Arterial–ventricular coupling with aging and disease

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INTRODUCTION
The population in the Western world is aging; by 2030, there will be 71 million individuals in the United States over 65 years of age, representing ≈20% of the U.S. population. Aging significantly increases cardiovascular morbidity even in the absence of other risk factors (e.g., hypertension, obesity, diabetes, hypercholesterolemia). Thus, the risk of death from heart disease is ≈60-fold greater in individuals in the eighth decade compared to individuals in the 4th decade of life. Not only does clinically overt cardiovascular disease increase dramatically with aging, but so do subclinical or occult diseases, such as silent coronary atherosclerosis. Therefore, the aging of the U.S. population is one of the major public health challenges that we face in the twenty-first century.

The cardiovascular system is modulated to provide sufficient pressure and flow to the tissues at rest and during exercise. Understanding the performance (pressure and flow output) of the left ventricle (LV) requires not only examining the properties of the LV itself (power and stroke capacity of the heart), but also investigating the modulating effects of the arterial system on left ventricular performance. These modulating effects of the vasculature include the capacitance and inertial properties of the aorta, along with the resistance capacity of the microcirculation. The interaction of the LV with the arterial system, termed arterial–ventricular coupling (EA/ELV), is a central determinant of cardiovascular performance and cardiac energetics. EA/ELV can be indexed by the ratio of effective arterial elastance (EA; a measure of the net arterial load exerted on the LV) to left ventricular end-systolic elastance (ELV; a load-independent measure of left ventricular chamber performance). Age-associated alterations in arterial structure and function, including diameter, wall thickness, wall stiffness, and endothelial dysfunction, contribute to a gradual increase in resting EA with age. Remarkably there is a corresponding increase in resting ELV with age, due to alterations to LV remodeling (loss in myocyte number, increased collagen) and function. These age-adaptations at rest likely occur, at least, in response to the age-associated increase in EA and ensure that EA/ELV is closely maintained within a narrow range, allowing for optimal energetic efficiency at the expense of mechanical efficacy. This optimal coupling at rest is also maintained when aging is accompanied by the presence of hypertension, and obesity, despite further increases in EA and ELV in these conditions. In contrast, in heart failure patients with either reduced or preserved ejection fraction, EA/ELV at rest is impaired. During dynamic exercise, EA/ELV decreases, due to an acute mismatch between the arterial and ventricular systems as ELV increases disproportionately compared to EA (≈200 vs. 40%), to ensure that sufficient cardiac performance is achieved to meet the increased energetic requirements of the body. However, with advancing age the reduction in EA/ELV during acute maximal exercise is blunted, due to a blunted increase ELV. This impaired EA/ELV is further amplified in the presence of disease, and may explain, in part, the reduced cardiovascular functional capacity with age and disease. Thus, although increased stiffness of the arteries itself has important physiological and clinical relevance, such changes also have major implications on the heart, and vice versa, and the manner in which they interact has important ramifications on cardiovascular function both at rest and during exercise. Examination of the alterations in arterial–ventricular coupling with aging and disease can yield mechanistic insights into the pathophysiology of these conditions and increase the effectiveness of current therapeutic interventions.

Keywords: left ventricular function, arterial system, exercise, aging, disease
its physiological consequences. Further, we will discuss potential therapeutic interventions to restore the coupling between the heart and arteries.

**ARTERIAL–VENTRICULAR COUPLING**

The gold standard to assess the arterial afterload that opposes left ventricular ejection independent of left ventricular function is through aortic input impedance derived from the Fourier analysis of aortic pressure and flow (Murgo et al., 1980). However, aortic input impedance is described in the frequency domain, whereas measures of left ventricular contraction are best described in the time domain; consequently making direct comparisons between arterial and left ventricular function difficult. The pioneering work from Sunagawa et al. (1983) conceived a measure of arterial load \( (E_A) \) that could be directly compared to a measure of left ventricular contraction \( (E_{LV}) \) in the same units (elastance; change in impedance, and systolic and diastolic time intervals). EA represents the negative slope of the line joining the end-diastolic volume and end-systolic pressure points (Figure 1).

The contractile function of the LV can also be expressed from the slope of the end-systolic pressure–volume relationship \( (E_{LV}; \text{Figure 1}) \), which can be obtained from a series of pressure–volume loops recorded while the preload of the heart is altered. \( E_{LV} \) reflects a relatively (within normal physiological limits) load-independent measure of left ventricular contraction (chamber stiffness at end systole). An increase in contractility is depicted by an increase in the slope and a shift in the end-systolic pressure–volume relationship to the left, which allows the ventricle to generate more pressure for a given left ventricular volume. However, in addition to the inotropic state, \( E_{LV} \) is also influenced by the geometric (structural remodeling) and biochemical properties (i.e., stiffness or compliance of myocytes, composition of muscle, fibrosis, collagen, etc., in the LV wall) that underlie end-systolic stiffness (Borlaug and Kass, 2008). Whereby a “stiffer” LV due to remodeling leads to a higher \( E_{LV} \). Thus, caution should be exercised in interpreting the significance of an elevated \( E_{LV} \), particularly when other measures of left ventricular systolic function are normal. \( E_{LV} \) should, therefore, be considered an integrated measure of left ventricular chamber performance that can be related to an integrated measure of arterial load (i.e., \( E_A \)). Importantly, \( E_A \) shares common units with \( E_{LV} \), and their ratio \( E_A/E_{LV} \) is a measure of the interaction between the heart and the arterial system and provides information about how blood pressure and flow change due to different loading conditions. See Chantler et al. (2008a) for a more detailed review of the methods to measure \( E_A \) and \( E_{LV} \).

