not all of the cardiac cycle (46). Unfortunately, this limits the assessment of diastolic tissue displacement and acquired images have low signal to noise ratio.

Feature tracking (FT) CMR analysis (FT-CMR) tracks points or features in the myocardium across successive imaging frames over the whole cardiac cycle, generating strain values and curves (Figure 2). FT-CMR is increasingly favored for the assessment of strain due its close correlation with echocardiographic speckle tracking (47) and myocardial tagging (48, 49). Moreover, the ability to utilize routine SSFP-based cine images simplifies clinical workflows. In a small study of 18 HFrEF patients compared to 18 age and sex-matched controls, GLS independently predicted abnormal relaxation index, Tau, by invasively measured pressure-volume loops (50). GLS measured by FT-CMR proved to be a powerful independent predictor of all-cause mortality in a multi-center study of 1,012 patients with HFrEF and a median follow-up of 4.4 years (51). In a study of 131 patients with HFrEF, GLS ≥ −8% by FT-CMR independently predicted HF hospitalization and cardiovascular death at 2.5 years follow up (52). Novel applications of FT-CMR include the assessment of LV torsion, rotation and diastolic recoil (53). Interestingly, different rotational mechanics are found in patients with amyloidosis and HCM, conditions which often masquerade as HFrEF (54).

Left atrial size

Changes in left atrial size (LA) are a hallmark of elevated LV filling pressures in patients with HFrEF (55). Maximal LA volume measured at end-systole in sinus rhythm and indexed to body surface area (BSA) has strong predictive value for adverse cardiovascular events (56). CMR provides a more accurate measurement of LA size compared to echocardiography, owing to inherently superior spatial resolution. LA size can be measured by contouring the LA in the short axis stack or, more commonly, by the area-length method or Simpson’s biplane in the 4-chamber and 2-chamber views. Although Simpson’s biplane is the reference standard to assess LA size by echocardiography, there is limited consensus on the preferred method by CMR (57, 58). Nevertheless, in a multicenter study of nearly 11,000 subjects with a median follow up of 4 years, increased BSA-indexed LA size measured by CMR was independently associated with all-cause mortality (59).

Left atrial function

LA function can be subdivided into its reservoir (LA filling during LV systole), conduit (passive LV filling in early to mid-diastole) and pumping (LA contraction to augment LV filling in late diastole) phases. LA conduit-filling capacity is calculated by maximum LA volume minus pre-atrial contraction volume whilst LA active filling capacity is calculated by pre-atrial contraction volume minus the minimum LA volume (60). LA stroke volume is defined as maximum minus minimum LA volume. Furthermore, LA ejection fraction (LAEF) can be derived from these measured volumes with both biplane and short axis methods showing good agreement in sinus rhythm (61). Left atrial ejection fraction is closely associated with LVEDP on cardiac catheterization (62).
Increased LA volumes, reduced LAEF, reduced LA reservoir and booster pump strains are all associated with diastolic dysfunction as well as its severity (63). In the multi-ethnic population-based Dallas Heart Study of 1,802 patients, lower LAEF was independently associated with increased mortality [hazard ratio per 1 SD (8.0%): 1.56 (1.32–1.87)] with superior and incremental predictive value over maximum LA volume index (64). These findings were further supported by the MESA study of 336 patients with T2D, in whom incident cardiovascular disease was strongly and independently associated with lower passive, active and total LAEF (65). Kanagala et al. demonstrated that LAEF is reduced in HFrEF (n = 140) compared to controls (n = 48) and that a lower LAEF was associated with an increased risk of all-cause mortality or first HF hospitalization (log-rank, all p < 0.028; sinus p = 0.036) (61, 66). Furthermore, the strong association with adverse outcomes was similar for LAEF derived by either the biplane or short axis methods during CMR (61).

