Recommendations on the use of echocardiography in adult hypertension: a report from the European Association of Cardiovascular Imaging (EACVI) and the American Society of Echocardiography (ASE)

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Hypertension remains a major contributor to the global burden of disease. The measurement of blood pressure continues to have pitfalls related to both physiological aspects and acute variation. As the left ventricle (LV) remains one of the main target organs of hypertension, and echocardiographic measures of structure and function carry prognostic information in this setting, the development of a consensus position on the use of echocardiography in this setting is important. Recent developments in the assessment of LV hypertrophy and LV systolic and diastolic function have prompted the preparation of this document. The focus of this work is on the cardiovascular responses to hypertension rather than the diagnosis of secondary hypertension. Sections address the pathophysiology of the cardiac and vascular responses to hypertension, measurement of LV mass, geometry, and function, as well as effects of treatment.

Keywords
Hypertension • Echocardiography

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Pathophysiology of cardiac responses to hypertension

Left ventricular hypertrophy

Size and geometry of the normal heart

The main contribution of echocardiography to the management of hypertension is the assessment of left ventricular (LV) mass (LVM).

Body habitus represents one of several factors that confound the association between hypertension and LVM. However, cardiac size is influenced by body size, and for any given size, men have larger hearts than women, athletes have larger hearts than non-athletes, and obese subjects have larger hearts than non-obese subjects. LVM and volumes bear an approximately quadratic (rather than approximately cubic) relationship with height in men and women.

In the enlarged heart, wall (fibre) stress increases with LV size (radius and volume). This increase is compensated by a proportional increase of wall thickness, so that wall stress remains matched with the systolic pressure. The ‘relative’ geometry of the ventricle appears to be similar across species and body size, with normal relative wall thickness [RWT, the ratio of twice the posterior wall thickness (PW) and the LV diastolic diameter] from 0.32 to 0.42.

Mass/volume ratios corresponding to the above-mentioned normal RWTs range between 1.1 and 1.3. RWT and M/V do not require correction for body size.

Effect of gender

Data from several studies indicate that after adjustment for blood pressure and anthropometric parameters, LV volume and LVM are higher in men than in women. These differences persist when values of LVM are corrected for fat-free mass. This sex difference may explain the surprising lack of consensus in appropriate indexation of LVM, as it impacts the optimal method for indexing LVM for body height. Figure 1 displays LVM, calculated by the Devereux formula (unidimensional 2D measurements) in the healthy reference subgroup of the Asklepios population. Using the allometric index 1.7, the body

Figure 1 Relationship between body height and LVM, calculated by the Devereux formula (unidimensional 2D measurements). Body height–LVM relationship in Asklepios reference participants assessed with nonlinear regression with and without accounting for the confounding effect of sex. The red line represents the body height–LVM relationship in men. The blue line represents the body height–LVM relationship in women. The black line represents the exaggeration of nonlinearity in the height–LVM relationship when the confounding effect of sex is neglected. This particularly leads to estimation problems at the extremes.
Both sexes. However, when an allometric exponent is computed for males and females considered together (thick black line) without adjustment for gender, there is an exaggeration of nonlinearity in the height–LVM relationship (allometric index 2.7). This has important clinical and epidemiological implications, resulting in marked overestimation of the prevalence of LV hypertrophy in short subjects and a marked underestimation in tall subjects. The appropriate indexation remains an issue of contention.

**Effect of age**
LV volumes are inversely associated with age. LVM decreases with age as well, albeit to a more limited extent than volume. As a consequence, RWT and M/V ratio increase. There is an age-related development of a concentric remodelling (see the Identification of LV Geometric Patterns section) with systolic and diastolic dysfunction.

**Effect of exercise and sport**
Isotonic exercise involves movement of large muscle groups. The profound vasodilatation of the skeletal muscle vasculature that is involved produces hypertrophy by increasing venous return to the heart and volume overload. This hypertrophy is characterized by chamber enlargement and a proportional change in wall thickness, with no changes in RWT. In contrast, isometric or static exercise involves developing muscular tension against resistance with little movement. Reflex and mechanical changes cause a pressure load on the heart rather than a volume load resulting in a slightly enlarged ventricle with increased RWT hypertrophy.

**Effect of obesity and diabetes**
Obesity is associated with increased LV volumes, increased LVM, and most typically increased RWT. In the Framingham study, an increase of body mass index over time was closely related to increased LVM and volumes. Insulin resistance, metabolic syndrome, and diabetes mellitus type II are similarly associated with increased LVM, RWT, and diastolic dysfunction. Diabetes patients have decreased systolic function as well. Correction of LVM for height preserves both the effects of obesity and elevated blood pressures on LVM. In contrast, correction of LVM for body surface area (BSA) effectively corrects for not only height but also obesity-related LV hypertrophy, which will remain undetected.

**Inherited and ethnic contributions**
Some of the variance in LV dimensions and mass may be explained by heredity, independent of the effects of sex, age, body size, blood pressure, heart rate, medications, and diabetes. Familial patterns of LV geometry were observed in subsequent generations of the Framingham study, but not in spouses. The greatest inheritable risk was found for concentric remodelling.

Normal ranges of LVM differ across races, being larger in African-Americans than in white Americans and/or Hispanics and smaller in Asian-Americans. Within one ethnicity, differences also exist between populations, e.g. Scandinavians being different from Mediterraneans. Only a part of these differences is accountable to ethnic variation in body size, and can be corrected by scaling. It is still unclear to what extent ethnic differences prevail when scaling for fat-free mass. It remains to be clarified to what extent these ethnic and population differences include a different prognosis and how to integrate ethnicities and populations in the definition of hypertrophy. At present, normal values and cutoffs should be adapted for each population.

**LV hypertrophy due to increased load**
Two basic patterns of cardiac hypertrophy occur in response to haemodynamic overload. In pressure overload (e.g., hypertension), pressure elevation most commonly leads to an increase in wall thickness and RWT, a phenomenon known as concentric remodelling (see the Identification of LV Geometric Patterns section). Eventually, an increase in systolic wall stress leads to concentric hypertrophy, caused by the addition of sarcomeres in parallel (hence, widening the cardiac myocytes), an increase in myocyte cross-sectional area, and an increase in LV wall thickening. In the Framingham Heart study, hypertensive patients had a greater increase in LVM and volume, and a smaller age-related reduction in LV size than individuals with normal blood pressure. In contrast, eccentric hypertrophy due to volume overload (e.g. with mitral regurgitation) is caused by increased diastolic wall stress. This leads to an increase in myocyte length with the addition of sarcomeres in series (hence, lengthening of cardiac myocytes), thereby engendering LV enlargement.

**Adaptation of LV function to increased load**
The complex changes that occur in the heart during LV remodelling cause alterations in LV size and geometry, but the process of LV remodelling also leads to alterations in contraction and relaxation, the volume of myocyte and non-myocyte components of the myocardium, the properties of the myocyte (sarcomeres, e.g. titin), and the extracellular matrix (balance of collagen types I and III, and collagen fraction). Diastolic function is influenced by alterations in LV systolic function and geometry, delayed myocardial relaxation, increased passive stiffness of the sarcomere and extracellular matrix, and altered myocardial tone.

Cardiac myocyte hypertrophy leads to foetal gene reactivation and decreased expression of a number of genes normally expressed in the adult heart. Depending on age, sex, duration of hypertension, severity, and treatment, differing cellular and molecular events may underlie the evolution from a ventricle with concentric hypertrophy to a more dilated failing ventricle (often presenting as HFpEF, heart failure reduced ejection fraction) or to a heavily fibrotic and non-dilated ventricle (presenting as HFrEF, heart failure preserved ejection fraction), according to the three stages in the hypertrophic process (overload, hypertrophy, and failure). Physiological hypertrophy (growth, pregnancy, and exercise) is characterized by normal organization of cardiac structure and normal or enhanced cardiac function, whereas pathological hypertrophy is commonly associated with upregulation of foetal genes, fibrosis, cardiac dysfunction, and increased mortality. The continuous vs. intermittent nature of overload in the settings of pathological and physiological hypertrophy is unlikely to account for the differences in response. In contrast to early-systolic load, late-systolic load delays myocardial relaxation and induces more maladaptive hypertrophy.
Morphology of the hypertensive heart

LV morphology

LV hypertrophy is defined on a normative basis; a definition based on 2 SD above the mean LVM in the general population will differ from a definition based on the healthy population without obesity or hypertension. Separate cutoffs are required for men and women. If LVM is corrected for BSA, it should be recognized that this corrects for obesity-related LVM, or for height. In the end-stage hypertensive heart, there is an increase in LV volumes and sphericity, a decrease in stroke volume, and finally a reduction in EF.

LA morphology

Left atrial (LA) volume may be calculated by either area-length or modified Simpson’s methods, and is usually scaled for BSA and expressed in mL/m²; the normal range is up to and including 34 mL/m². As with the LV, scaling by BSA corrects for an obesity-related increase in LA size that as a consequence will remain undetected. The LA is not symmetrical, and enlargement may occur non-uniformly, predominantly in one direction. Consequently, LA size is much better evaluated with 2D- or 3D-based LA volume rather than with M-mode. In hypertension and other situations where diastolic dysfunction occurs, reduction in early diastolic emptying is compensated by forceful atrial contraction. In addition, intermittent or permanent elevation of LV filling pressures leads to overfilling of the LA. The resulting LA enlargement is the ‘morpho-physiologic expression’ of chronic diastolic dysfunction, hypothesized to reflect the duration and severity of increased LA pressure. Although the presence of atrial fibrillation itself contributes to atrial size, LA enlargement is a well-known independent determinant of stroke, cardiovascular events, and death. Moreover, atrial fibrosis may be another endpoint of this process, predisposing to atrial remodelling and dysfunction with atrial fibrillation. This is a common endpoint that may be initiated by a number of aetiologies, including hypertension and diabetes mellitus. The main determinants of an increasing atrial size with age are the cardiovascular risk factors of elevated blood pressure and obesity. In hypertensive patients, LA enlargement is related to LVM (rather than the type of LV hypertrophy), overweight, higher fasting glucose, and metabolic syndrome.