At rest, in healthy individuals the properties of the heart and arteries are closely matched so that near maximal cardiac work, power, and chamber efficiency are achieved (Little and Cheng, 1991; De Tombe et al., 1993). Values obtained in isolated canine hearts show the efficiency and stroke work of the heart to be optimal over an \( E_A/E_{LV} \) ratio ranging 0.3–1.3 (Figure 2). Whereas, in healthy humans the optimal range of \( E_A/E_{LV} \) to cardiac efficiency and stroke work are generally ranging from 0.7 to 1.0 (Asano et al., 1992; Najjar et al., 2004, Redfield et al., 2005). During exercise, an acute mismatch between the arterial and ventricular systems occurs due to a disproportionate increase in \( E_{LV} \) vs. \( E_A \) (Najjar et al., 2004). As a result \( E_A/E_{LV} \) decreases to ensure that sufficient cardiac performance is achieved to meet the increased energetic requirements of the body. In the next section, we will review the alterations in \( E_A/E_{LV} \) at rest and during exercise due to aging and illustrate how the age–disease interaction accentuates the changes in \( E_A/E_{LV} \). We will also discuss therapeutic interventions aimed to restore the relationship between
E_A and E_LV, including the physiological meaning behind the adaptations.

**HEALTHY AGING AND ARTERIAL VENTRICULAR COUPLING AT REST**

Numerous studies have documented a gradual increase in E_A with advancing age (Cohen-Solal et al., 1996; Chen et al., 1998, Redfield et al., 2005). Two small clinical cohort studies calculated E_A from invasive methods of flow and pressure and reported a significant rise in E_A (44–73%, p < 0.001) from ≈20 to 70 years of age (Cohen-Solal et al., 1996; Chen et al., 1998). Although the strength of these data lies in the invasive assessment of E_A, the small sample size (Cohen-Solal et al., 1996; Chen et al., 1998), and the inclusion of patients with coronary artery disease, and individuals on chronic medications (Chen et al., 1998), provides limited insight into the influence of healthy aging on E_A. Non-invasive studies of E_A have also reported an increase in E_A with age. In a larger epidemiological study consisting of 623 individuals from Olmsted County, Minnesota study, Redfield et al. (2005) observed an age-associated increase in E_AI (E_A normalized for body surface area) in both men (r = 0.18, p < 0.01) and women (r = 0.13, p = 0.02). Inclusion of individuals with existing cardiovascular disease further increased the age-associated change in E_AI in both men (r = 0.28, p < 0.001) and women (r = 0.28, p < 0.001; Figure 3A).

The specific mechanisms for the increased E_A with age reflect the age-associated changes in the arterial properties of individuals. E_A is a lumped parameter incorporating mean resistance and pulsatile properties of the arterial load, and is therefore influenced by changes in arterial compliance, wave reflection, and characteristic impedance (Kelly et al., 1992a). During the past two decades, we have characterized the effects of aging on multiple aspects of arterial structure and function in a single study population in the Baltimore Longitudinal Study on Aging (BLSA) who are rigorously screened to exclude both clinical and occult cardiovascular disease. Although at rest, peripheral resistance is the dominant factor affecting E_A (Chemla et al., 2003) the age-associated actions of peripheral resistance are heterogeneous to the extent that only an age-associated increase in resting peripheral resistance is noted in healthy women but not men (Fleg et al., 1995). Indeed, the increase in E_A with age noted by Redfield et al. (2005) is not attributed to an increase in peripheral resistance as no relationship between age and peripheral resistance existed. E_A is also altered by arterial stiffening and blood pressure pulsatility increases with age (Franklin et al., 1997) thus raising the end-systolic pressure required to eject blood thereby increasing E_A. Findings from the BLSA indicate that healthy aging is associated with an increase in aortic root diameter, an increase in carotid wall intimal media thickness, and an overall stiffening of the large elastic arteries (Najjar et al., 2005). Redfield et al. (2005) proposed the increase in E_A with age is attributable to an increase in arterial stiffness, manifested by an increase in pulse pressure (a surrogate measure of arterial stiffening). Thus, the age-associated increase in E_A is largely attributed to an increase arterial stiffness.
The heart seems to respond to an increase in $E_A$ with a corresponding increase in $E_{LV}$ with age (Cohen-Solal et al., 1996; Chen et al., 1998; Redfield et al., 2005). Cohen-Solal et al. (1996) noted a 28% higher $E_{LV}$ in men ≈60 vs. 30 years of age. In the Olmsted County, Minnesota study, Redfield et al. (2005) noted a 10% ($r = 0.16, p < 0.001$) and 15% ($r = 0.23, p < 0.0001$) increase in $E_{LV}$ in men and women without cardiovascular disease, respectively (Figure 3B). These results would suggest that $E_{LV}$ increases to compensate for the increase in $E_A$ ensuring that their ratio, $E_A/E_{LV}$, is matched for maximal efficiency at rest. Indeed, significant relationships exist between $E_A$ and $E_{LV}$ ranging from correlations of 0.50 to 0.73 (Chen et al., 1998; Borlaug et al., 2009) suggesting that a change in $E_A$ accounts for a ≈25% change in $E_{LV}$ (Figure 4). This small percentage is not surprising given that both components have multiple different determinants. Nevertheless, older individuals with an increased arterial stiffness ($E_A$) are more likely to have an increased left ventricular stiffness ($E_{LV}$), i.e., a stiffer heart coupled to a stiffer vascular system. A consequence of the age-associated increase in $E_{LV}$, along with an increase in $E_A$, essentially permits the $E_A/E_{LV}$ ratio to remain relatively matched (coupled; Cohen-Solal et al., 1996; Chen et al., 1998, Redfield et al., 2005). However, Redfield et al. (2005) did note a slight sex difference in $E_A/E_{LV}$ with age whereby healthy women demonstrated a slight decline in $E_A/E_{LV}$ with age, reflecting a disproportionate increase in $E_{LV}$ compared with $E_A$. This suggests a greater impact of aging on ventricular vs. arterial properties in women compared to men, as indicated in Figure 4.

