Transthoracic echocardiography of hypertrophic cardiomyopathy in adults: a practical guideline from the British Society of Echocardiography.

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Article
Title: Transthoracic Echocardiographic of Hypertrophic Cardiomyopathy in Adults: A Practical Guideline from the British Society of Echocardiography

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Abstract

Hypertrophic cardiomyopathy (HCM) is common, inherited and characterised by unexplained thickening of the myocardium. The British Society of Echocardiography (BSE) has recently published a minimum dataset for transthoracic echocardiography detailing the core views needed for a standard echocardiogram. For patients with confirmed or suspected HCM, additional views and measurements are necessary. This guideline, therefore, supplements the minimum dataset and describes a tailored, stepwise approach to the echocardiographic examination, and echocardiography’s position in the diagnostic pathway, before advising on the imaging of disease complications and invasive treatments.
Intent behind update

These guidelines on hypertrophic cardiomyopathy (HCM) represent a five-year update. They contain a description of pertinent disease features and the critical echo parameters needed to evaluate the condition, alongside a recommended protocol. A specific HCM minimum data set, for use as an aide memoir when reporting, is provided.

The guideline also proposes an echocardiographic approach to diagnosis as well as information on the use of echo measurements for sudden death risk stratification. This guideline aims to enhance baseline knowledge and to allow echocardiographers to develop a systematic approach to the image acquisition and echocardiographic reporting of patients with proven or suspected HCM. The guideline-writing committee anticipates that readers armed with this knowledge will approach these examinations with confidence, extract as much information about each patient’s condition as possible and produce unambiguous, standardised reports. These actions will enhance clinical care by limiting the number of patients who are either under or over-diagnosed and highlight the sub-cohorts of patients who need additional investigations and treatments. The guidelines end with short sections covering the use of echo guidance for transseptal alcohol ablation and surgical myectomy as well as strain, stress and three-dimensional echocardiography in patients with HCM.

Hypertrophic Cardiomyopathy

HCM in adults is defined ‘by a wall thickness ≥15 mm in one or more left ventricular (LV) myocardial segments that is not explained solely by loading conditions’ (1), for example, hypertension. In a smaller number of cases, described in the next section, HCM may be associated with an abnormal wall thickness which measures less than 15mm. This dimension-based diagnosis covers a diverse group of diseases, both inherited and acquired, which differ in their pathophysiology and management.

Due to the challenges in certain aspects of diagnosis and management of this patient group, referral to specialist centres focused on inherited cardiac conditions and cardiomyopathies is recommended.
for patients with suspected or confirmed disease(1). Where possible, echocardiographers should obtain dedicated training in the scanning and interpretation of this patient group.

The condition affects between 0.2%(2) and 1.4% of individuals(3). Disease complications are reasonably common; in a multi-centre longitudinal study of patients with HCM, atrial fibrillation occurred in 20%, sudden cardiac death or resuscitated cardiac arrest in 4%(4) and left ventricular systolic dysfunction (ejection fraction <50%) in 8%(5).

The pattern of inheritance is autosomal dominant. A clinically meaningful gene change (found predominantly in MYBPC3 and MYH7) occurs in a fifth of patients where the family history is negative, and a half where it is positive(6). Finding a disease-causing gene change allows testing of family members using pre-symptomatic screening.

**Echocardiography’s Position in the Diagnostic Pathway – Wall thickness**

Accurate measurement of wall thickness is fundamental to decision-making. Because of this, the echocardiographic examination is a key component of the diagnosis pathway. Ancillary features such as left ventricular outflow tract obstruction (LVOTO) do not contribute.

Measurements should be made in short-axis views orthogonal to the circumference of the endocardium and epicardium, wherever maximal wall thickness occurs. Elements attached to but not incorporated in the septum should be excluded (Figure 1), as this will overestimate wall thickness and run the risk of misdiagnosis of HCM. The report should state if the study failed to visualise any part of the LV (often the basal anterior and anterolateral walls) and recommend alternative imaging modalities, specifically cardiovascular magnetic resonance.

The dimensional threshold for HCM depends on the location of hypertrophy as well as the clinical context. In apical HCM, where normal tapering of both cavity and epicardium is lost, the apical wall thickness may be less than 15 mm(7). One criterion defines apical HCM when the ratio between apical and basal wall thickness exceeds 1.3: 1(8). Visualisation of this area can be difficult and may require
the use of myocardial contrast (Figure 2). By ensuring the apical four, two and three-chamber view section the apex, the echocardiographer will avoid giving the impression of apical hypertrophy by foreshortening views. Apical wall thickness should be measured in the short-axis view, ensuring the cut is not oblique to the long axes of the LV.

In first-degree relatives - who have a 50% risk of inheriting the causative gene - the wall thickness threshold for diagnosis of HCM is ≥13 mm (1). The yield of positive screening examinations in first degree relatives vary based on the population tested; in one report, 5% of first-degree relative children were diagnosed with HCM(9), rising to 30 % of a mainly adult cohort in another, where many had a disease-causing gene(10). A feature of HCM is age-related penetrance, where the percentage of individuals carrying the disease-causing gene who express the condition increases with age. The yield of clinical screening is higher in families where the disease onset has been in childhood(9,10).

HCM featuring the so-called dilated-hypokinetic or ‘burnt-out’ phase (5), or due to specific gene mutations(11–13), can be associated with only mild increases in wall thickness.

Grey Cases

Ethnicity, hypertension, renal disease, significant aortic stenosis, increased body mass index and athletic remodelling all influence left ventricular hypertrophy. Increased LV wall thickness secondary to these processes may fall into a ‘grey zone’, overlapping with the degree of LV hypertrophy (LVH) seen in HCM (Figure 3). For example, a wall thickness of 15 -20 mm can occur in hypertensive heart disease in individuals of African/Afro-Caribbean ethnicity, whereas the same degree of hypertrophy in a Caucasian hypertensive patient would suggest HCM(1). LVH in hypertensive heart disease and athletic remodelling tends to be uniform and symmetrical.

In athletes, gender, in addition to ethnicity, is relevant. Wall thickness is lower in female athletes than their male counterparts and does not exceed 13 mm in Caucasian athletes or 15 mm in athletes of African/Afro-Caribbean ethnicity(14). In a study of athletes with HCM compared with athletes without
HCM(15), the diagnosis was definite in most individuals as maximal wall thickness was >16 mm, and often the LVH was distributed non-uniformly. The scenario in which there was uncertainty – where LVH was 13-16 mm and concentric (defined by a relative wall thickness of >0.42 (see BSE guidelines on normal reference intervals for cardiac dimensions and function for more information (16)) – cropped up in only 14% of athletes with HCM. Measures like left ventricular cavity size (previously reported to be a useful differentiator between HCM and athletic heart; being larger in the latter (17)) showed modest performance in picking out athletes with HCM. Additional tests were required to distinguish these individuals from athletes with physiological remodelling.

Recommended Language in Echocardiography Report

Echocardiography’s pivotal role means that a study’s interpretation can strongly influence the clinical team’s diagnostic decision. Because of this, we encourage the use of standardised language when reporting. In instances where there is uncertainty, ‘raises the possibility of HCM’ is recommended. In individuals undergoing screening, where there is no evidence of left ventricular hypertrophy, the conclusion should contain the following suggested phrase: ‘wall thickness is normal’. The proposed language provides an objective statement about the echocardiogram findings, rather than a definitive clinical assertion. Hence ‘wall thickness is normal’ is not the same as ‘does not have HCM’. Echocardiographers should exercise their judgement, but when the echocardiographic images show unequivocal evidence of HCM in an appropriate clinical context (clear-cut apical HCM, gross hypertrophy in a young patient and definite LVH in a screening echocardiogram), the phrase ‘consistent with HCM’ should be used.