Measurement of LVM

Linear echocardiographic dimensions

Acquisition and measurements

The measurement of LVM requires accurate measurements of wall thickness and chamber dimensions, as described in the Chamber Quantification update. The linear measurements of LV internal dimension (LVDd), septal (IVS), and PW are made from the parasternal long-axis acoustic window at the level of the LV minor axis, approximately at the mitral valve leaflet tips. M-mode recordings have excellent temporal resolution, and may be chosen from 2D images. However, even when directed by 2D guidance, it may not be possible to align the M-mode cursor perpendicular to the long axis of the ventricle (Figure 2). Software has been developed to...
reconstruct anatomical M-mode images from 2D images (Figure 3), but this is not yet universally available. Reference normal values for LV linear measurements are published in the Chamber Quantification update.29 Alternatively, chamber dimension and wall thicknesses can be acquired from the parasternal short-axis view using direct 2D measurements. The use of 2D-derived linear dimensions overcomes the common problem of oblique parasternal images resulting in overestimation of cavity and wall dimensions from M-mode (Figure 4).

When 2D measurements are used, the wall thicknesses and linear dimensions should be measured at the level of the LV minor dimension, at the mitral leaflet tips level. The upper limit of normal for LVDD is smaller than the M-mode measurement. Left ventricle internal dimension diastole (LVIDd), inter-ventricular septum diastole (IVSd), and posterior wall diastole (PWd) are measured at end-diastole from 2D or M-mode recordings, preferably on several beats.

Understanding the LVM literature is facilitated by recognizing various methods:

(i) The original American Society of Echocardiography (ASE) approach recommended that dimensions be measured from the leading edge to the leading edge of echocardiographic borders. This results in the inclusion of endocardial echoes from the IVS andPW, and the exclusion of endocardial echoes from the LVIDd.33 This was because the trailing edge of endocardial signals is dependent on gain settings. This may impact on LVM measurements, especially at the upper and lower extremes of these measurements.34 The simplified calculation of LVM with this approach is $LVM = 1.04[(IVS + LVIDd + PW)^3 - (LVIDd)^3] + 0.6 \, g$.

(ii) The subsequent Penn convention excluded endocardial echoes from IVS and PW dimensions, but included endocardial echoes in measurement of the LVIDd.35 As the Penn convention gives larger cavity dimensions and smaller wall thicknesses than the ASE convention, the use of this approach necessitates subtraction of 13.6 from the previous mass calculation.

(iii) The current ASE/European Association of Cardiovascular Imaging (EACVI) Chamber Quantitation Guidelines point out that refinements in image processing have allowed measurement of the actual visualized thickness of the ventricular septum and other chamber dimensions as defined by the actual tissue–blood interface, rather than the distance between the leading edge echoes, which had previously been recommended (Figure 5).29

All LVM algorithms (M-mode, 2D, or 3D echocardiographic measurements) are based on subtraction of the LV cavity volume

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**Figure 3** Reconstruction of anatomical M-mode images from 2D images. Overestimation of LV dimensions can occur through tangential imaging at an angle to the appropriate axis (A). When the echo window cannot be moved, an alternative means of obtaining accurate data may be provided by reconstructing the M-mode dataset from the 2D image—so-called anatomical M-mode (B). In this example, a small (1 mm) difference in LV dimension results in a 5 g difference in LVM. Tangential imaging may not just relate to selection of a longer than expected cross-section—it may underestimate the measurement by failure of the beam to pass through the axis of the ventricle (C). Again, the use of anatomical M-mode imaging may circumvent this problem (D).
from the volume enclosed by the LV epicardium to obtain the volume of the shell between the LV cavity and the epicardial surface. This shell volume is then converted to mass by multiplying LV wall volume by the specific gravity of myocardium (1.05 g/mL). The formula used for estimation of LVM from LV linear dimensions is based on modeling the LV as a prolate ellipse, and assumes that the major/minor axis ratio is 2 : 1: LVM = 0.8 × {1.04[(LVIDd + PW + IVSd)3 − (LVIDd)3]} + 0.6 g. Extensive validation of this formula has been performed from necropsy specimens.

Normal values
Table 1 summarizes the reported range of normal values for LVM by M-mode echocardiography.3,37–45 These values differ between men and women, with the latter systematically lower than the former, even when indexed for BSA (Table 1; see the section below—methods of indexation). The upper limits of normal ranges in the ASE chamber quantification update are > 95 g/m² (> 44 g/ht².7) in women and > 115 g/m² (> 48 g/ht².7) in men.29

Limitations
There are four principal limitations in the calculation of LVM using linear methods:

(i) The ‘Cube’ formula is not accurate in patients with major distortions of LV geometry (e.g. apical aneurysm, or any condition where the 2 : 1 axis ratio requirement is not met).
(ii) Because this formula involves cubing primary measurements, even small errors in these measurements may be magnified.
(iii) These measurements are insensitive to small changes in mass.
(iv) The measurements are highly dependent on imaging quality and observer expertise.

Two-dimensional echocardiography
The most commonly used 2D methods for measuring LVM are based on the area-length formula and the truncated ellipsoid
model, as described in detail in the previous ASE/EACVI chamber quantification document. In the presence of shape distortions, such as that caused by post-myocardial infarction (MI) remodelling, the geometric assumptions inherent in this approach remain problematic. Both methods were validated in the early 1980s in animal models and by comparing premorbid echocardiograms with measured LV weight at autopsy in human beings. Normal values are summarized in Table 2, and the degrees of abnormality are classified in Table 3. The main limitations relate to image quality and the temporal resolution of 2D imaging, compared with M-mode echocardiography. The limitations of M-mode regarding geometrical assumptions and the impact of small error on measurements are also applicable to 2D measurements. In addition, 2D imaging leads to frequent foreshortening due to inappropriate cut-planes.

Three-dimensional echocardiography

The benefit of three-dimensional echocardiography (3DE) is especially to obviate inaccurate geometric assumptions, inherent to 2D, that become exaggerated in remodelled ventricles. 3DE is a potentially attractive modality for the measurement of LVM, and normal ranges have been developed. The accuracy of 3DE is reportedly similar to cardiac magnetic resonance (CMR) imaging methods for measuring LVM. However, there are wide limits of agreement which primarily relate to difficulties in accurately tracing the LV epicardial border, particularly in dilated ventricles, and generally show that while 3DE is imperfect for LVM estimation—with a tendency to underestimate LVM compared with CMR imaging in patients with cardiac disease—the accuracy is more favourable than with alternative ultrasound methods. Normal values of M-mode, 2D mass, and 3D mass are given in Tables 1 and 2. Degrees of abnormality of LVM are summarized in Table 3 and the validation of all methods against reference techniques is summarized in Table 4. A later section describes the use of 2D and 3D for the assessment of LV function.

Identification of LV geometric patterns

While patients with early hypertensive disease will most likely have normal LV geometry, longstanding or untreated hypertension will result in changes in LV shape and eventually, a deterioration of systolic function. Broadly, the changes in LV geometry can be classified according to whether LVM is normal or increased and whether ventricular morphology (RWT) is altered (Table 5). RWT is variably reported as (PW + 2)/LVd or (IVS + PW)/LVd, of which we favour the former because septal measurements may be confounded by the presence of septal bulge. RWT is problematic and not reflective of true LV geometry in patients with asymmetric hypertrophy. The upper limit of normal RWT is 0.42.

Concentric LV hypertrophy

Concentric LV hypertrophy, probably most commonly associated with hypertension, is characterized by normal cavity size, uniformly increased LV wall thickness, and increased LVM (Figures 7 and 8). Cutoff values adopted by the ASE and EACVI are based on either overall LVM (g), LVM/BSA (g/m²), LVM/height (g/m), or LVM/height² (g/m²²) and while each has been shown to have limitations of either under- or overestimating LVM, each has been used successfully in characterizing LV hypertrophy in different patient populations.

Concentric LV hypertrophy is an adaptive response to high systemic pressure caused by hypertension or diseases such as aortic stenosis, coupled with high peripheral resistance. Concentric LV hypertrophy (LVH) and changes in LV geometry have been shown to affect both men and women regardless of age, and are also associated with changes in diastolic function, longitudinal and radial myocardial function, and atrial size.

Eccentric LV hypertrophy

In contrast to concentric LVH, eccentric hypertrophy is associated with volume, rather than pressure overload. This is usually due to significant valvular regurgitation or high cardiac index, as is seen in elite athletes (although eccentric hypertrophy may be the consequence of strength training). Systemic pressure is normal and peripheral resistance is not increased in patients with eccentric hypertrophy. Echocardiographic findings show increased LV cavity size, normal LV wall thickness, and increased LVM (Figure 9).

Patients with eccentric hypertrophy share similar changes in diastolic function and longitudinal and radial function as those with concentric hypertrophy. Unlike concentric hypertrophy, however, patients with eccentric LVH generally have low normal or mildly impaired systolic function due to chronic volume overload.

Changes in LV shape associated with LV enlargement have been quantified as sphericity index. This is a ratio between measured end diastolic volume (EDV) (preferably with 3DE) and a spherical volume based on the longitudinal dimension of the LV (4/3 × π × D²). This parameter has been shown to be a predictor of remodeling, but this is more in the setting of LV dysfunction after MI than in hypertensive heart disease.