Given that $E_{LV}$ is as a load-independent index of left ventricular contractility (Sagawa, 1978), one might suggest that an increase in $E_{LV}$ with age (and more so in women) reflects enhanced contractility. This is unlikely given that other measures of left ventricular function do not increase with age (Lakatta, 1993). Thus, what does the increase in $E_{LV}$ likely reflect? $E_{LV}$ is also influenced by the structural/geometric and biochemical properties (i.e., stiffness or compliance of myocytes, composition of muscle, fibrosis, collagen, etc., in the heart wall) that underlie left ventricular end-systolic stiffness (Borlaug and Cass, 2008; Chantler et al., 2008a). Advancing age is also associated with alterations in left ventricular structure. Most notably, there is a reduction in myocyte number, and there is an increase in left ventricular wall thickness and collagen deposition in the heart (Olivetti et al., 1995; Lakatta, 2003). Thus, the increase in $E_{LV}$ with age could represent differences in left ventricular geometry and/or structure between young and older individuals. For example, concentric remodeling leads to a higher $E_{LV}$ (Borlaug et al., 2009). However, the increase in $E_{LV}$ noted in Redfield et al. (2005) are unlikely to be due to differences in LV chamber size, as similar results are obtained when $E_{LV}$ is normalized to end-diastolic volume (a crude estimate of chamber size). An increase in the amount (focal increases) and a change in the physical properties of collagen (purportedly due to non-enzymatic cross-linking) also occur within the myocardium with aging (Lakatta and Levy, 2003). Thus, the increase in $E_{LV}$ with age is likely attributed to a combination of left ventricular remodeling but more so due to a loss of myocyte number and an increase in myocardial collagen content, thereby increasing myocardial stiffness.

