The pathophysiology of diastolic heart failure
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Abstract
Whilst resting disturbances of both diastolic and long-axis systolic function are observed in patients with heart failure who have normal left ventricular ejection fraction, recent evidence suggests that dynamic disturbances in cardiac function occur during exercise. A paradoxical slowing of left ventricular active relaxation during exercise limits cardiac filling and therefore stroke volume and appears to be due to the combination of cardiac energetic impairment and disturbed ventricular-vascular coupling.

Introduction and context
Many consider heart failure with preserved ejection fraction (HFpEF) to be a disorder of diastolic function (hence the term ‘diastolic heart failure’) [1], whilst others believe that it may be due to a combination of diastolic abnormalities with subtle disturbances of systolic function that are insufficient to reduce left ventricular ejection fraction (LVEF) [2]. In this review, HFpEF refers to heart failure patients with an LVEF of greater than 50% [3,4].

Resting left ventricular stiffness in HFpEF
Diastolic function is influenced by passive elastic properties of the left ventricle, by the highly energy-dependent process of active relaxation, and by the atrial contribution to filling. Zile et al. [5] demonstrated that patients with HFpEF had abnormal left ventricular (LV) relaxation and increased resting LV stiffness with a shift in the diastolic pressure-volume relationship upwards and to the left. However, other studies have shown that resting ventricular stiffness is not invariably increased in patients with HFpEF [6]. Increased myocardial mass or changes in the extra-myocardial collagen network [1] (increased collagen content and increased collagen cross-linking) may contribute to increased LV passive diastolic stiffness at rest [5]. A study by van Heerebeek et al. [7] examined differences in titin isoform expression between patients with systolic and ‘diastolic heart failure’. Titin is a large cytoskeletal protein that contributes to resting stiffness of the myocardium [8]. A shift toward the stiffer N2B isoform was observed in the diastolic heart failure group and toward the longer N2BA isoform in the systolic heart failure group when compared with previously published results from healthy controls [7]. A shift to expression of the shorter N2B isoform in response to increased arterial stiffness would increase ‘contractility’ (to compensate for increased aortic impedance) at the price of increased LV systolic and diastolic stiffness. More recently, myocardial biopsies from patients with aortic stenosis and patients with heart failure suggest that, rather than just a shift in titin isoforms, other mechanisms such as relative hypophosphorylation of the stiffer N2B titin isoform were a possible cause of increased LV stiffness in failing myocardium [9].

Resting large-artery stiffness and ventricular-vascular coupling
The interaction between the heart and the systemic vasculature, termed ventricular-vascular coupling (VVC), is essential for the heart to achieve maximal cardiac work, power and chamber efficiency while maintaining physiological blood pressures and cardiac outputs [10]. VVC is the ratio of arterial elastance ($E_a$) to end-systolic elastance ($E_{es}$). LV end-systolic elastance (stiffness) ($E_{es}$) is calculated from the slope of the end-systolic pressure-volume relationship. Arterial elastance (stiffness) ($E_a$) is...
a measure of impedance and is determined by the ratio of systolic pressure to stroke volume.

Ventricular-vascular interaction is important in the context of HfP EF because of its important effects on diastolic filling [11]. In patients with HfP EF, the resting VVC is lower than in younger individuals [12] but similar to asymptomatic hypertensive elderly patients [13] and falls within a range in which cardiac work and efficiency are not compromised [14]. However, although the resting VVC ratio is within the physiological range, the absolute values of $E_a$ and $E_s$ are considerably elevated, indicating increased arterial and ventricular systolic stiffness and this becomes important during exercise (see below).

**Systolic, diastolic and vascular function during exercise**

In young healthy subjects, exercise is associated with an increase in contractility and in the rate of LV active relaxation, although the latter is attenuated with increasing age [15]. In HfP EF, these physiological changes during exercise are profoundly deranged and this appears to be central to the pathophysiology of the disorder. In a small study by Kitzman et al. [16], HfP EF patients underwent invasive cardiopulmonary exercise testing. HfP EF patients exhibited a shift of the LV end-diastolic pressure-volume relationship upward and to the left at rest, and during exercise increases in LV filling pressure were not accompanied by increases in end-diastolic volume index, indicating a limitation to LV filling during exercise and a failure of the Frank-Starling mechanism [16]. More recently, a study conducted by Kawaguchi et al. [6] reported a dynamic impairment of LV active relaxation during isometric (handgrip) exercise in a group of HfP EF patients.

These findings suggest a potentially attractive link between increased large-artery stiffness and exercise-induced diastolic dysfunction. Animal studies have demonstrated that a large acute increase in afterload in healthy hearts of rabbits resulted in a marked slowing of active relaxation and impaired LV diastolic filling [17]. However, acute increase in afterload required to cause a slowing of active relaxation may be much less in a diseased heart compared with a healthy heart. The concept of ‘relative load’, which represents the ratio of systolic LV pressure to isovolumetric LV pressure [18], allows for the possibility that an increase in afterload required to cause a slowing of active relaxation may be much less in a diseased heart compared with a healthy heart. A similar systolic LV pressure represents a higher relative load in the failing heart than in the normal heart. When relative load is low, afterload reserve is still available, allowing the heart to face increased afterload without slowing of LV active relaxation. When relative load is high, afterload mismatch [19] occurs and a pronounced slowing of LV active relaxation is observed [18]. Other extra-cardiac factors that may contribute to the pathophysiology of HfP EF include volume overload in conditions such as anaemia, renal dysfunction and obesity [20]. In addition, previous data in an animal model have indicated that neurohormonal activation with intravascular volume expansion following relatively small degrees of ischaemic myocardial injury can lead to elevated LV end-diastolic pressure in the absence of diastolic dysfunction or reductions in maximal $dP/dt$ (rate of rise of LV pressure) or $EF$ [21].