Post-echocardiography Work-Up

In patients with suspected HCM, the clinical team will contextualise the echocardiography report with information regarding past medical and family history, blood tests and ECG results, and often cardiovascular magnetic resonance. In grey cases, clinicians judge whether the degree of hypertrophy matches the severity of the comorbidity (Figure 3). Clarification of the diagnosis in these instances is
possible after assessing the response of wall thickness and LV mass to a sustained period of reduced
afterload, for example, improved blood pressure control in the hypertensive patient, weight loss in
the obese individual, aortic valve replacement in the patient with severe aortic stenosis or cessation
of training in the athlete(18). In exceptional cases where there is non-apical hypertrophy measuring
less than 15 mm, and an ECG highly suggestive of underlying cardiomyopathy, the clinical team might
screen first-degree family members to look for clear-cut evidence of HCM. Given the likelihood of
finding a negative result on gene testing of confirmed cases, it is rarely used as a diagnostic tool when
there is ambiguity about the diagnosis.

Phenocopies

It is possible to find within the population of patients with hypertrophic cardiomyopathy rarer
conditions, called phenocopies or ‘mimics’(19). In general, these will come to light during clinical
evaluation of the patient’s medical history, family history, physical examination and the results of
blood tests, including genetics, and other imaging modalities. However, there are particular features,
termed ‘red flags’, which should alert the echocardiographer to the possibility of a phenocopy (Table
1). Of these, cardiac amyloidosis is the most obvious due to its classical signs: increased biventricular
wall thickness, poor long axis function, relative sparing of apical longitudinal contraction and global
longitudinal strain (although not pathognomonic of amyloid), interatrial and valvar thickening,
pericardial effusion, and mismatch between the degree of LVH seen on echo and low amplitude
voltages on ECG. Diagnosing HCM should be avoided immediately after an acute cardiac injury such
as myocarditis as the myocardium becomes oedematous and thickened; these changes resolve with
time.

Defining the pattern of hypertrophy in HCM

The echocardiographic report should detail the distribution of LVH using the schema described in
Figure 4 as this informs the clinical team of the likelihood of finding a disease-causing gene change;
being highest in patients with a reverse curvature pattern and lowest in those with a sigmoid septal
Right ventricular hypertrophy is present in around 20% of patients with HCM. The echocardiographer should report this as it occurs in disease mimics; however, it does not add to the likelihood of finding a disease-causing mutation.

Hypertrophy can also extend to the papillary muscles, which can contribute to mid-cavity obstruction. Additional morphological abnormalities of papillary muscles in HCM which can cause LVOT obstruction include antero-apical displacement, double bifid and anomalous papillary muscles which insert directly into the mitral valve leaflets. Bands running between the apex and basal anteroseptal wall are seen in HCM.

**Echo assessment in risk stratification and disease complications**

Risk stratification of sudden death is the process clinicians follow to decide which patients should receive an implantable cardioverter-defibrillator. Using the European Society of Cardiology (ESC) calculator, it is possible to generate an estimate of the five-year risk of sudden death and categorise patients into low, intermediate, and high-risk groups. Echocardiography provides three of the seven parameters required in the online tool (maximal wall thickness, LVOT gradient and 2D parasternal long axis left atrial size). To allow this critical information to be accessed rapidly by the referring clinician, the conclusion for every report in a patient with suspected or confirmed HCM should contain these parameters. Although not in the ESC risk calculator, the presence of left ventricular impairment and an apical aneurysm is also essential to include in the study conclusion as they modify risk of sudden cardiac death.

The importance of reporting cardiac rhythm in every echocardiogram report is particularly relevant in HCM as a significant proportion of patients will develop atrial fibrillation. The finding of new atrial fibrillation should be directly communicated to the referring team as anticoagulation is essential to prevent stroke or other embolic complications.
Heart failure can occur due to systolic impairment, diastolic dysfunction and LVOT obstruction. As a measure of systolic function, ejection fraction (EF) can be misleading in HCM being normal even when markers of systolic dysfunction such as abnormal regional wall motion and global longitudinal strain (26) (see the section below) are present. Nonetheless, the absolute value helps clinical teams to identify patients in whom systolic dysfunction is likely to develop (50-60%) and those in whom it is overt (<50%) (5). Accurately determining EF using Simpson’s biplane, and three-dimensional quantification where possible, and highlighting instances when this measurement is discordant with the systolic function will aid clinical management. Longitudinal systolic function should be assessed using tissue doppler imaging and in select cases strain (see section below), while radial systolic function should be assessed visually. Outcomes are generally adverse once the EF falls below 50% (5). Below this level, clinical teams should consider medications (1), heart transplant (1) and device therapy (27).

Diastolic dysfunction is common in HCM and results in elevated filling pressures and dilatation of the left atrium, whose diameter in the parasternal long axis is a predictor of sudden death in the ESC risk calculator (28), and of stroke and other thromboembolic events (29). Accurate classification of diastolic function grade is challenging in HCM due to the concomitant presence of left ventricular outflow tract obstruction and mitral regurgitation in many patients. Many independent echo variables have weak correlations with filling pressures. As such integration of several parameters is necessary to quantify diastolic function accurately. Diastolic function assessment should include Doppler tissue imaging and pulmonary vein Ar timings as per BSE guidelines (30).

It is essential to identify patients with preserved left ventricular ejection fraction but a restrictive diastolic filling pattern, which is often accompanied by pulmonary hypertension (31). These patients have adverse outcomes (32) and should be observed closely for evidence of deterioration as heart transplant is an option when symptoms related to heart failure are resistant to medical treatment (31).
Left ventricular outflow tract obstruction occurs as a result of a reduced cross-sectional area of the outflow tract due to hypertrophy, abnormalities of the mitral valve apparatus, and in most patients supranormal ejection, which drags the anterior mitral valve leaflet anteriorly towards the basal septum. The mitral valve coaptation is disrupted, with the resultant jet of mitral regurgitation in the majority of patients being directed posteriorly in mid-to-late systole (65% based on a recent study of patients undergoing myectomy with systolic anterior motion-related mitral regurgitation[33]). The same study found that posteriorly directed mitral regurgitation occurred in approximately a third of patients with intrinsic mitral valve disease.

There is a spectrum of LVOTO defined according to the severity and whether it is present at rest or with provocation (Table 2). Echocardiographers should try to provoke LVOT obstruction in every patient at the bedside by re-imaging while the patient is performing a Valsalva manoeuvre and in a seated and standing position. Obstruction in the mid and apical LV and right ventricle can also occur due to narrowing of the cavity as neighbouring myocardial walls contract towards each other. Accurate identification of the site of obstruction is relevant to guiding treatment strategies.

In patients who fail to respond to medical therapy directed at relief of LVOT obstruction, invasive septal reduction therapies (surgical myectomy and alcohol septal ablation) are considered. Given the potential complications of invasive therapies, it is important that patients fulfil the necessary clinical, anatomical, and hemodynamic criteria to determine suitability for a procedure, and this decision is based heavily on the echocardiographic assessment.