**DURING EXERCISE**

During exercise $E_A$ has been shown to increase (Najjar et al., 2004; Otsuki et al., 2006), decline (Asanot et al., 1992), or remain unchanged (Cohen-Solal et al., 1998). The response of $E_A$ during exercise is dependent on the changes in its components and has important consequences on the Frank–Starling mechanism (Chantler et al., 2011). $E_A$ is linearly related to heart rate and peripheral resistance, and inversely related to compliance (Segers et al., 2002; Chemla et al., 2003, Otsuki et al., 2006; Chantler et al., 2011). Both resistance and compliance usually decrease during exercise (reflecting less resistance to blood flow in the microcirculation, but increased stiffness of the conduit arteries) and the relative contribution of the pulsatile component (compliance) to $E_A$ increases, so that by 80% of peak exercise the resistive and the pulsatile components provide nearly equal contributions to $E_A$ (Otsuki et al., 2006). With advancing age, the ability to increase heart rate, and lower resistance during exercise is blunted. In addition the reduction in compliance (due to a greater increase in pulse...
pressure) is also limited during exercise (Lakatta, 1993; Fleg et al., 1995). Although the blunted maximum cardiovascular responses with age are, in part, due to older individuals achieving a lower maximal workload, similar cardiovascular deficits are also evident at submaximal workloads (Lakatta, 1993; Fleg et al., 1995). The only study to directly examine the age-associated change in $EA/ELV$ in the absence of cardiovascular disease reported that $EA$, in general, increases during exercise and that $E_A$ does not differ between young and older subjects (Najjar et al., 2004). Perhaps the blunted changes in resistance, compliance, and heart rate are compensated for by the greater increase in blood pressure during exercise in older vs. younger healthy individuals (Ogawa et al., 1992). Indeed, some of the components of the changes in $E_A$ seem to be related to each other (Chantler et al., 2012). That is, greater preservation of compliance during exercise is associated with a greater reduction in resistance (and a smaller increase in mean and pulse pressure). This suggests that the tandem changes in resistance and compliance appear to be linked, and raises the possibility of a crosstalk between central and peripheral arteries. Further, the change in $E_A$ during exercise is also linked to a specific pattern of change in ventricular volumes and function. The change in $E_A$ is inversely related to the recruitment of end-diastolic volume, and the enhancement of stroke volume and cardiac output with exercise. Indeed individuals expressing a large increase in $E_A$ during exercise demonstrate a blunted utilization of the Frank-Starling mechanism, irrespective of age, sex, and body size (Chantler et al., 2012). The cardiovascular system responds to exercise by increasing cardiac output predominately through an increase in heart rate, and in the upright position stroke volume is increased. The perfusion of the tissues depends both on the ability of the LV to produce flow and sustain perfusion pressure. Thus, the speed and force of left ventricular contraction increases during exercise and this is reflected by an increase in $E_{LV}$ during exercise (Little and Cheng, 1993). Unlike $E_A$, aging impairs the increase in $E_{LV}$ during exercise. Najjar et al. (2004) reports that the blunted increase in $E_{LV}$ with aging begins to appear at 50% of maximal workload and that at maximal exercise older men and women (>60 years) have $\approx 40-55\%$ smaller $E_{LV}$ compared to men and women <40 years of age. A consequence of the blunted increase in $E_{LV}$ during exercise in older healthy individuals is a corresponding blunted reduction in $E_{A}/E_{LV}$ (Najjar et al., 2004). Furthermore, the altered coupling noted at peak exercise with age is also associated with a blunted increase in ejection fraction. Indeed, $E_A/E_{LV}$ is inversely related to ejection fraction $\left[ E_A/E_{LV} \approx \left(1/ejection\;fraction\right) - 1 \right]$; Cohen-Solal et al., 1994]. The advantage of $E_A/E_{LV}$ over ejection fraction is that examining the components of $E_A/E_{LV}$ permits evaluation of whether alterations in $E_A/E_{LV}$ are due to alterations in arterial properties, left ventricular properties, or both. Consequently, advancing age is associated with a smaller $E_A/E_{LV}$ (and ejection fraction) reserve $\left( E_A/E_{LV} = 1 \text{ peak} - E_A/E_{LV} \text{ rest} \right)$ and $E_{LV}$ reserve $\left( E_{LV} \text{ peak} - E_{LV} \text{ rest} \right)$; Najjar et al., 2004). Unlike at rest when an increase in $E_{LV}$ could represent enhanced left ventricular contraction, decreased left ventricular mass (or concentric remodeling), or increased myocardial stiffness (due to alterations in biochemical properties), an increase in $E_{LV}$ during exercise likely reflects an increase in inotropy. This would suggest a decrease in left ventricular contraction during exercise with increasing age.

### Aging-CV Disease Interactions Affect Arterial Ventricular Coupling

Unfortunately in today’s society, for the most part, aging is highly linked to the occurrence of cardiovascular diseases. Further, cardiovascular disease risk factors (obesity, hypercholesterolemia, diabetes, and hypertension) often co-vary in number or severity with increasing age. Although measuring the age-associated changes in cardiovascular structure and function in a “healthy” aging population is important to provide insights into the normal aging process, the generalizability of these findings are sometimes limited. The age and disease interaction has important consequences on arterial ventricular coupling both at rest and during exercise as outlined below.

### Hypertension

The prevalence of hypertension markedly increases with advancing age, such that the relative risk of acquiring hypertension is $\approx 90\%$ of individuals over 40 years of age (Lloyd-Jones et al., 2009; Figure 5). Age-associated changes in arterial and left ventricular structure and function are accelerated in the presence of hypertension. Hypertensive patients exhibit greater carotid wall thickness (Arnett et al., 1996), central arterial wall stiffness (Amar et al., 2001), and reflected waves (Nichols et al., 1992) than normotensive subjects, even after adjusting for age. Furthermore, hypertension is associated with left ventricular remodeling and fibrosis (Mayet and Hughes, 2003). As such, $E_A$ and $E_{LV}$ are reported to be increased, between 15-60 and 16-95%, respectively, in hypertensive patients compared with normotensive controls (Cohen-Solal et al., 1994; Saba et al., 1999, Lam et al., 2007). As with aging, however, the coupling ratio $\left( E_A/E_{LV} \right)$ remains matched (coupled) between normotensives and hypertensives (Cohen-Solal et al., 1994; Saba et al., 1999). We found similar results (matched $E_A/E_{LV}$ due to tandem increase in $E_A$ and $E_{LV}$) when comparing men with predominantly systolic hypertension to normotensive men. In contrast, women with predominantly systolic hypertension, have a 23% lower resting $E_A/E_{LV}$ than normotensive women, a finding that persisted even after adjusting for age (Chantler et al., 2008b). The lower $E_A/E_{LV}$ in women with systolic hypertension is due to a disproportionate increase in $E_{LV}$ compared with $E_A$ (45 vs. 16%), suggesting an adaptation by these women to limit the impact of systolic hypertension on the vasculature or, alternatively, a more pronounced impact of systolic hypertension on ventricular vs. arterial elastance.