Concentric remodelling

Concentric LV remodelling is a late stage response of the LV and can be caused by chronic pressure, volume overload, or MI. It is most commonly associated with coronary artery disease, but is also associated with longstanding hypertension, especially untreated hypertension. Like eccentric hypertrophy, it is also associated with LV systolic dysfunction. Echocardiographic features show normal or small LV cavity size, usually increased LV wall thickness and normal
<table>
<thead>
<tr>
<th>Source</th>
<th>Year</th>
<th>Men</th>
<th>Women</th>
<th>Age (years)</th>
<th>Body size indexation</th>
<th>Measurement convention</th>
<th>LVM Men</th>
<th>LVM Women</th>
<th>Upper limit of LVMI Men</th>
<th>Upper limit of LVMI Women</th>
<th>Basis for upper limits</th>
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<tbody>
<tr>
<td>Henry et al.</td>
<td>1980</td>
<td>78</td>
<td>58</td>
<td>20–97</td>
<td>None</td>
<td>ASE</td>
<td>160 ± 25 g</td>
<td>17 g/m²</td>
<td>210 ± 25 g (140 g/m²)</td>
<td>210 ± 25 g (140 g/m²)</td>
<td>95% CL</td>
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<td>Devereux et al.</td>
<td>1981</td>
<td>106</td>
<td>120</td>
<td>39 ± 13</td>
<td>BSA</td>
<td>Penn</td>
<td>89 ± 21</td>
<td>19 g/m²</td>
<td>136 ± 21 g</td>
<td>136 ± 21 g</td>
<td>97th percentile</td>
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<td>Hammond et al.</td>
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<td>83</td>
<td>77</td>
<td>44 ± 13</td>
<td>BSA</td>
<td>Penn</td>
<td>155 ± 50 g</td>
<td>193 ± 23 g</td>
<td>134 ± 50 g</td>
<td>134 ± 50 g</td>
<td>Comparison with hypertensive population: LV determination</td>
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<td>Byrd et al.</td>
<td>1985</td>
<td>44</td>
<td>40</td>
<td>35 ± 10</td>
<td>BSA</td>
<td>–</td>
<td>148 ± 26 g</td>
<td>108 ± 21 g</td>
<td>200 g</td>
<td>200 g</td>
<td>95th percentile</td>
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<td>Levy et al.</td>
<td>1987</td>
<td>347</td>
<td>50</td>
<td>43 ± 12</td>
<td>Ht/BSA</td>
<td>ASE</td>
<td>208 ± 43 g</td>
<td>145 ± 27 g</td>
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<td>294 g</td>
<td>M + 2 SD</td>
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<td>Koren et al.</td>
<td>1991</td>
<td>167</td>
<td>86</td>
<td>47 ± 13</td>
<td>BSA</td>
<td>Penn</td>
<td>–</td>
<td>–</td>
<td>125 g</td>
<td>125 g</td>
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<td>de Simone et al.</td>
<td>1992</td>
<td>137</td>
<td>91</td>
<td>39 ± 14</td>
<td>None</td>
<td>Penn</td>
<td>155 ± 34 g</td>
<td>117 ± 28 g</td>
<td>223 g</td>
<td>223 g</td>
<td>M + 2 SD</td>
</tr>
<tr>
<td>Kuch et al.</td>
<td>2000</td>
<td>213</td>
<td>291</td>
<td>42 ± 12</td>
<td>None</td>
<td>Penn</td>
<td>89 ± 21 g</td>
<td>92 ± 18 g</td>
<td>127 ± 21 g (106 g/m²)</td>
<td>127 ± 21 g (106 g/m²)</td>
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<tr>
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<td>2001</td>
<td>651</td>
<td>1066</td>
<td>72 ± 5</td>
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<td>CV Health Study (Healthy Substudy)</td>
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<td>93</td>
<td>213</td>
<td>75 ± 4</td>
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<td>ASE</td>
<td>96 ± 27 g/m²</td>
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<td>Asklepios—total population</td>
<td>2007</td>
<td>1301</td>
<td>1223</td>
<td>46 (41–51)</td>
<td>None</td>
<td>Height, Ht, Height, BB</td>
<td>175 ± 39 g</td>
<td>121 ± 32 g</td>
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<td>243 g</td>
<td>95th percentile</td>
</tr>
<tr>
<td>Asklepios Healthy, Risk factor deprived</td>
<td>2007</td>
<td>198</td>
<td>414</td>
<td>43 (39–48)</td>
<td>None</td>
<td>Height, Ht, Height, BB</td>
<td>155 ± 36 g</td>
<td>108 ± 21 g</td>
<td>214 g</td>
<td>214 g</td>
<td>95th percentile</td>
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<td>CHS healthy subgroup: no prevalent HF, CVD, hypertension, obesity, or subclinical heart disease (i.e. normal aortic augmentation index and normal carotid intima-media thickness).</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
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</table>

Table I Normal limits of M-mode LVM
LVM (Figures 7 and 10). Concentric remodelling is also associated with changes in the shape of the LV—e.g. LV sphericity changes—and becomes more rounded, rather than bullet shape. The result of this is more dramatic degradation of diastolic function and loss of radial and longitudinal function.

Other classification

The limitation of the classical categories is the suboptimal categorization of dilated ventricles. Recently, Gaasch and Zile proposed a subdivision based on LVM (vertical axis), LV volume (horizontal axis), and RWT or M/V, represented by the oblique lines indicating the upper (full) and lower (dashed) limit of normality (Table 6 and Figure 7). Using this approach, the non-dilated ventricle is characterized as having normal morphology, concentric remodelling, or concentric hypertrophy, based on LVH and RWT (>0.42). Dilated ventricles without LVH are described as having eccentric remodelling if the RWT is <0.32. Dilated ventricles with LVH are described as having eccentric hypertrophy (RWT <0.32), mixed hypertrophy (RWT >0.42), or physiological hypertrophy (RWT 0.32–0.42). The resulting categories yield distinct functional behaviours and prognoses.

Natural history of LV geometry in hypertension

Left ventricular hypertrophy is caused by increased wall stress, either due to chronic pressure overload, as seen in hypertension, or the volume overload seen in valvular disease. However, in early, mild hypertension, LVH is usually absent and the first manifestation of
hypertension is diastolic dysfunction.\textsuperscript{58,62} This can be detected as grade 1 diastolic impairment, or impaired relaxation. Over time however, if left untreated, filling pressures continue to rise, ventricular hypertrophy develops as an adaptive response to chronic pressure, and more severe disturbances of diastolic filling are more commonly encountered. Eventually, LV remodelling will occur and left ventricular systolic function will become impaired. While the goal of hypertension management is to prevent any changes in LV

### Table 3 Degrees of abnormality of LVM

<table>
<thead>
<tr>
<th></th>
<th>Women</th>
<th>Mildly abnormal</th>
<th>Moderately abnormal</th>
<th>Severely abnormal</th>
<th>Men</th>
<th>Mildly abnormal</th>
<th>Moderately abnormal</th>
<th>Severely abnormal</th>
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<tr>
<td></td>
<td>Range</td>
<td></td>
<td></td>
<td></td>
<td>Range</td>
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<td></td>
</tr>
<tr>
<td>LV, g</td>
<td>67–162</td>
<td>163–186</td>
<td>187–210</td>
<td>( \geq 211 )</td>
<td>88–224</td>
<td>225–258</td>
<td>259–292</td>
<td>( \geq 293 )</td>
</tr>
<tr>
<td>LV/BSA, g/m(^2)</td>
<td>43–95</td>
<td>96–108</td>
<td>109–121</td>
<td>( \geq 122 )</td>
<td>49–115</td>
<td>116–131</td>
<td>132–148</td>
<td>( \geq 149 )</td>
</tr>
<tr>
<td>LV/height, g/m</td>
<td>41–99</td>
<td>100–115</td>
<td>116–128</td>
<td>( \geq 129 )</td>
<td>52–126</td>
<td>127–144</td>
<td>145–162</td>
<td>( \geq 163 )</td>
</tr>
<tr>
<td>LV/height(^2), g/m(^3)</td>
<td>18–44</td>
<td>45–51</td>
<td>52–58</td>
<td>( \geq 59 )</td>
<td>20–48</td>
<td>49–55</td>
<td>56–63</td>
<td>( \geq 64 )</td>
</tr>
<tr>
<td>Relative wall thickness, cm</td>
<td>0.22–0.42</td>
<td>0.43–0.47</td>
<td>0.48–0.52</td>
<td>( \geq 0.53 )</td>
<td>0.24–0.42</td>
<td>0.43–0.46</td>
<td>0.47–0.51</td>
<td>( \geq 0.52 )</td>
</tr>
<tr>
<td>Septal thickness, cm</td>
<td>0.6–0.9</td>
<td>1.0–1.2</td>
<td>1.3–1.5</td>
<td>( \geq 1.6 )</td>
<td>0.6–1.0</td>
<td>1.1–1.3</td>
<td>1.4–1.6</td>
<td>( \geq 1.7 )</td>
</tr>
<tr>
<td>Posterior wall thickness, cm</td>
<td>0.6–0.9</td>
<td>1.0–1.2</td>
<td>1.3–1.5</td>
<td>( \geq 1.6 )</td>
<td>0.6–1.0</td>
<td>1.1–1.3</td>
<td>1.4–1.6</td>
<td>( \geq 1.7 )</td>
</tr>
<tr>
<td>2D method</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LV, g</td>
<td>66–150</td>
<td>151–171</td>
<td>172–182</td>
<td>( \geq 193 )</td>
<td>96–200</td>
<td>201–227</td>
<td>228–254</td>
<td>( \geq 255 )</td>
</tr>
<tr>
<td>LV/BSA, g/m(^2)</td>
<td>44–88</td>
<td>89–100</td>
<td>101–112</td>
<td>( \geq 113 )</td>
<td>50–102</td>
<td>103–116</td>
<td>117–130</td>
<td>( \geq 131 )</td>
</tr>
</tbody>
</table>

BSA, body surface area; LV, left ventricular; 2D, two-dimensional. Bold italic values: recommended and best validated.

### Table 4 Correlation of all echocardiographic methods of LVM calculation vs. MRI

<table>
<thead>
<tr>
<th></th>
<th>End-diastole</th>
<th>End-systole</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>( r )</td>
<td>SEE (g)</td>
</tr>
<tr>
<td>1D Echo-Penn vs. CMR</td>
<td>0.725</td>
<td>25.6</td>
</tr>
<tr>
<td>2D Echo-AL vs. CMR</td>
<td>0.694</td>
<td>24.2</td>
</tr>
<tr>
<td>2D Echo-TE vs. CMR</td>
<td>0.687</td>
<td>21.8</td>
</tr>
<tr>
<td>3D Echo-PSR vs. CMR</td>
<td>0.882</td>
<td>10.4</td>
</tr>
</tbody>
</table>

CMR, magnetic resonance imaging; 1D Echo-Penn, M-mode echocardiographic method (Penn convention); 2D Echo-AL, two-dimensional echocardiographic area-length method; 2D Echo-TE, two-dimensional echocardiographic truncated ellipsoid method (8); 3D Echo-PSR, three-dimensional echocardiographic polyhedral surface reconstruction method.