Although a complete discussion of the work-up for these procedures is outside this guideline’s remit, pertinent echocardiographic features are summarised in Table 3. A clear description of the nature of LVH, mitral valve abnormalities, additional areas of obstruction, and aortic valve disease supports decision-making. The focus is on identifying those elements that point to the need for surgical intervention and not alcohol septal ablation. Surgery can address features aligned with the latter, but the converse is not true for alcohol septal ablation.
Alcohol septal ablation is performed through an angiographic percutaneous approach and provides a suitable alternative for patients of advanced age or with significant comorbidities that would lead to an increased surgical risk. Injection of alcohol via a septal perforator branch of the LAD is performed into the target myocardium. This site is the hypertrophied basal septum adjacent to the point of anterior mitral valve leaflet-basal septal (systolic anterior motion-septal) contact, creating an acute infarction and progressive thinning of the myocardium with scar formation over a 6-12-month period. Selective intracoronary injection of contrast is essential to guide the selection of the appropriate septal perforator vessel, ensuring that the selected branch supplies only the targeted area of the myocardium, with no enhancement of remote areas such as the papillary muscles, inferior wall of the LV, or right ventricular free wall. A decrease in resting and provokable LVOT gradients is seen immediately because of myocardial stunning, with a progressive reduction in resting and dynamic LVOT gradients over 3-6 months.

Finally, the examination should include careful evaluation for aneurysm formation and associated thrombi in patients with apical HCM using contrast when necessary (Figure 2). Table 4 describes the relevance of various parameters captured by the echocardiography examination and Table 5 the minimum data set. A protocol for the transthoracic echo study in HCM is described in Table 6.

**Stress Imaging in HCM**

By imaging the heart during controlled exercise, stress echocardiography can unmask latent obstruction in symptomatic patients whose baseline transthoracic echocardiography – despite the previously described physiological manoeuvres – has not shown LVOT gradients ≥ 50 mmHg. Symptom-limited exercise is safe using an exercise bike or treadmill. There is some evidence to suggest that treadmill exercise can provoke higher LVOT gradients compared to semi-supine bicycle exercise(34). Dobutamine is not employed in HCM since this infusion can induce LVOTO in normal subjects. When the patient has reached peak exercise, images are obtained within 60-90 seconds to detect obstruction which can be present before or after the patient’s heart rate reaches 85% of target
heart rate. The protocol in Table 7 suggests an optimal scanning order to utilise peak heart rate with minimal changes to the acoustic window. Table 8 illustrates the data acquired in each step of the protocol. In specific scenarios, the echocardiographer can employ additional measures to provoke LVOT obstruction. For patients with postprandial symptoms, exercise after eating is useful (35) while for those who cannot exercise, administering GTN spray can unmask obstruction (36).

Strain Imaging in HCM

Measurement of global longitudinal strain (GLS) by two-dimensional speckle tracking echocardiography is becoming more widely used in current practice. Strain is a measure of myocardial deformation in multiple directions throughout the cardiac cycle. Most commonly, analysis based on the Lagrangian method (derived from speckle tracking techniques) expresses strain as a fractional change in length. Shortening of the myocardium becomes a negative value and lengthening of the myocardium a positive value (37). In HCM, reduced overall left ventricular GLS occurred in individuals with preserved ejection fraction (38). A recent systematic review has shown an association between abnormal GLS and adverse outcomes (39).

However, the author group feel that several practical considerations limit routine use in every HCM patient. These include the expertise and experience needed to ensure the strain curves generated are accurate and the potential difficulties in tracking where there is gross hypertrophy, apical hypertrophy or apical insertion of the papillary muscles. Consequently, inter-observer variability may well be higher in HCM than for dilated cardiomyopathies. Finally, strain-based measures are yet to be adopted into clinical HCM guidelines and so will not routinely alter patient management.

For this reason, we recommend that GLS is used to help distinguish HCM from cardiac amyloidosis, and athletic remodelling. This position will be re-evaluated in the next update of the guideline as more evidence emerges and the technology evolves.
Three-dimensional echocardiography

Besides enabling accurate quantification of left and right ventricular volumes and ejection fraction, three-dimensional echocardiography also allows echocardiographers to describe mitral valve and LVOT morphology. Three-dimensional technology is also valuable in transoesophageal echocardiography to detail SAM’s features and underlying causes (40,41). We recommend that patients undergo transoesophageal echocardiography when the transthoracic study suggests significant abnormalities of the mitral valve apparatus, and to evaluate both the mitral valve and the LVOT when planning for invasive septal reduction.

Recommendations

The echocardiogram report conclusion should include:

1. The following suggested phrases: when there is uncertainty: raises the possibility of HCM; when there is unequivocal evidence of HCM: consistent with HCM; for screening scans with no LVH: wall thickness is normal.

2. The presence of red flags pointing to a phenocopy.

3. The pattern of LVH: sigmoid septal, reverse curvature, apical or neutral.

4. The values for maximal wall thickness, LVOT gradient and LA size.

5. The presence or absence of disease complications.
   a. Left ventricular cavity size
   b. Systolic dysfunction with EF 50-60%, EF<50%.
   c. Diastolic dysfunction, specifically the presence of a restrictive filling pattern with preserved ejection fraction.
   d. Systolic anterior motion, mitral regurgitation, LVOT obstruction and other forms of obstruction; at rest and with provocation. Evidence of intrinsic mitral valve disease.
   e. Aneurysm formation.
6. Image quality, completeness of LV visualisation and need for contrast and transoesophageal echocardiography, and cardiovascular magnetic resonance.

Conclusion

Transthoracic echocardiography plays an essential role in the assessment of patients with proven or suspected HCM, and their first-degree family members. The guideline writing committee hopes that this document equips readers with the knowledge and tools needed to perform and report these studies to a uniformly high level.
Declaration of Interest

The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of this guideline.

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Figure Titles

- Figure 1. Measurement of Wall Thickness by Echocardiography
- Figure 2. Use of Contrast in Apical Hypertrophic Cardiomyopathy
- Figure 3. Thinking Underlying Clinical Decision-Making in HCM
- Figure 4. Different Phenotypes of Left Ventricular Hypertrophy in Hypertrophic Cardiomyopathy

Figure Legends

- **Figure 1.** The challenges to accurate wall thickness measurement vary at each left ventricular chamber level. Dashed lines represent erroneous measurements and solid lines accurate measurements.
- **Figure 2.** An apical four-chamber acquisition enhanced with contrast to show apical hypertrophic cardiomyopathy complicated by aneurysm formation.
- **Figure 3.** This schematic demonstrates various scenarios and the corresponding likelihood of the condition. Once investigations are complete, and a full clinical picture is available, this information is weighed by clinicians to reach a final diagnosis. Between cases where the likelihood of the condition is the same as the background population (left-hand side, green shading) and definite disease (right-hand side, green shading), lies the diagnostic grey zone (grey shading).
- **Figure 4.** Echocardiographic images are displayed for the four main patterns of hypertrophy, accompanied by the criteria for each pattern.
References

1. 2014 ESC Guidelines on diagnosis and management of hypertrophic cardiomyopathy: The Task Force for the Diagnosis and Management of Hypertrophic Cardiomyopathy of the European Society of Cardiology (ESC). Eur Heart J. 2014 Oct 14;35(39):2733–79.

2. Maron BJ, Gardin JM, Flack JM, Gidding SS, Kurosaki TT, Bild DE. Prevalence of Hypertrophic Cardiomyopathy in a General Population of Young Adults: Echocardiographic Analysis of 4111 Subjects in the CARDIA Study. Circulation. 1995 Aug 15;92(4):785–9.