**Left atrial strain**

Application of CMR-FT to the LA generates strain data that have been evaluated as measures of LA function (67). Chirinos et al. (68) compared patients with HFrEF (n = 101), HFrEF (n = 120) and without HF (n = 640) demonstrating that conduit and reservoir LA strain measured using CMR-FT independently predicted risk of incident HF admission or mortality. In the MESA cohort, incident HF was predicted by lower longitudinal atrial strain (25 ± 11% vs. 38 ± 16%; p < 0.001) and lower LA emptying fraction (40 ± 11 vs. 48 ± 9%; p < 0.001) at baseline (69). In a small study of 22 HFrEF patients compared to healthy controls, LA conduit strain was significantly reduced in HFrEF and was associated with impaired oxygen uptake (VO2 max) during cardiopulmonary exercise testing and invasive measurements of impaired early LV filling (70).

**Right ventricle**

CMR is also the gold standard non-invasive method to assess right ventricular (RV) size and function. In a prospective, observational study the prevalence of RV dysfunction (RVEF < 47%) as determined by CMR was present in 19% of individuals with HFrEF (n = 135) (71). Furthermore, RV dysfunction was independently associated with death and HF hospitalization (adjusted HR 3.946, 95% CI 1.878–8.290, p = 0.0001). HFrEF is a recognized cause of elevated pulmonary artery pressures (PAP) and pulmonary hypertension. Resultant changes in the right heart readily assessed by CMR include increased right atrial size, RV hypertrophy and septal bowing. Assessment of diastolic dysfunction must take into account an estimation of PAP which is typically elevated in patients with HFrEF. Echocardiography can estimate systolic PAP by measurement of the peak tricuspid regurgitant (TR) velocity using the formula PAP = 4∗(TRV_max)2. While measurement of the tricuspid regurgitant jet peak velocity is readily assessed by echocardiography, this is less readily performed by CMR. However, systolic PAP can be indirectly estimated by identifying the peak TR velocity using phase contrast CMR flow analysis at the level of the tricuspid valve (72).

**Myocardial tissue characterization**

In HFrEF, alterations in the extracellular matrix with increased collagen deposition are thought to be a result of inflammation and increased oxidative stress (73). This process leads to myocardial fibrosis, which may contribute to the impaired relaxation that is observed in HFrEF. CMR has the unique capability to detect both focal replacement myocardial fibrosis using late gadolinium enhancement (LGE) and diffuse interstitial fibrosis through parametric mapping sequences (native T1 and extra cellular volume (ECV) quantification).
The presence of replacement fibrosis by LGE in patients at risk of HfP EF including in AF (74) and diabetes (75) has been shown to increase the risk of mortality. Furthermore, identifying LGE can provide additional risk stratification for patients at risk for hospitalization for HF regardless of etiology or LV systolic dysfunction (76). In an observational cohort study of 1,916 patients with AF, LA fibrosis by LGE was associated with an overall incidence of developing HF at 3.1% per year (77). Moreover, 80% of patients developed HfP EF (n = 63) rather than HFrEF (n = 20) after a median 2.7 years follow-up and that the incidence of HF increased with increasing LA fibrosis.

T1 measures the time taken for longitudinal relaxation of excited protons to return to equilibrium following application of a radiofrequency pulse. Colored maps can be generated so that pixel values represent the T1 in each voxel. T1 values in the myocardium and blood pool acquired pre- and post-contrast, and by accounting for hematocrit, can be used to calculate ECV. ECV quantifies the relative expansion of the extracellular matrix acting as a validated surrogate imaging biomarker for myocardial fibrosis (78, 79).

Both native T1 mapping and ECV are helpful in detecting inherited and acquired cardiomyopathies, particularly Fabry disease and cardiac amyloidosis (Figure 3). Diffuse myocardial fibrosis by native T1 mapping and ECV appears to be associated with diastolic dysfunction and LV stiffness in HfP EF (n = 62) but not in HFrEF (n = 40) or healthy controls (n = 22) (80). ECV has been demonstrated to correlate with invasive measures of load-independent passive LV stiffness (81, 82). In a study comparing patients with HfP EF, hypertension and healthy controls, ECV was significantly increased in HfP EF (35.9 ± 5.0%) compared to both hypertensive patients (31.9 ± 5.2%) and healthy controls (27.0 ± 4.3%) (83). Moreover, ECV was superior to GLS in differentiating between HfP EF and hypertension.