### Table 5 Classical description of LV geometry

<table>
<thead>
<tr>
<th>LV geometry</th>
<th>LVM</th>
<th>RWT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal</td>
<td>( \leq 115 \text{ g/m}^2 ) (men) or ( \leq 95 \text{ g/m}^2 ) (women)</td>
<td>&lt;0.42</td>
</tr>
<tr>
<td>Concentric hypertrophy</td>
<td>( &gt;115 \text{ g/m}^2 ) (men) or ( &gt;95 \text{ g/m}^2 ) (women)</td>
<td>&gt;0.42</td>
</tr>
<tr>
<td>Eccentric hypertrophy</td>
<td>( &gt;115 \text{ g/m}^2 ) (men) or ( &gt;95 \text{ g/m}^2 ) (women)</td>
<td>&lt;0.42</td>
</tr>
<tr>
<td>Concentric remodelling</td>
<td>( \leq 115 \text{ g/m}^2 ) (men) or ( \leq 95 \text{ g/m}^2 ) (women)</td>
<td>&gt;0.42</td>
</tr>
</tbody>
</table>

Measurements performed using 2D-directed M-mode.\textsuperscript{29}
geometry, the current ability of echocardiography to provide serial assessment of the LV response in the individual patient is compromised by variability of LVM measurement.

**Recommendations**

Description of LV geometry, using at the minimum the four categories of normal geometry, concentric remodelling, and concentric and eccentric hypertrophy, should be a standard component of the echocardiography report.

**Tissue characterization**

The haemodynamic disturbances and humoral stimulation that lead to the cardiac responses to hypertension\(^6^3\) do not necessarily progress in parallel.\(^6^4\) While measurement of LVM addresses the response to haemodynamic disturbance, this may not necessarily reflect the full physiological impact of hypertension on the heart. While not part of current guidelines, tissue characterization may provide information about myocardial remodelling, and allow targeted therapy against molecular changes, sarcoplasmatic failure, apoptosis, fibrosis, and disturbances of vascular structure and function.\(^6^5\)

Interstitial, perivascular, plexiform, and replacement fibrosis of necrotic tissue\(^6^6\) are likely responsible for disturbances of myocardial perfusion, synchrony, and rhythm.

An important reason for attempting to characterize myocardial tissue is that not all increments in LVM that occur in the setting of hypertensive heart disease are due to hypertension. The recognition of other causes of increased wall thickness, including athletic hypertrophy, valvular disease, infiltrative disorders (amyloid, Friedrich’s ataxia, and Fabry’s disease), non-compaction, and hypertrophic cardiomyopathy,\(^6^7\) has important treatment implications.

Tests of myocardial tissue characterization can be divided into processes that measure tissue reflection (and therefore tissue density), and functional changes that are due to the dynamic consequences of changes in myocardial ultrastructure (which are discussed in the section on LV function). The only echocardiographic marker of tissue density is integrated backscatter, a measure of ultrasonic scatter from small reflectors, which relate to tissue density.\(^6^8\) Calibrated integrated backscatter refers to a method whereby the amplitude of reflection is measured in relation to the amplitude deriving from a reference tissue, for example, blood within the LV cavity or the pericardium. The primary determinant of both scatter and attenuation in myocardial tissue is collagen.\(^6^9\) However, as scatter is also related to position and orientation of myofibrils relative to the ultrasound beam, variations in these measurements are not specific for...
Figure 9  Eccentric LVH. Parasternal long-axis view (left) and apical four-chamber view (right) of a 28-year-old female patient with a failed mitral valve repair showing eccentric LVH. LVDd 56 mm; LVDs 39 mm; IVS 12 mm; PW 12 mm; EF 50%; LVM 206 g.

Figure 10  Concentric LV remodelling. A 59-year-old male patient with concentric LV remodelling. LVEDD 47 mm; LVESD 36 mm; IVS 20 mm; PW 11 mm; EF 43%; LVM 270 g.

Table 6  Characterization of LV geometry based on LVM (vertical axis), LV volume (horizontal axis), and RWT, measured using 2D-directed M-mode

<table>
<thead>
<tr>
<th>LV geometric pattern</th>
<th>LV volume index (mL/m²)</th>
<th>LVM index (g/m²)</th>
<th>RWT</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal ventricle</td>
<td>≤75</td>
<td>≤115 (men) or ≤95 (women)</td>
<td>0.32–0.42</td>
</tr>
<tr>
<td>Physiological hypertrophy</td>
<td>&gt;75</td>
<td>&gt;115 (men) or &gt;95 (women)</td>
<td>0.32–0.42</td>
</tr>
<tr>
<td>Concentric remodelling</td>
<td>≤75</td>
<td>≤115 (men) or ≤95 (women)</td>
<td>&gt;0.42</td>
</tr>
<tr>
<td>Eccentric remodelling</td>
<td>&gt;75</td>
<td>≤115 (men) or ≤95 (women)</td>
<td>≤0.32</td>
</tr>
<tr>
<td>Concentric hypertrophy</td>
<td>≤75</td>
<td>&gt;115 (men) or &gt;95 (women)</td>
<td>&gt;0.42</td>
</tr>
<tr>
<td>Mixed hypertrophy</td>
<td>&gt;75</td>
<td>&gt;115 (men) or &gt;95 (women)</td>
<td>&gt;0.42</td>
</tr>
<tr>
<td>Dilated hypertrophy</td>
<td>&gt;75</td>
<td>&gt;115 (men) or &gt;95 (women)</td>
<td>0.32–0.42</td>
</tr>
<tr>
<td>Eccentric hypertrophy</td>
<td>&gt;75</td>
<td>&gt;115 (men) or &gt;95 (women)</td>
<td>≤0.32</td>
</tr>
</tbody>
</table>
fibrosis and backscatter and attenuation is also affected by angle of 
insolation (i.e. the same myocardial segment will have different ultra-
sonic characteristics when viewed from orthogonal windows—
parasternal long axis vs. apical) due to the alignment of myofibrils per-
pendicular or parallel to the ultrasound beam. Thus, feasibility can be 
limited. Moreover, these changes in early hypertensive heart 
disease may be subtle.71

Two other imaging methods are probably superior to echocar-
diography for myocardial tissue characterization. Late gadolinium en-
hancement with CMR has become widely used for the recognition 
of replacement fibrosis in ischaemic heart disease.72 The same has 
also been helpful in understanding the contribution of fibrosis in 
hypertrophy, where ~50% of patients with hypertensive LVH mani-
fest patchy late enhancement,73 which correlates with the presence 
of diastolic dysfunction.74 The problem with this technique is that it is 
based on defining a reference normal segment within the myocar-
dium, so it may be misleading for the detection of diffuse interstitial 
fibrosis. A potential solution is the use of T1 mapping, which allows 
the recognition of differences in T1 relaxation between the normal 
and fibrotic myocardium. Recent work has validated T1 mapping as an 
accurate marker of the extent of diffuse fibrosis.75 The 
final methods that are used in tissue characterization are ‘cardiac 
nuclear imaging’ procedures for molecular imaging of collagen76 
and detection of apoptosis.77

Other echocardiographic markers—for example tissue Doppler 
and strain—have been used as markers of fibrosis.66 It should be 
recognized that these functional parameters may be confounded by 
myocardial processes that parallel the development of fibrosis, 
and may not be optimal for this purpose.78

### Recommendations

**Myocardial characterization using CMR can identify non-hypertrophic 
causes of LV thickening.** It should be considered when (i) the degree 
of LV thickening is at least moderate, (ii) severity of the LV thickening 
is inconsistent with the severity of hypertension, (iii) there is evidence 
of LV dysfunction despite appropriate BP control, (iv) other features 
rise the prospect of an infiltrative process (severe thickening, 
alteration of tissue density on fundamental imaging, or e’ velocity 
< 5 cm/s).

### Arterial function and ventriculo-arterial matching

#### Arterial function

**Arterial afterload**

Arterial afterload is characterized by both steady and pulsatile compo-
nents of blood pressure.79 This parameter is determined by imped-
ance, compliance, or resistance, derived from aortic pressure ($P_{ao}$) 
and flow waveforms ($F_{ao}$), both of which can be assessed non-invasively 
by means of applanation tonometry and ultrasound, respectively.

A variety of measurements have been created to better under-
stand the process of displacement of blood from the LV into the ar-
terial tree (Figure 11). If the arterial system was composed of rigid 
tubes without any storage capacity, blood would be accelerated in 
systole throughout the complete arterial tree, which would give 
rise to very large intra-arterial pressure differences (and a high 
load on the heart). Owing to the elasticity of the large arteries, 
however, part of the stroke volume is locally stored in the aorta in 
systole (the ‘windkessel’ function), buffering the pulsatility of blood 
flow and providing a more continuous blood flow in the distal circu-
lation. This reduces the importance of inertial forces. Characteristic 
impedance ($Z_c$) reflects the interplay between these inertial effects 
and the local storage of blood in the proximal aorta and the load ini-
tially experienced by the ventricle upon opening of the aortic valve. It 
is calculated by plotting the relation of time-varying $P_{ao}$ (aortic pres-
sure) vs. time-varying $F_{ao}$ (aortic flow) during the ejection phase of 
the cardiac cycle; the slope provides $Z_c$ [in mmHg/(mL/s)]. This par-
arameter is dependent on blood pressure and aortic size; a stiff and 
narrow aorta leads to high $Z_c$, a distensible, wide aorta to a low $Z_c$. 
While $Z_c$ determines the upstream of pressure, pulse pressure is 
mainly determined by the total arterial compliance (TAC) of the ar-
terial tree in combination with the systemic vascular resistance (SVR). 
The simplest approximation of TAC is the ratio of the stroke volume 
and pulse pressure (mL/mmHg), although this leads to systematic 
overestimation. TAC is highly size-dependent, depends non-linearly 
on arterial pressure, and that there are systematic differences 
between different methods, making TAC a parameter difficult to 
standardize (Figure 11).