3. Massera D, McClelland RL, Ambale-Venkatesh B, Gomes AS, Hundley WG, Kawel-Boehm N, et al. Prevalence of Unexplained Left Ventricular Hypertrophy by Cardiac Magnetic Resonance Imaging in MESA. J Am Heart Assoc. 2019 Apr 6;8(8).

4. Ho CY, Day SM, Ashley EA, Michels M, Pereira AC, Jacoby D, et al. Genotype and Lifetime Burden of Disease in Hypertrophic Cardiomyopathy. Circulation. 2018 Oct 2;138(14):1387–98.

5. Marstrand P, Han L, Day SM, Olivotto I, Ashley EA, Michels M, et al. Hypertrophic Cardiomyopathy With Left Ventricular Systolic Dysfunction: Insights From the SHaRe Registry. Circulation. 2020 Apr 28;141(17):1371–83.

6. Alfares AA, Kelly MA, McDermott G, Funke BH, Lebo MS, Baxter SB, et al. Results of clinical genetic testing of 2,912 probands with hypertrophic cardiomyopathy: expanded panels offer limited additional sensitivity. Genet Med. 2015 Nov;17(11):880–8.

7. Flett AS, Maestrini V, Milliken D, Fontana M, Treibel TA, Harb R, et al. Diagnosis of apical hypertrophic cardiomyopathy: T-wave inversion and relative but not absolute apical left ventricular hypertrophy. International Journal of Cardiology. 2015 Mar;183:143–8.

8. Suzuki J, Shimamoto R, Nishikawa J, Yamazaki T, Tsuji T, Nakamura F, et al. Morphological onset and early diagnosis in apical hypertrophic cardiomyopathy: a long term analysis with nuclear magnetic resonance imaging. J Am Coll Cardiol. 1999 Jan;33(1):146–51.

9. Norrish G, Jager J, Field E, Quinn E, Fell H, Lord E, et al. Yield of Clinical Screening for Hypertrophic Cardiomyopathy in Child First-Degree Relatives. Circulation. 2019 Jul 16;140(3):184–92.

10. van Velzen H, Schinkel A, Baart S, Oldenburg R, Frohn-Mulder I, van Slegtenhorst M, et al. Outcomes of Contemporary Family Screening in Hypertrophic Cardiomyopathy. Circulation: Genomic and Precision Medicine. 2018 Apr 1;11(4):e001896.

11. Coppini R, Ho CY, Ashley E, Day S, Ferrantini C, Girolami F, et al. Clinical Phenotype and Outcome of Hypertrophic Cardiomyopathy Associated With Thin-Filament Gene Mutations. J Am Coll Cardiol. 2014 Dec 23;64(24):2589–600.

12. van Velzen HG, Schinkel AFL, Oldenburg RA, van Slegtenhorst MA, Frohn-Mulder IME, van der Velden J, et al. Clinical Characteristics and Long-Term Outcome of Hypertrophic Cardiomyopathy in Individuals With a MYBPC3 (Myosin-Binding Protein C) Founder Mutation. Circ Cardiovasc Genet [Internet]. 2017 Aug [cited 2020 Nov 15];10(4). Available from: https://www.ahajournals.org/doi/10.1161/CIRCGENETICS.116.001660
13. Page SP, Kounas S, Syrris P, Christiansen M, Frank-Hansen R, Andersen PS, et al. Cardiac myosin binding protein-C mutations in families with hypertrophic cardiomyopathy: disease expression in relation to age, gender, and long term outcome. Circ Cardiovasc Genet. 2012 Apr;1(2):156–66.

14. Sheikh N, Papadakis M, Carre F, Kervio G, Panoulas VF, Ghani S, et al. Cardiac adaptation to exercise in adolescent athletes of African ethnicity: an emergent elite athletic population. Br J Sports Med. 2013 Jun;47(9):585–92.

15. Sheikh N, Papadakis M, Schnell F, Panoulas V, Malhotra A, Wilson M, et al. Clinical Profile of Athletes With Hypertrophic Cardiomyopathy. Circ Cardiovasc Imaging. 2015 Jul;8(7):e003454.

16. Harkness A, Ring L, Augustine DX, Oxborough D, Robinson S, Sharma V. Normal reference intervals for cardiac dimensions and function for use in echocardiographic practice: a guideline from the British Society of Echocardiography. Echo Research and Practice. 2020 Mar;1(7):G1–18.

17. Caselli S, Maron MS, Urbano-Moral JA, Pandian NG, Maron BJ, Pelliccia A. Differentiating Left Ventricular Hypertrophy in Athletes from That in Patients With Hypertrophic Cardiomyopathy. The American Journal of Cardiology. 2014 Nov;114(9):1383–9.

18. Pelliccia A, Maron BJ, De Luca R, Di Paolo FM, Spataro A, Culaslo F. Remodeling of Left Ventricular Hypertrophy in Elite Athletes After Long-Term Deconditioning. Circulation. 2002 Feb;105(8):944–9.

19. Rapezzi C, Arbustini E, Caforio ALP, Charron P, Gimeno-Blanes J, Heliö T, et al. Diagnostic work-up in cardiomyopathies: bridging the gap between clinical phenotypes and final diagnosis. A position statement from the ESC Working Group on Myocardial and Pericardial Diseases. Eur Heart J. 2013 May;34(19):1448–58.

20. Binder J, Ommen SR, Gersh BJ, Van Driest SL, Tajik AJ, Nishimura RA, et al. Echocardiography-guided genetic testing in hypertrophic cardiomyopathy: septal morphological features predict the presence of myofilament mutations. Mayo Clin Proc. 2006 Apr;81(4):459–67.

21. Kwon DH, Setser RM, Thamilarasan M, Popovic ZV, Smedira NG, Schoenhagen P, et al. Abnormal papillary muscle morphology is independently associated with increased left ventricular outflow tract obstruction in hypertrophic cardiomyopathy. Heart. 2008 Oct;94(10):1295–301.

22. Klues HG, Roberts WC, Maron BJ. Anomalous insertion of papillary muscle directly into anterior mitral leaflet in hypertrophic cardiomyopathy. Significance in producing left ventricular outflow obstruction. Circulation. 1991 Sep;84(3):1188–97.

23. Lentz Carvalho J, Schaff HV, Morris CS, Nishimura RA, Ommen SR, Maleszewski JJ, et al. Anomalous papillary muscles-Implications in the surgical treatment of hypertrophic obstructive cardiomyopathy. J Thorac Cardiovasc Surg. 2020 Apr 15;

24. Gruner C, Chan RH, Crean A, Rakowski H, Rowin EJ, Care M, et al. Significance of left ventricular apical–basal muscle bundle identified by cardiovascular magnetic resonance imaging in patients with hypertrophic cardiomyopathy. Eur Heart J. 2014 Oct;35(39):2706–13.
25. Rowin E, Maron B, Haas T, Garberich R, Wang W, Link M, et al. Hypertrophic Cardiomyopathy With Left Ventricular Apical Aneurysm: Implications for Risk Stratification and Management. J Am Coll Cardiol. 2017 21;69(7):761–73.

26. Hiemstra Y, Debonnaire P, Bootsma M, van Zwet E, Delgado V, Schalij M, et al. Global Longitudinal Strain and Left Atrial Volume Index Provide Incremental Prognostic Value in Patients With Hypertrophic Cardiomyopathy. Circulation: Cardiovascular Imaging. 2017 Jul 1;10(7):e005706.