ECV may have a unique role to play in identifying HfP EF phenotypes at higher risk of cardiovascular events and death (84). In a cohort of 410 patients at risk for or diagnosed with HfP EF, ECV correlated with BNP levels and outcomes of heart failure hospitalization or death (85). This may indicate that the degree of myocardial fibrosis occurs in a continuum of severity and may precede overt clinical features of HfP EF. These findings are corroborated by other studies demonstrating that a lower post-contrast T1 time (median <388 ms) is a strong predictor of adverse events (86). Furthermore, both focal and diffuse myocardial fibrosis was noted to be more prevalent in HfP EF (n = 140), compared to age and sex matched control subjects. ECV indexed to body surface area correlated with LV mass:volume ratio, RV end-diastolic volume index and maximum LA volume index and independently predicted adverse outcomes (87). Interestingly, a recent study demonstrated that anterior RV insertion point fibrosis measured by increased native T1 times significantly correlated with markers of increased LV end-diastolic pressures and filling (88).

**Stress perfusion and exercise stress CMR**

Patients with HfP EF may only develop dyspnea on exertion and have limited indices of diastolic dysfunction at rest. The gold standard for diagnosing the effects of HfP EF during exertion is invasive right heart catheterization during exercise (89), although it is more often assessed indirectly by exercise stress echocardiography (90). Recent studies have shown a role for stress perfusion or CMR combined with exercise to increase the diagnostic yield for HfP EF (91–93). In particular, there appears to be significant coronary microvascular dysfunction in HfP EF patients as revealed by myocardial perfusion reserve following stress perfusion imaging (94). Stress perfusion CMR identifies patients at higher risk of major cardiovascular events in HfP EF without known coronary artery disease (95, 96).

In a feasibility study, exercise stress combined with real-time CMR was able to detect HFpEF, confirmed by right heart catheterization, with high accuracy (93). The authors also demonstrated that LA longitudinal shortening was the most accurate parameter to detect HFpEF. In future, exercise stress CMR may play a prominent role alongside exercise echocardiography in the workup of HFpEF patients, without the need for invasive tests.

**Myocardial energetics**

The high energy requirement of the heart and minimal capacity to store energy suggests that an imbalance of energy supply and demand may predispose to the development of myocardial dysfunction. MR spectroscopy has the capability to assess myocardial energetics by measuring phosphocreatine to adenosine-triphosphate (PCr/ATP) ratio, and cardiac steatosis by measuring myocardial triglyceride content (MTG). In a small study of 12 patients with T2D and diastolic dysfunction, there were reductions in PCr/ATP ratio when compared to controls (97). Burrage et al. (98) investigated patients with a spectrum of diastolic dysfunction including T2D (n = 9), HfP EF (n = 14) and cardiac amyloidosis (n = 9) and controls (n = 11). Across the spectrum of HfP EF, there was a decrease in PCr/ATP ratio in parallel to increases in E/e′, NT-proBNP and lower LV diastolic filling rates with low workload exercise. Moreover, patients with HfP EF and cardiac amyloidosis had transient pulmonary congestion with exercise as revealed by pulmonary proton density mapping. In HfP EF, significantly greater MTG and therefore myocardial steatosis correlates with reductions in CMR measured diastolic strain and VO2max (99).

Recent clinical trials have identified beneficial effects of therapeutics on cardiac metabolism. Evidence has been growing for the sodium–glucose cotransporter 2 (SGLT2) inhibitor, empagliflozin, for the treatment of HfP EF (100). The mechanism of these benefits have not been fully elucidated.
but an improvement in cardiac metabolism has been shown following in a significant increase in PCR/ATP in a small study of 18 patients with T2D compared to 10 healthy volunteers (101). The authors also found an increase in mean LVEF of 7% and a 3% increase in GLS.