There are a limited number of studies that have examined the coupling response during dynamic exercise in hypertensive individuals. Borlaug et al. (2010) showed that hypertensive individuals express similar changes in $E_A$, $E_{LV}$ and $E_A/E_{LV}$ at submaximal and maximal exercise compared to normotensive individuals matched by age and sex. In contrast, Chantler et al. (2008b) reported that the effects of predominantly systolic hypertension on the changes in $E_A/E_{LV}$ are similar between normotensive men and systolic hypertensive men and women at 50% of maximal workload and at peak upright bicycle exercise. In men, this is because $E_A$ and...
ELV are proportionally higher at peak exercise in systolic hypertensive compared with normotensive individuals. In women, this is because EA and ELV are similar at peak exercise between systolic hypertensive and normotensive women. Thus, EA/ELV reserve is similar between systolic hypertensive and normotensive men, but 61% lower in systolic hypertensive vs. normotensive women because of the uncoupling in EA/ELV at rest in these systolic hypertensive women.

**OBESITY**

About one-third of U.S. adults (33.8%) are obese and the prevalence of overweight/obesity increases with advancing age. Obesity-related conditions include heart disease, stroke, type 2 diabetes, and certain types of cancer, some of the leading causes of death (Flegal et al., 2010). The physiological changes that occur with obesity are similar to those in hypertension, and often hypertension and obesity co-exist in a given individual complicating interpretations due to obesity alone (Stamler et al., 1978, 1991). Indeed, Zebekakis et al. (2005) identified that obese individuals (14% on antihypertensive medications) have increased carotid, brachial, and femoral diameter with age, which remained significant after adjusting for differences in BP, medications, and smoking status. Further, increasing body mass index is associated with a reduced arterial distensibility (Zebekakis et al., 2005) and increased arterial stiffness (Toto-Moukouo et al., 1986; Zebekakis et al., 2005). In individuals with uncomplicated obesity (without the presence of cardiovascular disease), an increase in body mass index (45 \( \pm \) 5 vs. 21 \( \pm \) 2) is associated with an increase in left ventricular mass (irrespective of normalized to height or body surface area), and end-diastolic volume (unadjusted or indexed to height; Rider et al., 2011). Although such alterations in arterial and left ventricular structure/function increase resting EA and ELV in obese vs. non-obese controls, their coupling (EA/ELV), however, remains matched, irrespective of body size.

Rider et al. (2011) noted a similar resting ejection fraction, which is inversely related to EA/ELV, between individuals of various body mass index categories. However, whether the increase in EA and ELV noted in obese individuals represents the pathologic effects of excess adipose tissue on cardiovascular function, or the normal physiological relationship between body size and cardiovascular function is important to decipher. In order to have meaningful clinical and scientific comparisons, differences in body size must be accounted for (Chantler et al., 2005; Chantler and Lakatta, 2009). Indeed, in the unadjusted form, resting EA and ELV are slightly reduced (4 and 3%, respectively) in obese vs. healthy non-obese controls (Chirinos et al., 2009). In contrast, when EA and ELV are scaled either ratiometrically or allometrically to body surface area, EA and ELV are between 16–20 and 18–19% higher \((p < 0.001)\) in obese vs. controls. Irrespective of the scaling technique employed there was no relationship between obesity and EA/ELV (Chirinos et al., 2009). Unfortunately, the changes in EA and ELV, and subsequently EA/ELV have not been reported during dynamic exercise in obese vs. non-obese individuals.

**HEART FAILURE**

Heart failure (HF) is a syndrome that is characterized by an inability of the heart to pump a sufficient amount of blood to meet the demands of the metabolizing tissues, or can do so only at the expense of elevated filling pressures (Adams et al., 2006). The inability to meet the tissues’ demands is attributed to the ability, or lack thereof, for the LV to fill (diastolic properties) or to eject (systolic properties) blood. Although there is considerable confusion as to the pathological mechanisms that describes individuals with systolic (HF with reduced ejection fraction), or diastolic (HF with preserved ejection fraction) the coupling between the heart and arteries does seem to be impacted differently by the type of HF.
HF WITH A REDUCED EF
Heart failure patients with systolic dysfunction are characterized by a diminished resting ejection fraction and left ventricular contractility (Asanai et al., 1989). Patients with HF and a reduced ejection fraction have a downward and rightward shift of the end-systolic pressure–volume relationship, reflecting a reduced ELV (range 0.6–2.6 mmHg·ml−1·m−2; Asanai et al., 1989). Patients with HF and a reduced ejection fraction have an augmented EA (range 1.7–3.7 mmHg·ml−1·m−2) due to a decrease in stroke volume and increases in heart rate and peripheral resistance (Asanai et al., 1989). The increase in EA and decrease in E LV result in a three to fourfold increase in EA/ELV (range 1.3–4.3; Asanai et al., 1989; Sasayama and Asanai, 1991). This suboptimal coupling reflects diminished cardiovascular performance and efficiency of the failing heart.