27. Maron MS, Rowin EJ, Wessler BS, Mooney PJ, Fatima A, Patel P, et al. Enhanced American College of Cardiology/American Heart Association Strategy for Prevention of Sudden Cardiac Death in High-Risk Patients With Hypertrophic Cardiomyopathy. JAMA Cardiol. 2019 Jul 1;4(7):644.

28. O’Mahony C, Jichi F, Pavlou M, Monserrat L, Anastasakis A, Rapezzi C, et al. A novel clinical risk prediction model for sudden cardiac death in hypertrophic cardiomyopathy (HCM risk-SCD). Eur Heart J. 2014 Aug 7;35(30):2010–20.

29. Guttmann OP, Pavlou M, O’Mahony C, Monserrat L, Anastasakis A, Rapezzi C, et al. Prediction of thrombo-embolic risk in patients with hypertrophic cardiomyopathy (HCM Risk-CVA). Eur J Heart Fail. 2015 Aug;17(8):837–45.

30. BSE Guidelines for Diastolic Function - in press.

31. Rowin E, Maron B, Kiernan M, Casey S, Feldman D, Hryniewicz K, et al. Advanced Heart Failure With Preserved Systolic Function in Nonobstructive Hypertrophic Cardiomyopathy. Circulation: Heart Failure. 2014 Nov 1;7(6):967–75.

32. Biagini E, Spirito P, Rocchi G, Ferlito M, Rosmini S, Lai F, et al. Prognostic Implications of the Doppler Restrictive Filling Pattern in Hypertrophic Cardiomyopathy. The American Journal of Cardiology. 2009 Dec 15;104(12):1727–31.

33. Hang D, Schaff HV, Nishimura RA, Lahr BD, Abel MD, Dearani JA, et al. Accuracy of Jet Direction on Doppler Echocardiography in Identifying the Etiology of Mitral Regurgitation in Obstructive Hypertrophic Cardiomyopathy. Journal of the American Society of Echocardiography. 2019 Mar 1;32(3):333–40.

34. Reant P, Dufour M, Peyrou J, Reynaud A, Rooryck C, Dijos M, et al. Upright treadmill vs. semi-supine bicycle exercise echocardiography to provoke obstruction in symptomatic hypertrophic cardiomyopathy: a pilot study. European Heart Journal - Cardiovascular Imaging. 2018 Jan 1;19(1):31–8.

35. Feiner E, Arabadjian M, Winson G, Kim B, Chaudhry F, Sherrid MV. Post-Prandial Upright Exercise Echocardiography in Hypertrophic Cardiomyopathy. Journal of the American College of Cardiology. 2013 Jun;61(24):2487–8.

36. Zemánek D, Tomašov P, Homolová S, Linhartová K, Veselka J. Sublingual isosorbide dinitrate for the detection of obstruction in hypertrophic cardiomyopathy. Eur J Echocardiogr. 2011 Sep 1;12(9):684–7.

37. Hoit BD. Strain and Strain Rate Echocardiography and Coronary Artery Disease. Circ Cardiovasc Imaging. 2011 Mar;4(2):179–90.
38. Haland TF, Almaas VM, Hasselberg NE, Saberniak J, Leren IS, Hopp E, et al. Strain echocardiography is related to fibrosis and ventricular arrhythmias in hypertrophic cardiomyopathy. Eur Heart J Cardiovasc Imaging. 2016 Jun;17(6):613–21.

39. Tower-Rader A, Mohananey D, To A, Lever HM, Popovic ZB, Desai MY. Prognostic Value of Global Longitudinal Strain in Hypertrophic Cardiomyopathy: A Systematic Review of Existing Literature. JACC: Cardiovascular Imaging. 2019 Oct 1;12(10):1930–42.

40. Nampiaparampil RG, Swistel DG, Schlame M, Saric M, Sherrid MV. Intraoperative Two- and Three-Dimensional Transesophageal Echocardiography in Combined Myectomy-Mitral Operations for Hypertrophic Cardiomyopathy. Journal of the American Society of Echocardiography. 2018 Mar;31(3):275–88.

41. Vainrib A, Massera D, Sherrid MV, Swistel DG, Bamira D, Ibrahim H, et al. Three-Dimensional Imaging and Dynamic Modeling of Systolic Anterior Motion of the Mitral Valve. J Am Soc Echocardiogr. 2020 Oct 12;

42. Gersh B, Maron B, Bonow R, Dearani J, Fifer M, Link M, et al. 2011 ACCF/AHA Guideline for the Diagnosis and Treatment of Hypertrophic Cardiomyopathy. Circulation. 2011 Dec 13;124(24):e783–831.

43. Nistri S, Olivotto I, Betocchi S, Losi MA, Valsecchi G, Pinamonti B, et al. Prognostic significance of left atrial size in patients with hypertrophic cardiomyopathy (from the Italian Registry for Hypertrophic Cardiomyopathy). Am J Cardiol. 2006 Oct 1;98(7):960–5.

44. Maron MS, Olivotto I, Betocchi S, Casey SA, Lesser JR, Losi MA, et al. Effect of Left Ventricular Outflow Tract Obstruction on Clinical Outcome in Hypertrophic Cardiomyopathy. New England Journal of Medicine. 2003 Jan 23;348(4):295–303.

45. Harris K, Spirito P, Maron M, Zenovich A, Formisano F, Lesser J, et al. Prevalence, Clinical Profile, and Significance of Left Ventricular Remodeling in the End-Stage Phase of Hypertrophic Cardiomyopathy. Circulation. 2006 Jul 18;114(3):216–25.

46. Augustine DX, Coates-Bradshaw LD, Willis J, Harkness A, Ring L, Grapsa J, et al. Echocardiographic assessment of pulmonary hypertension: a guideline protocol from the British Society of Echocardiography. Echo Res Pract. 2018 May 11;5(3):G11–24.

47. American Heart Association Writing Group on Myocardial Segmentation and Registration for Cardiac Imaging:; Cerqueira MD, Weissman NJ, Dilsizian V, Jacobs AK, Kaul S, et al. Standardized Myocardial Segmentation and Nomenclature for Tomographic Imaging of the Heart: A Statement for Healthcare Professionals From the Cardiac Imaging Committee of the Council on Clinical Cardiology of the American Heart Association. Circulation. 2002 Jan 29;105(4):539–42.

48. Ho C, Sweitzer N, McDonough B, Maron B, Casey S, Seidman J, et al. Assessment of diastolic function with Doppler tissue imaging to predict genotype in preclinical hypertrophic cardiomyopathy. Circulation. 2002 Jun 25;105(25):2992–7.