**Major recent advances**

Borlaug and colleagues [22] recently reported that hypertensive patients with or without HfP EF had increased arterial and LV stiffness but similar VVC at rest. However, despite the apparently ‘preserved’ $EF$, patients with HfP EF had depressed cardiac contractility compared with asymptomatic hypertensive subjects [22]. This was consistent with other studies that had used tissue Doppler imaging and had demonstrated the presence of diastolic or systolic dysynchrony or both in patients with HfP EF [23, 24].

A study by Westermann et al. [25], involving pressure-volume loop analysis with and without atrial pacing in 70 HfP EF patients and 20 matched controls, found that enhanced LV stiffness can result in increased end-diastolic pressure during handgrip exercise. In addition, during atrial pacing, pacing with HfP EF displayed decreased stroke volume, but it should be noted that these patients were resting in a supine position and thus to artificially increase heart rate by atrial pacing might not reflect true physiological exercise conditions. At the very least, these data suggest that LV stiffness can modulate cardiac function in HfP EF patients. Another group, also using pressure-volume loop analysis, demonstrated that in HfP EF patients there was indeed increased LV passive stiffness and prolonged LV relaxation at rest and, interestingly, that during atrial pacing, LV relaxation reserve was limited but the cardiac contractility was normal. The net result was a blunted cardiac output during atrial pacing [26]. Increased LV stiffness in HfP EF patients may reflect not only increased myocardial mass and changes in extra-myocardial collagen (e.g., titins) but also LV fibrosis as demonstrated by Martos and colleagues [27], who found marked serological evidence of active fibrotic processes in patients with HfP EF.

Using echocardiographic techniques, our group recently demonstrated that, at rest, LV torsion and strain patterns
in HFPpEF patients were similar to those in age-related controls [28] but that during exercise HFPpEF patients had reduced systolic and diastolic function as well as evidence of delayed LV untwisting and LV suction [29]. In a separate study, using radionuclide ventriculography, we studied 37 HFPpEF patients and 20 control subjects during cycle exercise and demonstrated that HFPpEF patients had marked disturbances of VVC and of both systolic and diastolic function which appeared to be responsible for exercise limitation. LV active relaxation was paradoxically slowed. $E_a$ fell less and $E_a\prime$ increased much less during exercise than in age-matched controls, the latter indicating a failure of contractile reserve [30]. The impaired LV diastolic filling during exercise may be partly compensated by increased left atrial contribution during the final stages of diastolic filling [31] until atrial failure and eventually atrial fibrillation supervene later in the natural history of the disease.

A key coupler of load-dependent LV relaxation is troponin I-protein kinase A (TnI-PKA) phosphorylation. This energy-dependent process of phosphorylation of troponin I by PKA decreases myofibrillar calcium sensitivity and increases the rate at which calcium dissociates from troponin C and this can lead to an increased rate of LV relaxation by increasing the rate of thin filament deactivation. Using magnetic resonance spectroscopy, we recently demonstrated that HFPpEF patients had reduced myocardial energetic reserve at rest (decreased phosphocreatine/adenosine triphosphate [PCr/ATP] ratio), and this might explain why these HFPpEF patients are particularly prone to impaired LV active relaxation during exercise and impaired contractile reserve [30].

The cause for this resting energy deficit may relate to insulin resistance, to impaired mitochondrial function as a result of ageing, and to neuro-endocrine activation and aberrant substrate metabolism. In addition, a recent study in patients with hypertrophic cardiomyopathy found that reduced myocardial PCr/ATP correlated with the presence of fibrotic area in the myocardium of the left ventricle [32].

**Future directions**

There are consistent data demonstrating markedly abnormal diastolic responses to exercise in HFPpEF. In our view, the detection of HFPpEF patients cannot rely solely on resting parameters and we need to consider some form of exercise testing such as metabolic exercise testing or exercise radionuclide ventriculography or echocardiography scans in the future.

Future studies should begin to translate key pathophysiological findings in HFPpEF to an effective clinical therapy. An effective therapy for HFPpEF patients has yet to be developed. However, the finding that HFPpEF patients have reduced myocardial energetic status may provide the rationale to assess the therapeutic value of ‘metabolic agents’ (e.g., perhexiline and trimazadidine) that increase cardiac energy status by altering cardiac substrate use [33]. LV stiffness and fibrosis could be addressed by drugs such as aldosterone antagonists (e.g., eplerenone), which may help to alleviate aldosterone-induced cardiac fibrosis, which causes increased stiffness, impaired LV relaxation and impaired LV filling.

**Abbreviations**

$E_a$, arterial elastance (stiffness); $E_a\prime$, end-systolic elastance (stiffness); EF, ejection fraction; HFPpEF, heart failure with preserved ejection fraction; LV, left ventricular; LVEF, left ventricular ejection fraction; PCr/ATP, phosphocreatine/adenosine triphosphate; PKA, protein kinase A; VVC, ventricular-vascular coupling.

**Competing interests**

MF has received honoraria from Medronic (Minneapolis, MN, USA), St Jude Medical (St Paul, MN, USA), and Biotronik (Berlin, Germany) (amount of less than $10,000) and has served as a consultant and advisory board member of Medronic, St Jude Medical, Menarini (Florence, Italy) and Biotronik (amount of less than $10,000) and on the speaker panel for Menarini (amount of less than $10,000). TTP declares that he has no competing interests.

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