#### Arterial afterload: pulse wave velocity and wave reflection

The above section simplifies the arterial system to a simple ‘windkes-
sel’ system. Cardiac contraction gives rise to pressure- and flow 
waves travelling through the arterial tree. The stiffer the arteries, 
the higher the pulse wave velocity (PWV; Figure 12). PWV is propor-
tional to the intrinsic mechanical properties of the arterial wall 
(stress–strain relationships), the ratio of wall thickness to lumen 
diameter, and inversely proportional to the density of blood 
(which is virtually constant). Thus, PWV is independent of size 
and only varies with arterial remodelling or changes in arterial 
tissue properties (note that these are pressure-dependent). The 
carotid and femoral artery is the most commonly used measuring 
locations, with time delay derived from either pressure (tonometry), 
ultrasound- (pulsed Doppler), or CMR-based (phase contrast) 
signals. As the carotid and femoral artery is not along a single un-
equivocal trajectory, the latest consensus is that distance is approxi-
mated as 0.8 times the linear distance measured directly between the 
carotid and femoral sites. Age-specific normal values for carotid– 
femoral have been reported (Figure 13), but have the disadvantage 
of obscuring the important effect of age.81 Numerous studies have 
now demonstrated an association between increased arterial stiff-
ness and increased cardiovascular risk. Although PWV provides an 
overall estimate of the elastic properties of the aorta and central ar-
teries, it also depends on functional and dynamic properties, including 
production of nitric oxide. It is also possible to assess the local elastic 
properties at the carotid or femoral artery, and several ultrasound-
based techniques exist for this purpose (e.g. wall tracking to 
measure arterial distension) or are under investigation (pulse wave 
imaging and shear wave imaging).

Wave dynamics are too complex to resolve in full detail in an in 
vivo setting and are commonly simplified, considering only one forward 
(generated by the heart) and one backward wave (due to reflections
in the periphery). The timing and magnitude of these waves can directly be linked to cardiovascular pathophysiology. Recent studies have reported an association between augmentation index (a rather poor measure of wave reflection) and cardiovascular risk, although there is disagreement about the prognostic value of this information. An increased magnitude of wave reflection, measured with the wave decomposition technique, is an independent prognostic determinant of cardiovascular risk and a powerful and independent predictor of incident heart failure.

Ventriculo-arterial interaction

The classical approach to ventriculo-arterial matching

The most widespread paradigm for the assessment of ventricular–vascular coupling is the ventricular (Ees)—arterial (Ea) elastance framework, which links mechanical performance of the ventricle to its oxygen consumption.

For an efficient energy transfer, the LV should develop an elastance that is greater than the arterial elastance. Arterial elastance is commonly calculated as end-systolic pressure/stroke volume and is a measure of resistive, not pulsatile load. Ees is the end-systolic elastance (slope of the end-systolic pressure–volume relation), a measure of ventricular contractility. Ea stands for arterial elastance (ratio of end-systolic pressure and stroke volume), although it is an imperfect measure of arterial properties, being highly sensitive to the heart rate. Resting Ea/Ees ratios of ~0.62–0.82 are observed across species and in human populations. The LV generates maximal stroke work when Ea/Ees = 0.80, while it operates at maximal energetic efficiency with an Ea/Ees of 0.70. The normal Ea/Ees values seen in the Asklepios cohort and the Olmsted cohort suggest that normal subjects’ Ea/Ees values approximate this optimal value. Values >1 indicate an ‘ill-matched’ ventricle and arterial system. While the framework is essentially based on pressure–volume loop analysis—and hence restricted to an invasive setting—it has been simplified to make it suitable for application in clinical settings, approximating Ees as the ratio of end-systolic pressure and the end-systolic volume (ESV) or via the use of single-beat methods that take advantage of the relatively small variability in the shape of the normalized time-varying left ventricular elastance curve over the cardiac cycle.

Novel approaches to ventriculo-arterial matching

The standard Ea/Ees analysis does not involve any evaluation of time in the analysis. Using cardiac ultrasound and applanation tonometry (Figure 13), myocardial stress can be expressed as a function of time...
throughout systole. Peak stress occurs in early systole, before important contributions of reflected waves to central pressure and correlates directly with SVR and Zc. The greater peak and end-systolic wall stress and higher ejection phase stress-time integral in women may relate to the susceptibility of women to heart failure.

Wave intensity analysis is a new method of assessing ventriculo-vascular interaction. There are three aortic waves: (i) a wave reflecting LV contraction, generating a forward wave increasing pressure and flow; (ii) a reflected wave, generally increasing pressure and lowering blood flow, and (iii) a late-systolic wave due to LV relaxation, lowering blood pressure and flow. Current research is seeking whether this wave-based analysis can be used to quantify cardiac systolic and diastolic performance.

Assessment of the aorta

Hypertension is an important contributor to aortic disease, and any echocardiogram performed for the evaluation of end-organ disease should include assessment of the aorta. Echocardiographic views are usually limited to the ascending aorta between the coronary sinuses and main pulmonary artery, the aortic arch (in the suprasternal view), the descending aorta in the far-field of the parasternal, suprasternal, and foreshortened apical two-chamber view, and the abdominal aorta in the subcostal view. In particular, this simple step adds an incremental value in screening men > 65–70 years old for abdominal aortic aneurysm, especially if they are smokers. Coarctation of the aorta is a well-known structural abnormality that can lead to hypertension and LV hypertrophy and may go undetected by clinical assessment, particularly in younger adults. The echocardiogram is central to making this diagnosis, so younger patients presenting with hypertension should undergo 2D imaging, colour and Doppler assessment of the distal arch and upper descending aorta. Further information about echocardiography and aortic disease, including normal aortic dimensions, are described in the EACVI recommendations for clinical practice.

Recommendations

Blood pressure should be obtained at the time of the examination and integrated into the report.
Aortic dimensions should be reported in all studies of hypertensive subjects.
Measurement of pulse wave velocity should be considered as a marker of vascular health and risk in primary prevention patients.
Assessment of ventriculo-arterial mismatch is currently a research rather than a routine clinical investigation.
LV systolic function in hypertension

Parameters from linear measurements
LV linear dimensions for the calculation of LVM are widely used in the setting of hypertensive patients. The use of these measurements for the evaluation of endocardial fractional shortening (FS) has been superseded by more accurate and reliable measures. Likewise, the Teichholz or Quinones methods for measurement of EF from linear measurements are dependent on geometric assumptions and are not recommended.

Two-dimensional measurements
While the process of tracing LVM (above) and volumes are similar, the prognostic independence of LVM and function justifies their separation. The techniques and reference normal values for obtaining EF from tomographic 2D echocardiography are summarized in the Chamber Quantification update.29 The biplane method of discs (modified Simpson’s rule obtained from apical four- and two-chamber views) is the most accurate in abnormally shaped ventricles.46,91

In the pre-harmonic and pre-digital era, the main sources of inter-study variability included repeated echo recordings, repeated video measurements, and measurements made by different investigators.92 Similar analyses have not been performed by harmonic imaging, which may be an important distinction for two reasons. The use of lower frequencies (required for the creation of a wider broadband) implies a reduction of spatial resolution, with apparently thicker structures and potential effects on the measurement of wall thickness. On the other hand, the use of harmonic imaging improves the reproducibility of 2D LV volumes.93 When compared with CMR, 2D determination of LV volumes shows higher interstudy variability which reaches statistical significance for LV ESV (4.4–9.2% vs. 13.7–20.3%, $P < 0.001$).94 and results in higher calculated sample sizes (increases of 55–93% in comparison with CMR) to show clinically relevant changes in LV size.

The ejection phase indices (FS, EF, stroke volume, and cardiac output) cannot determine the relative contribution of each of these variables to LV pump function. In particular, load dependency of these parameters may induce inaccurate estimation of intrinsic myocardial contractility in chronic pressure overload conditions. The estimation of LV afterload may help in determining whether or not observed LV pump function is representative of actual myocardial contractile performance. The most direct measurement of LV afterload is end-systolic stress (ESS).95 Two main types of ESS can be measured, meridional and circumferential ESS (cESS), each acting as counter forces to fibre shortening.95 Longitudinal shortening of endocardial fibres is limited by longitudinal (meridional) ESS, which can be measured using a catheterization-validated formula which incorporates end-systolic LV internal diameter (LVIDs) and wall thickness coupled with simultaneous cuff blood pressure.36

Three-dimensional measurements
Assessment of LV volumes by 2DE is limited by foreshortening, malrotation, angulation, and a reliance on geometric assumptions for volumetric calculation, resulting in an underestimation of the true
volumes, particularly in remodelled ventricles.36,97 Transthoracic 3DE provides a rapid and accurate method for quantifying LV volumes and EF (LVEF).98,99 It has a superior reproducibility to 2DE, with a closer correlation to CMR-derived volumes.50,96 For these reasons, the ASE and EACVI recently recommended 3DE, rather than 2DE, for the routine assessment of LV volumes and EF.100 Two recent studies have addressed normal ranges of 3D measurements but identified somewhat different normal values, emphasizing racial, gender, and age differences.49,101 A recent meta-analysis of validation studies comparing 3DE and CMR demonstrated that considerable variability still exists in the measurement of LV volumes (±34 mL for EDV, ±30 mL for ESV, and ±12% for EF), although it is less than that observed between 2DE and CMR.97 Moreover, both 2DE- and 3DE-derived volumes are less accurate in dilated LVs.102 Several sources of 3D volume acquisition and measurement error are discussed in the recent ASE/EACVI guidelines, including difficulty in imaging the anterior and lateral walls because of interference from ribs, low line density (and therefore lower spatial resolution—which may be partly addressed with the use of LV opacification), low temporal resolution (which may be addressed by using multiple subvolumes—but at the risk of stitching artefacts), and time-consuming off-line analysis.100 Recently, a fully automated endocardial contouring system combined with real-time full-volume 3DE has been described as providing accurate and reproducible volumes.103

### Midwall function

#### Rationale
LV systolic function is commonly assessed through the use of EF and FS. However, because these measurements are performed at the endocardial surface, their appropriateness has been questioned in patients with LV hypertrophy. The inner layer of the LV has been shown to move inward further than the outer layer, a difference markedly increased in hypertrophic walls due to the ‘cross-fibre shortening’ phenomenon which, in hypertrophic LVs, achieves normal systolic wall thickening despite reduced shortening of individual myocardial segments.104–106 Hence, LVEF and FS often lead to overestimation of LV systolic performance yielding normal or even supranormal results not matching the individual’s clinical situation and prognosis, since they take into account geometric changes that do not accurately reflect the actual contractile function of the myocardium.104,107–109 The greatest proportion of ventricular myocardial fibres is located in the myocardial midwall, the region responsible for circumferential left ventricular contraction and where cross-fibre shortening is less significant.110–112 Consequently, indices representing LV midwall mechanics have received increasing attention lately, as they have shown to better reflect myocardial contractile function in several clinical scenarios, through better prediction of cardiovascular outcomes than indices based on endocardial measurements and better correlation with patients’ clinical status.120–123