49. Patel P, Dhillon A, Popovic ZB, Smedira NG, Rizzo J, Thamilarasan M, et al. Left Ventricular Outflow Tract Obstruction in Hypertrophic Cardiomyopathy Patients Without Severe Septal Hypertrophy: Implications of Mitral Valve and Papillary Muscle Abnormalities Assessed Using Cardiac Magnetic Resonance and Echocardiography. Circ Cardiovasc Imaging. 2015 Jul;8(7):e003132.
# Tables

## Table 1. Echocardiographic Clues to the Presence of Phenocopies

| Condition                  | Echocardiographic ‘Red Flags’ which raise the possibility of a phenocopy * |
|----------------------------|---------------------------------------------------------------------------|
| Cardiac amyloidosis        | Thickened interatrial septum, mitral and tricuspid valves and right ventricular free wall, mild to moderate pericardial effusion. Ground-glass appearance of the myocardium. Global hypokinesia (with and without LV dilatation) in TTR amyloidosis. Markedly reduced longitudinal function, relative sparing of apical longitudinal contraction/global longitudinal strain, a mismatch between LVH on echo and low amplitude voltages on ECG. |
| Fabry disease              | Thickened mitral and tricuspid valves and right ventricular free wall, concentric LVH, Global hypokinesia (with and without LV dilatation). |
| Myocarditis                | Thickened right ventricular free wall, mild to moderate pericardial effusion, global hypokinesia (with and without LV dilatation) |
| Danon disease              | Extreme concentric LVH, global hypokinesia (with and without LV dilatation) |
| Pompe disease              | Extreme concentric LVH |
| PRKAG2 mutations           | Global hypokinesia (with and without LV dilatation) |
| Glycogenosis               | Concentric LVH |
| Mitochondrial disease      | Global hypokinesia (with and without LV dilatation) |
| Noonan syndrome and associated disorders | Right ventricular outflow tract obstruction |

*Adapted from Rapezzi et al(19) and Elliott et al(1).
Table 2. Definition of LVOT obstruction.

| LVOT gradient at rest and with physiological provocation | Definition                          |
|---------------------------------------------------------|-------------------------------------|
| Gradient $\geq 30$mmHg at rest                          | Basal or resting obstruction        |
| Gradient $\leq 30$mmHg at rest and $\leq 30$mmHg after provocation | Non-obstructive                     |
| Gradient $\leq 30$mmHg at rest but $> 30$mmHg with physiological provocation | Labile, provable or dynamic obstruction |

Adapted from Gersh et al. (42)

Table 3. Use of Echocardiography When Determining Optimal Invasive Septal Reduction Approach

| Favours surgical myectomy | Aligned with alcohol ablation strategy | Unfavourable for either                  |
|----------------------------|--------------------------------------|-----------------------------------------|
| Septal thickness $> 25$mm  | Focal basal septal hypertrophy or sigmoid septal morphology | Apical hypertrophy                       |
| Central or anteriorly-directed mitral regurgitation due to intrinsic valve disease | Posteriorly-directed mitral regurgitation secondary to systolic anterior motion |                                         |
| Abnormal mitral subvalvar apparatus contributing to obstruction | | Mid-cavity obstruction |
| Concomitant aortic valve disease or coronary artery disease necessitating CABG | | |
Table 4. Rationale for Key Echo Parameters in Hypertrophic Cardiomyopathy

| Feature                        | Prognostic Relevance                                                                 | Role in ESC HCM Guidelines(1)                                                                 |
|-------------------------------|--------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------|
| Left atrial diameter          | Sudden cardiac death (28), with >48 mm predicting all-cause mortality(43). Risk of thromboembolism increases exponentially (29) | In risk calculator. If LA >45mm, for six to twelve monthly ambulatory monitoring               |
| Indexed Left atrial volume    | >34 mL/m² predicts all-cause mortality, heart transplantation, sudden cardiac death, and appropriate implantable cardioverter-defibrillator therapy (26) |                                                                                                 |
| Mitral valve filling pattern  | Restrictive filling pattern in HCM patients with heart failure with preserved ejection fraction carry adverse prognosis HCM(32) |                                                                                                 |
| Left ventricular wall thickness| Sudden cardiac death (28)                                                            | In risk calculator.                                                                                                                                         |
| Left ventricular outflow tract obstruction | >30 mmHg predictor of sudden cardiac death and heart failure(28,44) | In risk calculator. If the patient has symptoms and > 50 mmHg LVOTO resistant to medical therapy, invasive septal reduction indicated |
| Left ventricular function      | Ejection fraction <50% associated with unfavourable outcome(45)                      | When ejection fraction <50% in patients with NYHA III-IV despite optimal medical therapy, heart transplant indicated |

Table 5. Minimum Dataset

| Structure and Function          | Measurement                                                                 |
|--------------------------------|-----------------------------------------------------------------------------|
| Left atrium size               | Diameter (mm)                                                               |
| Mitral valve inflow Doppler    | E wave (m/s)                                                                |
| Pulmonary venous Doppler       | Systolic wave (m/s)                                                         |
| Mitral regurgitation           | Severity                                                                    |
| Systolic anterior motion       | Yes/No                                                                       |
| Left ventricle wall            | Septum at basal                                                             |
|                               | Indexed biplane volume (mL/m²)                                              |
|                               | A wave (m/s)                                                                 |
|                               | Diastolic wave (m/s)                                                        |
|                               | Mechanism                                                                   |
|                               | Valvular or chordal                                                          |
|                               | Lateral wall at basal                                                        |
|                               | Inferior wall at basal                                                       |
thickness in short axis view | level, papillary muscle level and apex level (mm) | basal level, papillary muscle level and apex level (mm) | level, papillary muscle level and apex level (mm) | level, papillary muscle level and apex level (mm)
--- | --- | --- | --- | ---
LV dimensions | End diastolic dimension (cm) | End systolic dimension (cm) |  |  
LV volumes | End-diastolic Volume (ml), indexed to body surface area (ml/m²) | End-systolic Volume (ml), indexed to body surface area (ml/m²) | Systolic Volume (ml) |  
LV systolic function | Ejection fraction by Simpson’s Biplane (%) | Ejection fraction by visual assessment when Simpson’s Biplane cannot be calculated (%) | Global longitudinal strain in selected cases (%) |  
Tissue Doppler Imaging | Anterolateral annulus (Sm, E’, A’) (cm/s) | Inferoseptal annulus (Sm, E’, A’) (cm/s) | Anterior annulus *(Sm, E’, A’) (cm/s) | Inferior annulus* (Sm, E’, A’) (cm/s) 
LVOT or intra-cavity obstruction (defining which) | Resting (mmHg) | Valsalva (mmHg) | Sitting (mmHg) | Standing (mmHg) 
Right ventricle (RV) | Size and function | RV hypertrophy (mm) | RV outflow tract obstruction (mmHg) |  
Tricuspid regurgitation and inferior vena cava | Severity | Probability of pulmonary hypertension(46) | Inferior vena cava size and collapse response |  

1. In individuals being screened for HCM

2.