Epicardial adipose tissue (EAT) surrounding myocardium and within the pericardium is metabolically active providing energy to myocardium through the breakdown of triglycerides and also generates pro-inflammatory mediators (102). Furthermore, EAT may affect mechanical properties of ventricular function. In obese HFpEF (n = 99) compared to non-obese HFpEF (n = 71), there was significantly higher total epicardial heart volume [945 ml (831–1,105 ml) vs. 797 ml (643–979 ml) and 632 ml (517–768 ml); p < 0.0001] and EAT thickness (10 ± 2 vs. 7 ± 2 and 6 ± 2 mm; p < 0.0001) measured by CMR (103). Furthermore, the larger epicardial heart volume in obese HFpEF was associated with increased pericardial restraint and ventricular interdependence.

### Novel methods in assessing HFpEF by CMR

Recent advances in CMR have generated novel sequences and measures to assess HFpEF. Diastolic dysfunction results from altered LV compliance leading to changes in LV filling patterns. LV time-volume and peak filling rate curves generated using CMR acquired LV volumetric datasets associate with the severity of echocardiographic-derived diastolic dysfunction (104). Patients with HFpEF have a lower peak filling rate adjusted for end diastolic volume (105), prolonged peak filling rates (106) and greater diastolic volume recovery (proportion of diastole required for recovery of 80% of stroke volume) by CMR (107).

A fundamental aspect of HFpEF assessment is to determine PAP. Vortices of blood, measured by 3D or 4D phase-contrast CMR, appear in the main pulmonary artery in pulmonary hypertension (108, 109). Measurement of the duration of blood vortices allows estimation of pulmonary artery pressures (110). A vortex duration ≥15% corresponds to an invasive mean PAP ≥25 mmHg. Using 4D flow analysis there is good correlation to Doppler echocardiographic estimates with potentially higher diagnostic yields in the detection of raised pulmonary artery pressures (111).

Central transit times have long been known to be increased in HF (112). Using first pass perfusion, Cao and colleagues (113) demonstrated that global central transit times from the right atrium to aorta were significantly prolonged in HFpEF, which correlated with increased PCWP. Measurement of central transit time may therefore act as an additional marker of HFpEF for patients undergoing CMR though data on its use remains limited.

Experimental CMR sequences in the pipeline may further serve to characterize the myocardium and alterations in diastolic function potentially adding to the diagnostic role of CMR in the assessment of HFpEF. Examples include diffusion tensor imaging (114), MR elastography (115), artificial intelligence and radiomics (116), which are beyond the scope of this review article.

### Conclusion

Assessment by CMR enables refinement of a clinical HFpEF diagnosis into underlying cardiac conditions such as...
HCM and cardiac amyloidosis. Furthermore, the strengths of CMR over other imaging modalities include the capability for sub-categorization into pathophysiologically sub-types such as increased myocardial fibrosis, left atrial dysfunction or microvascular dysfunction. The comprehensive evaluation of HFpEF by CMR may enable risk profiling of patients with HFpEF and perhaps allow focused, targeted therapies in the future. Advances in CMR sequences and postprocessing are generating novel indices of diastolic dysfunction which not only aid diagnosis but may hold prognostic risk prediction. The role of CMR in assessment of diastolic dysfunction continues to evolve at a rapid rate but large-scale studies are still required to permit technical reproducibility as well as clinical validation across different populations and subcategories of HFpEF. Moreover, future research could focus on the challenge of identifying which established and novel indices should be routinely incorporated into clinical workflows. It remains to be determined how such advances compare to a predominant echocardiographic based approach to clinical diagnosis and management of HFpEF, especially whether the use of CMR can lead to improved prognostication and outcomes.

Author contributions

CL drafted the manuscript, performed literature review, and created figures. ME created a figure and reviewed the manuscript. PK and JK critically appraised the manuscript and made constructive modifications and suggestions. JA coded the topic for the manuscript, critically appraised the manuscript, and made constructive modifications and suggestions. SH devised the idea for the manuscript, led the whole project, critically appraised the manuscript from inception, and co-wrote the manuscript from the outset. All authors contributed to the article and approved the submitted version.

Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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