During exercise, the traditional reduction in EA/ELV due to a substantial increase in E LV vs. any change in EA are virtually absent in HF patients with a reduced ejection fraction (Cohen-Solal et al., 1998). Thus the limited capacity of systolic HF patients to augment their cardiovascular function during times of stress (such as exercise) involve marked deficits in both left ventricular and arterial elastance reserves.

HF WITH A PRESERVED EF
Heart failure patients with a preserved ejection fraction (≥50%) represent ≈40% of patients with HF (Owan et al., 2006). HF with a preserved ejection fraction is more prevalent with advancing age, in women, and in individuals with systolic hypertension (Klahpholz et al., 2004). Recent interest has refocused attention on examining the coupling between the heart (systolic and diastolic properties) and arterial system in order to gain further insights into the pathophysiological mechanisms that underlie HF with a preserved ejection fraction. Some of the pathophysiological adaptations that occur in HF with a preserved ejection fraction can be attributed to normal adaptations evident with aging. Indeed the relative risk (controlled for gender, race, medical history, and admission characteristics) of developing HF with a preserved ejection fraction is ≈10–27% between 65 and 90 years of age (Masoudi et al., 2003). However, aging alone does not account for all the physiological changes noted in HF with a preserved ejection fraction. For example, compared to age-matched normotensive controls, HF patients with a preserved ejection fraction typically have an EA that is ≈40% higher and an E LV that is approximately twofold greater, and thus a lower EA/ELV is noted in HF patients with a preserved ejection fraction vs. healthy controls (Kawaguchi et al., 2003). However, when comparing HF patients with a preserved ejection fraction with age, and hypertensive matched controls without HF E A/ELV is similar because of a tandem rise in both E A (≥40%) and E LV (≥50%) in HF patients with a preserved ejection fraction (Kawaguchi et al., 2003). In larger epidemiological studies in whom E A/ELV is examined non-invasively, HF patients with a preserved ejection fraction have matched increases in E A and E LV (and thus similar E A/ELV) compared to non-hypertensive controls without HF (Lam et al., 2007; Borlaug et al., 2009). Further, no differences in E A and E LV, or E A/ELV are found between individuals with hypertension without HF and HF patients with a preserved ejection fraction (Lam et al., 2007; Borlaug et al., 2009). Also the increase in E LV in HF patients with a preserved ejection fraction compared to non-hypertensive controls is not due to differences in left ventricular remodeling but more likely due to an increased passive myocardial stiffening (Borlaug et al., 2009). However, the vast majority of HF patients with a preserved ejection fraction in these epidemiological studies are hypertensive (>95%), which may contribute to the matched increases in E A and E LV. Indeed, some patients with HF patients with a preserved ejection fraction who are normotensive have values of E A and E LV that are similar to those of healthy normotensive controls (Maurer et al., 2005). These findings highlight the difficulty in understanding the pathophysiological mechanisms that are evident in a disease that is comprised of a very heterogeneous group of individuals.

Recent studies have highlighted the importance of altered arterial–ventricular interactions during exercise. HF patients with a preserved ejection fraction who have increased E A and E LV at rest exhibited a markedly hypertensive response and elevated diastolic pressures to sustained handgrip exercise (Kawaguchi et al., 2003). Further, during upright cycle ergometry exercise at maximal effort, HF patients with a preserved ejection fraction have a threefold smaller increase in E LV and a reduced ability to lower their peripheral resistance and increase their heart rate during exercise compared with hypertensive controls with left ventricular hypertrophy (Borlaug et al., 2006). Some of the peak cardiovascular deficits noted between HF patients with a preserved ejection fraction and hypertensive controls could be related to the fact that the HF patients with a preserved ejection fraction are unable to attain the same level of exercise workload compared to controls. In other words, are the peak cardiovascular deficits a mechanism or consequence of exercise limitation? However, similar deficits are also noted between HF patients with a preserved ejection fraction and hypertensive controls at matched submaximal workloads (Borlaug et al., 2010). Thus, HF patients with a preserved ejection fraction are characterized by a diminished increase in E LV and a smaller reduction in E A/ELV compared to controls without hypertension, and to those with hypertension but without HF, both at submaximal exercise workloads (20 W) and at maximum exercise (Borlaug et al., 2010). In addition, the depressed reserve responses correlated with reduced exercise capacity and greater subjective symptoms at low-level workloads. Since female sex, systolic hypertension, and older age are risk factors for HF with a preserved ejection fraction (Klahpholz et al., 2004), and as the pathophysiology of HF with a preserved ejection fraction involves a limited cardiovascular reserve (Kitzman et al., 1991), the diminished E A/ELV reserve observed in older systolic hypertensive women without HF (Chantler et al., 2008b) suggests that they may be exhibiting signs of subclinical (Stage B) HF. This raises the possibility that they may be on a trajectory to progressive exercise intolerance and perhaps functional limitations.