3.
Table 6. Transthoracic HCM protocol
| Measurement                          | View  | Modality | Explanation                                                                                                                                                                                                 | Image |
|-------------------------------------|-------|----------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------|
| LA diameter                         | PLAX  | 2D       | Measure LA dimension at end-systole just after the aortic valve closes using 2D acquisition as per BSE normal reference intervals guidelines (16). LA diameter is one of the criteria used in ESC risk calculator of SCD. Record in report conclusion. |       |
| SAM                                 | PLAX  | M-mode   | Place M-mode cursor through the MV leaflet tips, ensuring image is on-axis. Involves MV leaflets and/or chordae.                                                                                              |       |
| Feature of LVOT obstruction         | PLAX  | M-mode   | Mid-systolic notching and coarse systolic fluttering of the aortic valve are ancillary echocardiographic features in LVOTO.                                                                               |       |
| Contact plaque                      | PLAX, A3C | 2D     | Increased echogeneticity occurs in the basal anteroseptal wall due to fibrosis where leaflet contact occurs due to SAM.                                                                               |       |
| LV wall thickness measurements | SAX MV level | 2D | Freeze 2D image at end-diastole. Calliper diameter of maximal wall thickness – wherever it occurs - in the anterior, septum, inferior and lateral walls at the basal, mid-ventricular and apical levels(47). Be careful not to include right ventricular (RV) wall, papillary muscles, trabeculations or moderator band. The thickest segment may not be in the septum. Maximal wall thickness is one of the criteria used in ESC risk calculator of SCD. Record in report conclusion. |
|---------------------------------|-------------|----|--------------------------------------------------------------------------------|
| LV Simpson’s Biplane volumes and ejection fraction | A4C, A2C    | 2D Units: mL/m^2 and % | LV volumes should be obtained using 2D imaging from A4C and A2C, and wherever possible 3D imaging. Trace the endocardial border. LV length is defined as the distance between the midpoint of the mitral valve level line and the most distal point of the LV apex. Take care to ensure the LV is not foreshortened. Papillary muscles and trabeculations are excluded from the volumes and considered part of the chamber. Measure at end-diastole (onset of QRS complex) and end-systole (the frame before MV opens, where AV just closes)(16). Volumes are indexed to |
| LA biplane volume | A4C, A2C | 2D biplane volume using independent A4C and A2C views. Units: ml/m² | LA volume should be obtained from apical 4- and 2-chamber windows (separated by 60° of rotation), optimised for LA assessment, using the biplane Simpson’s method. Maximal LA volume should be obtained from the frame immediately prior to mitral valve opening. Values should be reported after indexing for BSA (16,30). Trace the inner aspect of the left atrial wall. At the mitral valve level, the contour is closed by a straight line between along the plane of the mitral valve annulus. Exclude left atrial appendage and pulmonary veins. |

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| TDI velocities | PW TDI | Systolic (Sm), early (E') and atrial (A') relaxation velocities at anterolateral and inferoseptal walls(30). In screening studies, there is an argument for averaging E' across anterolateral, inferoseptal, inferior and anterior LV annulus as a value <13.5 cm/s can be useful in identifying genotype positive phenotype negative individuals(48). |
|----------------|--------|------------------------------------------------------------------------------------------------ |
| in all four    | A4C, A2C |                                                                                                  |
| walls          | Units: cm/s |                                                                                                  |
|                |        |                                                                                                  |
| Global longitudinal strain (GLS) | A4C, A2C, A3C | 2D |  
|-------------------------------|---------------|----|  
| Units: -%                     | This is recommended when cardiac amyloidosis or athletic remodelling are being considered. Average global longitudinal strain (GLS) is calculated using the apical long axis (A3C), four chamber A4C and two chamber A2C standard views. High quality image acquisition, maintaining a frame rate of 40 to 90 frames/second at a stable heart rate is key. Clear endocardial and epicardial definition (seen throughout the cardiac cycle) is required to ensure adequate segmental tracking during systole and diastole. Markers are placed in each of the respective basal and apical regions, utilising automated tracking where possible to maintain reproducible results. ROI should be manipulated as required to fit the myocardium. Automated tracking should also be combined with a visual assessment of tracking in each view across the whole region of interest including the endocardial and epicardial border. If more than two segments in any one view are not adequately tracked, the calculation of GLS should be avoided. |
| LVOT or intra-cavity obstruction gradients | A4C, A5C | CW Doppler (or PW with HPRF as a significant gradient will alias on PW Doppler). Sampling PW Doppler throughout the LV cavity is a useful tool to pinpoint the exact location of obstruction if unclear on colour. Units: mmHg | Assess obstruction gradients at rest, with Valsalva manoeuvre and in sitting and standing positions. Align CW Doppler through entire turbulent colour flow for peak obstruction velocity. Peak LVOT obstruction gradient is one of the criteria used in ESC risk calculator of SCD. Record in report conclusion. |
| Multiple LV gradients | A4C, A5C | CW Doppler Colour flow mapping Units: mmHg | Intra-cavity obstruction at the apex produces an additional Doppler signal to the LVOTO signal. |
| MR versus LVOT obstruction | A4C, A5C | CW Doppler Colour flow mapping Units: mmHg | When mitral regurgitation occurs in the context of SAM or prolapse, its onset is later in mid to late systole. Otherwise, its onset is in early systole helps distinguish it from the LVOT signal which begins later in systole (see right hand image). LVOT obstruction is dagger-shaped due to the progressive decrease in LVOT orifice size as systoles progresses but of |
Mitral regurgitation compared to LVOTO. The lower image shows superimposed CW envelopes in a patient with mitral regurgitation and LVOTO. In this case mitral regurgitation starts later in systole, so timing of onset is a less useful discriminator. However, the velocity for the mitral regurgitation signal is far higher than for LVOTO.

| Condition                        | View(s)       | Method               | Description                                                                                     |
|----------------------------------|----------------|----------------------|-------------------------------------------------------------------------------------------------|
| Mitral regurgitation secondary to SAM | PLAX, A4C, A5C | Colour flow mapping | Mitral regurgitation quantification may be limited as the PISA dome may merge with turbulent LVOT flow. Mitral regurgitation secondary to SAM is mainly posteriorly directed. When quantitative assessment of MR is precluded by LVOTO, other indicators of MR severity should be considered. For example, an E velocity of < 1.3 m/s and an E/A ratio <1 are strongly suggestive of non-severe MR. |
| Abnormal MV anatomy (elongated AMVL) | PLAX, A4C, A3C | 2D                  | Describe MV anatomy; elongation of both leaflets, presence of SAM (and which leaflet(s) it involves), aberrant chordae running from anterior mitral valve leaflet to LVOT, anomalous papillary muscles running directly into the mitral valve leaflets and displacement of the papillary muscles antero- apically. If the anterior mitral valve leaflet is elongated (>16 mm), this increases the likelihood of LVOT. |
| Pulmonary venous Doppler | A4C       | PW Units: cm/s | Measure peak systolic (S) velocity, peak diastolic (D) velocity, the S/D ratio, peak atrial reversal (Ar) velocity in late diastole and the duration of the Ar velocity. In the apical 4-chamber view, superior angulation of the transducer and use of colour flow will help locate the pulmonary veins. This angle often brings the aorta into the visualised plane. The right upper is usually easiest and is next to the atrial septum. If the signal is weak, ask the patient to adopt a more supine position. Place the PW Doppler sample volume (1–3 mm) 1–2 cm into the right upper vein. Wall filter settings should be lowered (100–200 MHz). Aim to include clear visualisation of the atrial reversal velocity waveform. Measurements should be averaged over 3 cardiac cycles, at end expiration. Additional parameters for diastolic function should include A wave duration on transmitral inflow. For the measurement of the mitral valve A wave duration, the PW Doppler sample should be placed at the level of the |
annulus rather than at the leaflet tips. This provides a cleaner signal for the start and end of the wave. See BSE guidelines for diastolic function(30).