### Limitations
Some of the limitations of midwall function assessment include the fact that FS_{mw} is based on a limited region of the LV, which could hinder its application to patients with variable LV geometries.124 Another potential limitation is the need for manual tracking, which introduces the problems of time-consuming analysis and potential interobserver variability. However, new indices and calculations partly overcome these limitations through the analysis of 2D and 3D midwall mechanics, introducing the concepts of 2D and 3D midwall EF,125,126 Finally, advanced echocardiographic techniques are modifying the understanding of the hypertensive heart. Disturbances of longitudinal strain of the endocardial layer precedes the alteration of circumferential strain, which is attributed to the midwall layer.127 This is important because LV longitudinal dysfunction plays a role in mediating the effect of LV geometry on LV diastolic impairment.116

### Tissue Doppler assessment of systolic function
Tissue Doppler was the first widely available myocardial imaging technique, and is credited with improving the feasibility of longitudinal ventricular function measurement. Several studies have shown tissue Doppler—using either pulsed-wave or colour mapping—to be a reliable tool for the assessment of LV systolic function. This method has been validated against other methods for the assessment of myocardial systolic performance and regional coronary blood flow, as well as with histological findings.128–131 Its high temporal resolution enables accurate determination of myocardial velocity and acceleration even when overall image quality is deficient and endocardial delineation is poor.132,133 Technical considerations related to tissue Doppler have been considered in depth in an ASE/EAE consensus statement, and will not be replicated here.134 In hypertensive heart disease, the tissue relaxation velocity (e’) is
Assessment of myocardial function by strain

Strain, strain-rate, and twist imaging (deformation imaging) are relatively recent non-invasive methods for the assessment of regional and global myocardial function, allowing discrimination between active and passive myocardial tissue motion. Assessment of strain and twist is extracted from images using the commercially available software, providing sensitive echocardiographic measures to detect early subclinical evidence of ventricular dysfunction. This information can be gathered using tissue Doppler echocardiography or speckle tracking and has been described in detail in a recent ASE/EACVI Consensus Statement. The measurement of strain has been well validated with sonomicrometry, three-dimensional tagged CMR, and cyclically compressed tissue-mimicking gelatin phantom. Among the different deformation (strain) components, longitudinal strain has gained an important value in this context. Longitudinal strain corresponds to the function of the endocardial layer of myocardium, where longitudinal fibres are subjected to the negative impact of early development of fibrosis in hypertensive heart disease. However, strain is highly sensitive to increased afterload, and the relative degree of impairment of strain that is due to LV dysfunction vs. that is due to hypertension may be difficult to tease apart. Reported normal values of global longitudinal strain vary from −15.9 to −22.1% (mean, −19.7%; 95% CI −20.4 to −18.9%). This technique has been used to differentiate between different causes of increased wall thickness. In addition to the degree of reduction of strain, the pattern of strain reduction is also important. For example, amyloidosis is characterized by a particular pattern of apical sparing not seen in other causes of hypertrophy, and hypertrophic cardiomyopathy is associated with deformation disturbances at the site of hypertrophy with less abnormal deformation elsewhere. The morphology of the longitudinal strain signal may also be important in recognizing myocardial scarring. A characteristic double peak in the strain-rate signal has been identified in patients with scar tissue associated with hypertrophy in hypertrophic cardiomyopathy, Fabry’s disease, and aortic stenosis. This phenomenon presumably reflects a degree of post-systolic shortening in the presence of fibrosis. Thus, although the functional markers are non-specific for the diagnosis of hypertensive heart disease, they may demonstrate specific patterns and degrees of disturbance that distinguish between hypertensive hypertrophy and other aetiologies, as well as recognizing the contribution of fibrosis. Longitudinal strain can be even used to differentiate hypertensive heart disease from functional myocardial changes in the athlete’s heart.

Finally, CMR may be used for quantifying myocardial function, using techniques that measure myocardial deformation. It is not clear that these are superior to the echo techniques, as they are obtained at lower temporal resolution. This may be particularly pertinent for the identification of post-systolic shortening or disturbances of diastolic function.

Prognostic significance of LV function in hypertension

Chamber function

The prognostic significance of LV function is well established. It is known that heart failure is a common consequence of hypertension and in the majority of patients is related to impaired LV systolic function, which accounts for about half of heart failure cases. However, hypertension is not necessarily associated with a reduced systolic function—this may be increased in the initial stages. EF, a global measure of LV chamber function, is used to distinguish systolic (EF <50%) from diastolic HF (EF ≥50%), and is a reliable method for predicting primary cardiac events and cardiac mortality in individuals. Endocardial FS is a good measure of LV global systolic function; however, its use in the setting of hypertension is discouraged, especially in the presence of LV hypertrophy. As discussed above, both EF and FS are constrained...
because they measure endocardial function, whereas the true parameter of interest is midwall function. In addition, the limited field of view with M-mode leads to an under-appreciation of regional wall motion. Wall motion abnormalities can identify adults without known cardiovascular disease (CVD) who are at 2.4- to 3.4-fold higher risk of CVD morbidity and mortality.\textsuperscript{151}

In contrast, 2D strain has been shown to be abnormal in hypertensive patients with normal EF.\textsuperscript{127} as well as in prehypertension.\textsuperscript{152} Although EF is accepted as a prognostic marker, its prognostic value in the range that is close to normal is limited. Strain does not seem to share this limitation,\textsuperscript{46} and this may be of value in discerning the progression from hypertensive heart disease to heart failure.

### LV midwall function in hypertension

The calculation of LV midwall shortening (FS\textsubscript{MW}) has been discussed above. Depressed FS is associated with increased LV RWT and LVM, and FS may be impaired in hypertensive patients with normal LVEF.\textsuperscript{153} FS predicts adverse outcomes,\textsuperscript{108} but there are limited data about the relative ability of FS to predict cardiovascular events independent of known established risk factors (LVM and BP).\textsuperscript{122} Indeed, some authorities question the incremental information provided by the assessment of LV systolic function to LVM in hypertensive heart disease.\textsuperscript{154} Nor has it been shown that improved treatment of LV chamber systolic function (in those with a normal EF and depressed LV FS\textsubscript{MW}) is associated with lower CVD morbidity and mortality, independent of change in BP and LVM in treated hypertensive patients.\textsuperscript{122}

### Prognostic significance of mitral inflow patterns

The main prognostic importance of Doppler-derived LV filling is in patients with systolic HF, where mitral inflow measurements correlate with LV filling pressure, functional classes, and prognosis.\textsuperscript{158} In hypertension, normal in-treatment transmitial flow pattern indicates a low risk for heart failure (HR 0.22 [95% CI 0.05–0.98, P = 0.048], independent of blood pressure.\textsuperscript{159} However, the intermediate range of E/A ratio (from 0.6 to 1.5) do not stratify prognosis in hypertensive subjects,\textsuperscript{17} probably because normal and pseudonormal patterns are combined. Although antihypertensive treatment in patients with LVH results in improvement of mitral inflow patterns, this was not associated with reduced cardiovascular morbidity and mortality.\textsuperscript{159}

### Tissue Doppler assessment of myocardial diastolic function

### Acquisition and measurements

Guidance on the technical requirements for tissue Doppler acquisition has been provided regarding sample volume location, angulation, and respiratory phase.\textsuperscript{160} In hypertensive heart disease, early diastolic tissue velocity (e') is reduced by reduction in LV relaxation. However, it is also influenced by preload, systolic function, and LV minimal pressure. The other basic measured parameter is late (atrial) diastolic velocity (a', influenced by LA function and LV end diastolic pressure), E/e' has been used as a measure of LA driving pressure or LV filling pressure.\textsuperscript{161} However, there are a number of situations where e' and E/e' may be misleading,\textsuperscript{162} including reduced septal e' velocity due to inferior infarction or annular calcification, and increased transmitial E velocity due to mitral regurgitation. Averaging septal and lateral e' may reduce some of this variability, but does not address all of the limitations of the parameter.

### Normal values

Normal values are reported in Table 7. Diastolic filling patterns are classified by the combined quantitative analysis of E/A ratio, DT, tissue Doppler, and LA volume in particular.\textsuperscript{156} With increasing age, impairment of LV relaxation leads to low E velocity, high A velocity, and decreased E/A ratio with prolongation of DT.\textsuperscript{157} Delayed relaxation occurs in uncomplicated systemic arterial hypertension. Because of the load dependence of these measurements, the pseudonormal filling pattern cannot be recognized on the basis of simple evaluation of mitral inflow pattern, but needs additional assessment during a Valsalva manoeuvre (low reliability) or the additional assessment of pulmonary venous flow (intermediate reliability) or pulsed Tissue Doppler-derived e' velocity of the mitral annulus (highest reliability). The detection of LA enlargement is a marker of longstanding increase of LA pressure in hypertensive heart disease.
Prognostic significance of tissue Doppler parameters

Annular tissue velocities are strong predictors of outcome in a variety of settings. In a 2-year follow-up study of >500 patients, 35% of whom had hypertension, Wang et al. showed that a pulsed-wave e’ of <3 cm/s was associated with a 5.3-fold increment of hazard. As these data were gathered from colour-coded tissue Doppler, they represent unusually low values for e’ velocity, analogous to pulsed-wave signals in the range of <5 cm/s. Similar findings have been described using an e’ of <3.5 cm/s in hypertension and LV hypertrophy. It has to be acknowledged, however, that velocities <5 cm/s are quite extreme, and less usual in hypertensive heart disease than hypertrophic cardiomyopathy or infiltration.

Likewise, E/e’ has prognostic implications, with E/e’ ≥ 15 having been shown to add an independent prognostic value to B-type natriuretic peptide and EF. Although studies have been more focused on post-MI and heart failure than hypertension, Sharp et al. recently demonstrated the prognostic value of E/e’ ratio in uncomplicated hypertensive patients, independent of LVM. On these grounds, the 2013 ESC/ESH guidelines on arterial hypertension promote the use of E/e’ in the detection of cardiac target organ damage in hypertensive heart disease.

Recommendations

All echocardiography reports in patients with hypertension should include specific comments about diastolic function grade, left atrial volume, and about normal vs. elevated LV filling pressure (usually based on E/e’).