In summary, the pathophysiological mechanisms that contribute to HF with a preserved ejection fraction are due to the accumulation of multiple cardiovascular impairments that are expressed during exercise, reflecting impaired inotropic, chronotropic, lusitropic, and vasodilatory responses that impair E A/ELV (Kawaguchi et al., 2003; Borlaug et al., 2010).
CONSEQUENCES OF ALTERATIONS IN $E_A/E_L$ WITH AGING, HYPERTENSION, OR HEART FAILURE

In a young, healthy individual the coupling between the arteries and heart are well matched to: (1) maintain an optimal transfer of blood from heart to periphery without excessive changes in blood pressure and; (2) provide optimal cardiovascular flow reserve without compromising arterial pressures (Kass, 2005). However, the increased stiffness noted in both the arteries and heart with age, which is further exacerbated in the presence of disease (hypertension, HF, etc.) reflects a coupling disease (Kass, 2005). That is the stiff arteries and LV interact to limit the cardiovascular response to stress (reduced performance) and generates clinical symptoms (Kass, 2005).

As illustrated in Figure 6, a stiffer heart-arterial system increases load-sensitivity even if the coupling ratio is normal. That is, an increased resting $E_A$ and $E_L$ means that systolic pressures are much more sensitive to changes in left ventricular volume. This is clearly observed in young vs. old individuals, whereby a decreased preload in younger individual results in a modest drop in systolic pressure but in older individuals there is a much greater change in systolic pressure (Chen et al., 1998). This exaggerated systolic pressure response is also evident in hypertensives and HF patients with a preserved ejection fraction compared to controls (Kawaguchi et al., 2003). Consequently, the stroke work (myocardial demand) required to perform this task is increased and can potentely influence systolic and diastolic function, including coronary flow (i.e., greater dependence upon systolic pressure for coronary flow; Kawaguchi et al., 2003). Thus, older individuals are working at a higher setpoint regarding changes in pressure for a given change in loading conditions and this disadvantage is further exaggerated in hypertensives and HF patients with a preserved ejection fraction.

In addition to enhanced load-sensitivity, the global (systolic and diastolic) reserve capacity becomes blunted with arterial and left ventricular stiffening. Increased $E_L$ at rest translates into less effective changes in $E_L$ during exercise (or stress) thereby limiting cardiovascular performance (Borlaug et al., 2010). For example, individuals who start at a higher $E_L$ have a limited capacity to further increase stroke volume, and the limited stroke volume response is further exacerbated when a stiff heart is connected to a stiff artery (Chen et al., 1998; Kawaguchi et al., 2003). It is therefore not surprising that a stiffer LV and arterial system are linked to a reduced aerobic capacity (Borlaug et al., 2010). Further, in an isolated canine heart model, Kass et al. (1989) reported, larger reductions in $E_L$ after a myocardial infarction in hearts with a higher resting $E_L$. This greater mechanical vulnerability to an ischemic insult may help to explain why older individuals may experience worse outcomes following a myocardial infarction (Maggioni et al., 1993).

Another a major consequence of arterial stiffening is an increased pulse pressure, which also increases cyclic changes of arterial flow. As such, the microcirculation receives larger pulsatile pressures which can damage the vascular beds and in turn, cause damage to the end organs (such as the kidney and brain). Indeed increased arterial stiffness is independently associated with dementia (Hanon et al., 2005). Increased arterial stiffness can also lead to endothelial dysfunction and an abnormal vasodilation response to stress.

THERAPEUTIC INTERVENTIONS

Interventions related to improving arterial ventricular coupling span from pharmacological to lifestyle (exercise and diet) approaches. We will briefly highlight some important interventions in this area. The abnormalities in combined arterial and ventricular stiffening leading to a mismatch in their interaction, as highlighted above, has important physiological consequences. Numerous approaches have been taken to reverse the age and disease associated changes. By reducing the increase in arterial and left ventricular systolic stiffness with age or disease we may improve arterial-ventricular interactions and thus cardiovascular performance by being more efficient blood pressure regulation for a given volume of blood. For example, acute administration of sodium nitroprusside, a balanced vasodilator, acutely reduces $E_A$ (10%) and increases $E_L$ (47%) at rest in older (70±8 years) individuals (Chantler et al., 2011). Further, at peak exercise, sodium nitroprusside leads to an increase in $E_L$ (68%) without a change in $E_A$ and consequently the normal reduction in $E_A/E_L$ during exercise is enhanced (36%). Importantly, sodium nitroprusside acutely attenuates the age-associated deficits in $E_A/E_L$ and $E_L$ during exercise (Chantler et al., 2011). Similarly, acute intravenous verapamil (calcium-channel blocker) in healthy older (70±10 years) volunteers reduces resting arterial-ventricular stiffening and $E_A$ during exercise (though a decline in heart rate, pulsatile, and resistive arterial load), and improves (13%) aerobic exercise performance (Chen et al., 1999). These results in older persons highlight that, at least acutely, the abnormalities in arterial-ventricular interactions can be partly restored. Of note, verapamil also improved arterial-ventricular interactions and exercise capacity in HF patients with a preserved ejection fraction and hypertrophic cardiomyopathy (Brown et al., 1985; Setaro et al., 1990). However, whether the acute effects of sodium nitroprusside or verapamil identified can be maintained with chronic drug administration is unknown.