| TR jet velocity and probability of pulmonary hypertension | RV inflow, PSAX, A4C | CW Colour flow mapping |
|----------------------------------------------------------|----------------------|------------------------|
|                                                          | Units: Vmax m/s, peak gradient mmHg |
|                                                          | See BSE PHTN guidelines for risk of pulmonary hypertension(46). |

| RV Hypertrophy | Subcostal view, PLAX | 2D |
|----------------|----------------------|----|
|                | Units: mm            |
| Freeze the PLAX or subcostal view of the RV free wall, scroll to end diastole and calliper the RV wall thickness. |

| RVOT obstruction | PSAX view | 2D Colour flow mapping CW Doppler. |
|------------------|-----------|-----------------------------------|
|                   | Units: mmHg |
| Modify both the RV inflow and outflow to assess for RV hypertrophy and RV outflow tract obstruction. Use colour box as a guide for highest RVOT velocity. |
| Phenotype | Imaging VIEWS | Description |
|-----------|---------------|-------------|
| Septal myectomy and septal ablation | PLAX, PSAX MV level, A4C, A3C, subcostal views. | 2D Basal septum has scalloped appearance and is hypokinetic/akinetio. Colour flow Doppler should be applied to the area of myectomy to assess for iatrogenic VSD (systolic flow), and a denuded septal perforator vessel (diastolic flow). The pre-procedure HCM morphology cannot be determined in patients who have undergone a septal myectomy or septal ablation. |
| Aneurysmal apex | A4C, A2C, A3C, PSAX apex level. +/- ultrasound enhanced echo with contrast | 2D colour flow mapping Contrast Apical HCM can be accompanied by an apical aneurysm which encourages thrombus formation (see non-contrast image on right). Have a low threshold for giving contrast (far right image) if endocardial definition is poor at the apex. |
| HCM Phenotypes | A4C, A2C, A3C, PLAX, PSAX. | 2D Four distinct phenotypes describe the distribution of left ventricular hypertrophy. Comment on morphology in the report conclusion. |
Table 7. Stress Echocardiography protocol in HCM

| View | Modality | Explanation |
|------|----------|-------------|
| LVOT or intra-cavity obstruction | A5C/A3C (view which obtained the highest gradient at rest). | CW Doppler (or PW with HPRF as a significant gradient will alias on PW Doppler). Sampling PW Doppler throughout the left ventricular cavity is a useful tool to pinpoint the exact location of obstruction if unclear on colour. Increase in stroke volume with exercise. Use colour box as a guide to aim CW Doppler beam through area of turbulence obtaining the highest gradient. Assessment of LVOT obstruction assessment is performed prior to LV assessment it can be a short-lived phenomenon. |
| MR | A4C, A3C | Colour mapping CW doppler Be careful to differentiate mitral regurgitation from LVOT obstruction. |
| MV | A4C | PW Doppler Units: cm/s Peak exercise and intermediate stage (100-120bpm). Pulse at MV leaflet tips to obtain inflow Doppler. Description of MV morphology and SAM at intermediate and peak. |
| TR | A4C (alternative views are RV inflow or PSAX, however time consuming as requires a different window) | CW Doppler Units: mmHg (m/s) To exclude exercise induced pulmonary hypertension. |
| LV size and systolic function | A4C, A2C, A3C, SAX | 2D imaging Systolic TDI velocities in anterolateral and inferoseptal walls Units: cm/s A4C and A2C for LV volumes and Simpson’s Biplane EF. Small LV cavity may make measuring volumes difficult at intermediate and peak stress. |
| LV diastolic function | A4C, A2C | Diastolic TDI parameters in anterolateral and inferoseptal walls MV inflow flow Doppler E/e’ average Units: cm/s Peak exercise and intermediate stage (100-120bpm). E/A fusion will occur at high heart rates. Intermediate imaging with supine bicycle only. |
### Table 8. Illustrated Guide to Stress Echocardiography in HCM

| COLUMN C | ROW 1 (C1) | HCM stress echo protocol – Quick guide |
|---|---|---|
| 1. Echo data – rest | Resting BSE HCM guidelines 2020.  
Exclude contraindications to exercise test. |

| COLUMN C | ROW 2 (C2) |  |
|---|---|---|
| 2. Resting haemodynamics | Perform a resting ECG.  
Obtain resting BP and standing BP. |

| COLUMN C | ROW 3 (C3) |  |
|---|---|---|
| 3. Resting spirometry | Obtain resting spirometry tests if performing combined CPEX.  
CPEX data is used to establish exercise capacity and true exercise limitations. |

| COLUMN C | ROW 4 (C4) |  |
|---|---|---|
| 4. Exercise modality | Bicycle or treadmill method of exercise  
Treadmill – resting echo images obtained on echo bed.  
Bicycle – resting echo images obtained whilst patient on bike to ensure comparable echo windows. |

| COLUMN C | ROW 5 (C5) | Image 1  
Image 2 |
|---|---|---|
| 5. Exercise haemodynamic data | Continuous monitoring of ECG and BP throughout study.  
Pay particular attention to arrhythmias, ST changes and potential BP drop at peak exercise. |

| COLUMN C | ROW 6 (C6) |  |
|---|---|---|
| 6. Transition from treadmill to bed | Treadmill – stopped immediately at peak exercise, patient is carefully guided back onto the echo bed.  
Bicycle – peak images are obtained whilst patient is still on bicycle. |

| COLUMN C | ROW 7 (C7) |  |
7. Echo data – peak

- Peak exercise images are obtained within 60-90s.
- This is before preload decreases and before the patient’s heart rate recovers below 85% of THR.
- See table 1 for echo parameters collected at peak exercise.
- Echo measurements are calculated post acquisition to utilise time at peak HR.

| Peak | Obstruction (mmHg) | PVR | TR |
|------|--------------------|-----|----|
| LV volumes | EDV | ESV | SV | EF |
| TDI | E | E' | A' |
| Septal | | | |
| Lateral | | | |
| Anterior | | | |
| Inferior | | | |

8. Report

- CPEX, echo and haemodynamic data are combined to produce a clinical report.
| SEGMENT | BASAL                                                                 | MID-VENTRICULAR                                                                 | APICAL                                                                 |
|---------|------------------------------------------------------------------------|--------------------------------------------------------------------------------|-------------------------------------------------------------------------|
| PITFALLS (depicted by dashed lines) | Erroneously including septomarginal trabeculation and aberrant LV papillary muscles in measurement | Underestimating hypertrophy at the confluence of the mid inferior and inferoseptal wall, including RV papillary muscles | Missing apical hypertrophy, especially lateral wall (dotted line with arrows) |
| APPROACH (depicted by solid lines) | Exclude these components at their attachments using continuation of endo- and epicardial curve (dotted lines) | Exclude these components at their attachments using continuation of endo- and epicardial curve (dotted lines) | Clue in often abnormal ECG   <br>Look for ‘akinetiic’ apex<br>Use contrast to confirm |

Figure 1.

338x190mm (96 x 96 DPI)
Figure 2

338x190mm (96 x 96 DPI)
| CLINICAL INFORMATION PROVIDED AND ECHO FINDINGS | GREY ZONE | CLINICAL INFORMATION PROVIDED AND ECHO FINDINGS |
|-----------------------------------------------|-----------|-----------------------------------------------|
| Any age 14 mm Severe aortic stenosis           | Over 60 year old Afro-Caribbean patient 15-20 mm with poorly controlled hypertension | Any age 13 mm aortic wall Gross T wave inversion on ECG |
| Older patient 14 mm Basal septal hypertrophy Hypertension | Over 60 year old Afro-Caribbean patient >20 mm Hypertension | Younger patient 215 mm No past medical history |
| Younger patient 13 mm Severe myocarditis 16 mm wall thickness | Over 60 year old caucasian patient 15-20 mm Hypertension High BMI Gross ECG changes | Abnormal ECG First degree family member of someone with confirmed HCM ≥ 13 mm No past medical history |
| Athlete, normal ECG and no relevant family history | Same as background population risk | Likely |

Unlikely

Definite
338x190mm (96 x 96 DPI)