Cardiac impact of hypertension treatment

LV hypertrophy regression

LV hypertrophy represents an important end-organ consequence of hypertension. Population-based studies using echocardiography have demonstrated hypertrophy to be closely linked with adverse events, including stroke, renal impairment, left ventricular dysfunction, atrial and ventricular arrhythmias, and sudden cardiac arrhythmia or premature death. The eventual development of complications from LVH represents long-term effects that are too final to guide clinical therapy, and too slow as a research outcome. Therefore, LVH has been proposed as a surrogate marker of outcome. LVH has been shown to be reversed or prevented by a variety of haemodynamic, non-haemodynamic, and pharmacological factors.

Nonetheless, the use of repeat imaging to document changes in LVM has been difficult to incorporate into standard practice for at least two reasons. The first relates to the inherent variability of LVM measurements with echocardiography. While reductions in the ventricular mass have been associated with improved outcome across populations, in studies which identify regression of hypertrophy on an individual basis, large populations are required to overcome the variability of these measurements. Thus, while the association between LVH regression and improved outcome has now been recognized in a number of studies, because of the test–retest limitations of echocardiography, CMR may be more accurate to demonstrate this effect. This role of echocardiography may be improved by the enhancement and clinical use of 3DE, which has been validated against CMR.

The second limitation is that hypertrophy occurs in 36–41% of hypertensive subjects, but hypertension is not the only cause of this problem. Hypertrophy may be influenced by obesity, diabetes, the metabolic syndrome, and renal impairment, among other aetiologies. Progression of the condition may lead to ischaemia, both due to concurrent coronary artery disease as well as failure of vascular proliferation to match myocardial proliferation, vascular compression, and the effect of raised LV pressure on subendocardial flow.

Change in LV geometry

Changes in LV geometry have been associated with improved blood pressure control, reflecting the impact of afterload on LV remodelling. Again, however, the variability of 2DE has been a limitation in understanding the association of reverse remodelling with improved survival, using conventional techniques. Recent evidence has indicated that use of CMR (or potentially 3DE) provides a means of measuring sphericity on a serial basis, and therefore documents remodelling changes in response to blood pressure control.

Change in systolic function

LV systolic function, measured by EF, is normally preserved until late in the course of hypertensive heart disease. Indeed, although EF is associated with outcome in patients with moderate LV impairment,
the association of mild or borderline impairment with adverse outcome has been more difficult to show. Likewise, volumetric and EF changes in heart failure have been associated with improvements in outcome, but this information is difficult to apply to hypertensive heart disease in which EF is either preserved or borderline reduced.

Change in diastolic function

Diastolic dysfunction, particularly in the later stages of hypertensive heart disease, is associated with prognosis. However, most patients with hypertensive heart disease have grade I diastolic dysfunction, and changes in this finding are intrinsically ambiguous. When the E/A ratio is <1 and moves towards unity, this may occur because of recovery of function and improvement in LV suction, or it may occur because of raised filling pressures and transition of grade I to grade II disease. Documentation of changes in diastolic function is difficult to interpret in any patient, and no less in those with hypertensive heart disease. In a randomized study of angiotensin receptor blockade, no significant change in e’ was witnessed between valsartan and the control group. Nonetheless, other studies have shown that improvements in LV geometry after treatment in hypertensive patients with ECG evidence of LV hypertrophy have been associated with parallel improvements in Doppler-derived indices of diastolic function.

Recommendations

While echocardiography has been key in demonstrating the beneficial effects of hypertension treatment in large cohort studies, routine reassessment of echocardiograms to examine treatment response in hypertensive subjects is not recommended, due to the limited reproducibility of measurements on an individual patient basis. Follow-up echocardiograms may be of value to assess changes in symptom status.

Echocardiography in clinical management of hypertension

Stratification of risk in hypertension

The value of transthoracic echocardiography is recognized in the 2013 ESC/ESH guidelines, where it is listed as a class II indication (level of evidence B) for cardiovascular risk assessment in asymptomatic adults with hypertension. Transthoracic echocardiography received a high appropriate use criteria score of 8 (scale 1–9) for the initial evaluation of suspected hypertensive heart disease. In this document, LV hypertrophy, LV diastolic dysfunction, and LA enlargement are described as specific signs of hypertensive heart disease. LV hypertrophy is recognized as evidence of target organ damage in hypertension by the Joint National Committee for the prevention, detection, and evaluation of high blood pressure (JNC 7) of the National High Blood Pressure Education Program (National Heart Lung and Blood Institute).

In patients with hypertension, the type of LV remodelling (concentric remodelling, eccentric hypertrophy, and concentric hypertrophy) is predictive of the incidence of CV events. In particular, the presence of LVH on echocardiography identifies hypertensive heart disease with a higher sensitivity and specificity compared with electrocardiography. Several population cohort studies have shown that LVH is predictive of cardiovascular and all-cause mortality, independent of blood pressure, and across all racial groups that have been studied. In the predominantly white population of the Framingham Study, for every 50 g/m² higher left ventricular mass index, there was a relative risk of death of 1.73 (95% CI 1.19–2.52), independent of blood pressure level. In African-Americans enrolled in the ARIC study, LVH was associated with an increased risk of cardiovascular events (HR of 1.88 in men and 1.92 in women). Similarly, for Native Americans enrolled in the Strong Heart Study, echocardiographic LVH also had additive discriminatory power over ECG LVH; the prevalence of LVH on echocardiography was 9.5% and was associated with a seven-fold increase in cardiovascular mortality and a four-fold increase in all-cause mortality. Hispanic Americans showed a similar association of LVH and CVD mortality.

International studies have also confirmed a similar risk for CVD in hypertensive patients with LVH. Concentric LVH on echocardiography identifies a high risk phenotype with abnormal flow-mediated dilatation and decreased myocardial flow reserve.

In symptomatic adults with hypertension, the echocardiogram provides additional assessments for systolic and diastolic dysfunction, as well as evaluation of wall motion abnormalities to detect underlying coronary artery disease. The use of echocardiography during treadmill or pharmacological testing is indicated in hypertensive patients with symptoms suggesting CHD and/or to estimate prognosis in patients with known concomitant coronary artery disease as well as those with known or suspected valvular heart disease. Patients with LVH, as well as related problems (abnormal resting ECG, left bundle-branch block, electronically paced rhythm, and digoxin therapy), also warrant pharmacological stress echocardiography.

Investigation of chest pain symptoms

Chest pain in patients with hypertension may signify concurrent coronary artery disease or may simply reflect subendocardial ischaemia due to LV hypertrophy and increased afterload. The diagnosis of coronary artery disease has particular challenges in this setting, because ‘false-positive’ results may occur when subendocardial ischaemia causes abnormal stress ECG or myocardial perfusion scan in the absence of flow-limiting epicardial coronary disease. A normal stress electrocardiogram, performed to a high workload, has a high negative predictive value, but an abnormal or ambiguous test warrants further evaluation. There is some evidence in favour of preferential use of stress echocardiography for this purpose, because stress-induced wall motion abnormalities are highly specific for coronary artery disease, while perfusion defects in hypertensive patients may arise from abnormal myocardial flow reserve not due to epicardial coronary disease. The lack of specificity of the coronary flow signal for epicardial coronary artery disease is also a problem when stress echocardiography is combined with the assessment of coronary flow reserve in hypertensive patients. Finally, although hypertensive patients are at increased risk of coronary artery disease, screening for coronary disease is not recommended in asymptomatic patients because of the risk of false-positive results and uncertain management responses.
Effects of antihypertensive agents on LVM and other echocardiographic surrogate endpoints (e.g. LA size and diastolic function) have been extensively studied. Several large studies sponsored by the National Institutes of Health and the US Veterans Administration Cooperative Studies program have evaluated the effects of antihypertensive monotherapy. In general, it appears likely that there are differences between the efficacy of antihypertensive drugs and their effects on LVH. LVH regression does not adversely affect cardiac function and may be associated with improvements in diastolic function. However, although the finding of increased LVM on echocardiography could potentially guide selection of initial or intensity of therapy in hypertensive patients, JNC 7 recommendations do not risk stratify patients for treatment on the basis of target organ damage. Current guidelines recommend the use of combination treatment to get blood pressure to goal, thus blood pressure remains the primary target of therapy.

A part of the problem with getting a more central role for echocardiography to guide therapy is that despite the adverse prognosis associated with LVH in hypertension, there are inconsistent data from numerous studies that have evaluated the comparative efficacy of specific antihypertensive agents in LVH regression, as well as survival benefits associated with LVH regression. In a meta-analysis of 39 trials of antihypertensive therapy, angiotensin-converting enzyme inhibitors were the most effective agents, leading to a 13.3% reduction in LVM compared with 9.3% for calcium channel blockers, 6.8% for diuretics, and 5.5% for beta-blockers. However, in a comparison of enalapril and long-acting nifedipine in patients with essential hypertension, the PRESERVE (Prospective Randomized Enalapril Study Evaluating Regression of Ventricular Enlargement) trial, systolic and diastolic pressures, as well as LV mass were reduced to a similar degree with both agents. On the other hand, the LIFE (Losartan Intervention For Endpoint Reduction in Hypertension) trial echocardiographic sub-study demonstrated superior LVM reduction (21.7 g/m²) in patients treated with the angiotensin receptor blocker losartan compared with those treated with the beta-blocker atenolol (17.7 g/m²). Finally, despite a 20% incidence of LVH regression with placebo, diuretic therapy with chlorothalidone and hydrochlorothiazide, respectively, demonstrated greater LVH regression over alternative agents in both the TOMHS (Treatment of Mild Hypertension Study) and Department of Veterans Affairs Cooperative Study Group on Antihypertensive Agents. Similarly, left atrial size (itself a predictor of adverse outcomes) was reduced with hydrochlorothiazide.

In the recently defined category of pre-hypertension (systolic blood pressure 130–140 mmHg and/or diastolic blood pressure 80–90 mmHg), JNC 7 recommends intensive lifestyle modification in such patients. Clinicians may obtain echocardiography to evaluate the presence of LV hypertrophy in such patients with pre-hypertension, particularly where there is a strong family history of hypertension and cardiovascular complications including stroke, heart failure, or dialysis. The community practice consensus is that the presence of LVH in such patients should lead to more aggressive approaches to lifestyle modification. There is increasing recognition that data on target organ involvement, including echocardiographic LVH, may be important for young adults whose lifetime risk for hypertension is currently underestimated by most risk stratification models. However, no studies have examined whether a patient’s knowledge of echocardiography demonstrating LVH will improve adherence to lifestyle modifications or pharmacological treatment of hypertension.