![FIGURE 6](image-url)
Hypertensive patients on optimal brachial and central blood pressure antihypertensive therapies shifts arterial–ventricular coupling from blood flow maximization to left ventricular mechanical efficiency optimization (Osranek et al., 2008). Further, the effects of antihypertensive monotherapy on $E_A/E_LV$ examined in 10,670 patients over a 6-month period indicated that angiotensin converting enzyme inhibitors (ACEI), angiotensin II receptor blockers (AIIRA), and dihydropyridine calcium antagonists decrease $E_A/E_LV$, whereas diuretics, $\alpha$-blockers, both $\beta$-blocker groups (with and without intrinsic sympathomimetic activity), and non-dihydropyridines significantly increase $E_A/E_LV$ compared to baseline measurements (Figure 7; Iakovou et al., 2004). Thus various antihypertensive drugs have a differential effect on $E_A/E_LV$ with ACEI, AIIRA, and dihydropyridine calcium antagonists have the most favorable effect on this index, likely through their actions on causing vasodilatation and by actually inhibiting the vasoconstrictive results of neurohormonal activation (Iakovou et al., 2004). However, the effects of these drugs on the components of $E_A/E_LV$ are not reported.

Other therapies shown to improve $E_A/E_LV$ are exercise training. In healthy older men, 24–32 weeks of aerobic endurance exercise training does not alter baseline ejection fraction (inverse of $E_A/E_LV$) or left ventricular contractility (systolic blood pressure/end-systolic volume), but increases peak ejection fraction, suggesting that $E_A/E_LV$ would have further decreased during exercise, due to a greater peak left ventricular contractility (Schulman et al., 1996). In the same study, eight master athletes stopped their endurance training for 12 weeks, which tended to decrease peak ejection fraction and LV contractility (Schulman et al., 1996). In patients with coronary artery disease, 12 months of aerobic endurance exercise training did not alter resting $E_A/E_LV$, or $E_LV$, but produced a slight 13% reduction in $E_A$ at rest (Rinder et al., 1999).

Further, exercise training led to a 37% increase in $E_{LV}$ and a 23% decrease in $E_A/E_{LV}$ during handgrip exercise performed at 30% of maximal voluntary contraction. However, the change in $E_A$ during handgrip exercise remained unaltered after the exercise training. The results of this study suggest that long-term endurance exercise training induces significant cardiovascular adaptations both in the basal state and during an afterload stress in patients with coronary artery disease.

One year of progressive and vigorous endurance training in sedentary healthy older (70 ± 3 years) individuals resulted in slight reductions in $E_A$ at rest (14%) and peak exercise (20%). This coincided with an improvement in compliance at rest (Fujimoto et al., 2010). However, the exercise training failed to reverse cardiac stiffening. One possible reason for the lack of changes in left ventricular stiffness with exercise training is the development of cross-linked advanced glycation end products in the left ventricular wall along with a loss in the number and the volume of cardiac myocytes that occur with older age and which are pathologically irreversible once formed (Aronson, 2003). Thus any improvement in left ventricular function could have been constrained by cross-linked collagen. A phase II drug, alagebrium, is a cross-link breaker and is known to improve left ventricular and arterial stiffness in animals (Kass, 2003), and a clinical trial is currently underway to examine the effects of exercise and alagebrium combined on cardiovascular stiffening in the elderly (NIH clinicaltrials.gov identifier NCT01014572). However, alagebrium administered to patients with HF and systolic dysfunction does not improve exercise tolerance or cardiac function (Hartog et al., 2011). Whether HF patients with a preserved ejection fraction, in whom the largest effect of alagebrium could be expected, would benefit from an advanced glycation end products breaker therapy remains unknown.

**FIGURE 7** | Arterial–ventricular coupling (AVC) percentage changes after antihypertensive monotherapy in 10,670 patients over 6 months of treatment. Abbreviations: ACE inhibitors, angiotensin converting enzyme inhibitors; AIIRA, angiotensin II receptor antagonists; ISA, intrinsic sympathomimetic activity. From Iakovou et al. (2004) with permission.
CONCLUSION

Although increased stiffness of the arteries itself has important physiological and clinical relevance, such changes also have major implications on the heart, and vice versa, and the manner in the way they interact has important ramifications on cardiovascular function both at rest and during exercise. Examination of the alterations in arterial–ventricular coupling with aging and disease can yield mechanistic insights into the pathophysiology of these conditions and increase the effectiveness of current therapeutic interventions. Future studies should identify agents to chronically reverse increases in $E_a$ and $E_L$ that occur with age and disease. Furthermore, longitudinal studies are needed to evaluate whether alterations in $E_a/E_L$, $E_a$, and $E_L$ provide any prognostic information for adverse outcomes, such as HF.

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