According to the National Health Service and National Institute for Health and Clinical Excellence (NICE) recent guideline update on the clinical management of primary hypertension in adults, there is uncertainty about how to assess the impact of blood pressure treatment in people aged <40 years with grade 1 hypertension and no overt target organ damage or CVD. In particular, it is not known whether those with untreated hypertension are more likely to develop target organ damage and, if so, whether such damage is reversible. The writers of the NICE guideline further observe that target organ damage as surrogate or intermediate disease marker for CVD or hypertensive heart disease is the only indicator that is likely to be feasible in younger people because traditional clinical outcomes are unlikely to occur in sufficient numbers over the timeline of a typical clinical trial.

The decision to intensify treatment of hypertension is currently guided by monitoring of clinic as well as home blood pressures. In patients who have hypertensive heart disease with LVH and normal systolic function, the value of periodic echocardiographic follow-up is not established; the Appropriate Use Task Force gave a score of 4 (may be appropriate) based on insufficient data for a stronger recommendation, regarding the re-evaluation of known hypertensive heart disease without a change in clinical status or cardiac examination. However, echocardiography may be helpful in several scenarios. Patients with hypertensive heart disease who become symptomatic require follow-up echocardiography to evaluate systolic and diastolic function. Dissociation between blood pressure measurements and LV hypertrophy is an indication for further testing. The detection of high blood pressure without hypertrophy should lead to consideration of overestimation of the severity of hypertension, including ambulatory blood pressure monitoring or measurement of central aortic pressure. When there is apparent LV hypertrophy in the setting of apparent blood pressure control, more detailed blood pressure evaluation (e.g. for masked hypertension) or identifying other causes of wall thickening such as infiltrative diseases should be considered.

There is no current indication for the use of echocardiography to routinely monitor antihypertensive therapy, except as indicated and described in the section above for symptomatic patients or for patients with poor control of blood pressure. A recent intersocietal consensus document on the appropriate use of echocardiography in clinical practice characterized its routine use for patients with hypertension without symptoms or signs of heart disease as ‘rarely appropriate’ with a value score of 3 out of 10.
Relevance of hypertension to echocardiographic interpretation

Afterload is an important determinant of the assessment of cardiac function from ejection phase indices. Consequently, hypertension may have an important effect on the assessment of LV function in a variety of conditions. For example, an increment of blood pressure between visits may lead to an apparent deterioration of LV function when serial echocardiograms are being performed during chemotherapy or in the evaluation of valvular heart disease. In the assessment of aortic stenosis, arterial hypertension and the stenotic valve behave like serial resistors, and their combined impedance may explain symptom status.194

Likewise, in stress echocardiography, hypertension—especially a hypertensive response to stress—may provoke wall motion abnormalities or global LV dysfunction in the absence of coronary disease.197 However, the impact of hypertensive LVH is probably less than in perfusion scintigraphy, where abnormal coronary flow reserve may produce false-positive perfusion abnormalities in the context of normal wall motion.198

Recommendations

At present, decisions regarding the initiation, intensification, or monitoring of response to antihypertensive therapies are made based on clinical parameters.

Given the progressive nature of hypertensive cardiomyopathy, periodic evaluation of cardiac function and morphology by echocardiography may be warranted, especially if symptoms change.

Recommendations for clinical laboratories

The value of echocardiography as a research tool in hypertension is uncontested. In a relatively short time, it has defined the cardiac structural and functional effects of hypertension, determined the prevalence of LVH and LV remodelling, determined the cardiac effects of antihypertensive therapy, and in epidemiological studies, provided fundamental insights into the relationships between blood pressure, genetic susceptibility, and LV mass. However, although it has been suggested that treatment choices in individual patients should be guided by echocardiographic findings, the value of echocardiography in the clinical management of hypertension is unproven.

The benefits of echocardiography will depend on its value in affecting treatment decisions, and in early identification and intervention in patients at risk who would not otherwise be treated. Moreover, demonstration of a value requires that the impact of echocardiography on clinical decisions is accompanied by improvement in patient outcome. Importantly, any consideration of the utility of echocardiography is contingent upon its reliability for the assessment of target measures such as LVM. However, little information is available on the impact of echocardiographic data on physician behaviour, or on patient outcomes in hypertension.

Previously, World Health Organization-International Society of Hypertension (WHO-ISH) suggested that drug treatment could be withheld in hypertensive individuals with low cardiovascular risk based on non-echocardiographic criteria. However, echocardiographic findings in such individuals increases risk classification in 29% of such cases,199 suggesting a role for echocardiography in risk profiling. However, recommendations for drug therapy at lower blood pressure levels may have made this application of echocardiography moot.

Another important limitation on the wider use of echocardiography is cost, both relative to benefit, and in competition for economic resources. In the USA alone, approximately 76.4 million adults have hypertension.200 At even the arguably modest current Medicare/Medicaid reimbursement for echo of $238, one echo per patient with hypertension would cost $181.2 billion. Justification for this expenditure as an additional billable item would be difficult to provide. However, the development of hand-held ultrasound would allow suitably trained practitioners to obtain LV wall thickness and dimension information as part of the office visit. The effectiveness of this strategy remains unproven, especially in the light of training requirements, concern about interobserver variability, lack of standard quality assurance standards, and the increase in time for an office visit.

Given the above considerations, it has been recommended that echocardiography be reserved for those individuals with hypertension in whom hypertensive cardiac disease or cardiac disease in association with hypertension comorbidities is suspected. In such cases, a complete 2D and Doppler study should be performed, and the study not limited to evaluation of LVM/LVH. While calculation of LVM can readily be performed utilizing standard methods,46 variability can be quite large, and current evidence does not support using LVM measurement to either initiate or modify hypertension treatment.

Recommendations for research studies and clinical trials

Table 8 lists some potential areas where echocardiography (or other imaging) may help to guide management decisions in hypertension. The role of imaging in these settings is unproven and warrants further study.

Acquisition and interpretation of echocardiograms for research purposes in hypertension poses some special challenges. Even for clinically experienced sonographers, there is a significant learning curve present in recording technically adequate echocardiographic studies for the assessment of LVM, particularly in older subjects. In a Framingham analysis of M-mode echocardiograms performed in over 6000 subjects aged 17–90, the ability to record acceptable quality echocardiograms in subjects older than 60 years rose from a minimum of 28% during the first 5 months of the study to a maximum of 74–81% during studies 2 years later. Hence, echocardiography ‘drop-outs’ may not be randomly distributed, leading to the possibility of bias in data interpretation. Two-dimensional echocardiographic measurements were even more problematic than 2D-guided M-mode.

In previous large echocardiography trials, major differences in echo quality have existed between field centres. For example, in a 15 centre ventriculo-arterial trial of antihypertensive monotherapy,192 the percent of readable echocardiograms for LVM varied...
from ~30 to 85%. This was not due to differences between centres in the proportion of easy or difficult patients. While the use of suboptimal equipment in some cases contributed to poor studies, the intercentre differences were mostly because of variation in technical performance. Importantly, extensive previous clinical experience in echocardiography was no guarantee of high-quality echocardiograms for research purposes.

There is potential for differences in image acquisition styles that may exist between field centres in epidemiology studies, potential effects of instrumentation, continuing improvement in quality images obtained with newer generations of echo machines, and temporal intrareader drift in echo measurements and interpretations (e.g. LV walls may be read as thicker, or thinner at the beginning than the end of the clinical trial or observational study). All of these may produce not just large random variability in measurements and qualitative assessments, but substantial biases. For example, a temporal drift where readers might tend to read smaller wall thicknesses after months to years of experience with the study and patients receiving several treatments in the absence of a placebo control (common if not ubiquitous in clinical hypertension trials) may lead to the mistaken conclusion that both treatments are associated with decreased LVM and decreases in the proportion of individuals with LVM.

Several principles learned from clinical trials are applicable for echocardiography.201 The acquisition of reproducible, correctly oriented images requires sonographer training. Monitoring of study quality is important. The inclusion of ‘control’ subjects is a protection against apparent changes due only to regression to the mean. The use of sample echocardiograms is a means of ensuring that all team members are applying the same methodology, and to prevent ‘drift’ over time.

**Recommendations for echocardiography in hypertension clinical trials**

Given the large confidence intervals that may exist for measurement of LVM, it could be argued that treatment trials should recruit participants with markedly increased LVM. However, selection of participants with values for LVM values substantially above (or below) the population mean can result in subsequent tests that reflect regression to the mean. Therefore, higher than ‘true’ values for LVM on an initial determination will tend to decrease on subsequent measurement. It is recommended that if possible, partition values for LVM not be used as requirements for entry into the study. If such values are used, then batch reading at completion of the study (with continuous monitoring of studies for acquisition quality) should be done. This may not be practical in long-term studies.

In creating categorical variables (e.g. LVH, LA enlargement, and abnormal annular tissue velocity), it is advisable when possible to use comparative control subjects from the same study to generate partition reference values. This is often possible in observational or epidemiologic studies, where participants without clinically prevalent disease (or better still—free of subclinical disease as well) can be utilized to derive partition values for continuous variables. Where those values are affected by age, body weight, height, gender, etc.—reference values can be derived from regression models used to derive a predicted value (with confidence limits) and express abnormality of a parameter by determining its ratio to this predicted value. However, this may not be possible in many clinical trials.

In large multicentre observational studies and clinical trials where all studies are read by a single core laboratory, the volume of studies can quickly become overwhelming. Special considerations exist re: management of workflow, but also vetting of site sonographers, participation in trial design, statistical power estimations, provision of ongoing quality assurance and improvement, data transmission to the statistical core, and issues regarding participant and investigator clinical alerts for abnormal findings. Specific considerations regarding core laboratory best practices have been described in a previous ASE EACVI expert consensus statement.202,203

**Conflict of interest:** This report is made available by EACVI and ASE as a courtesy reference source for members. This report contains recommendations only and should not be used as the sole basis to make medical practice decisions or for disciplinary action.
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