CASE REPORT

Routine orthostatic LVOT gradient assessment in patients with basal septal hypertrophy and LVOT flow acceleration at rest: please stand up

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Summary

A 70-year-old female with exertional dyspnoea was found to have basal septal hypertrophy (BSH), or a ‘basal septal bulge’, with evidence of mild left ventricular outflow tract obstruction (LVOT) at rest on her initial echocardiogram. She was usually fit and well with no significant past medical history. She had no history of hypertension. She had never smoked. There was no family history of hypertrophic cardiomyopathy (HCM). A cardiac MRI did not demonstrate any typical features of HCM. ECG showed sinus tachycardia with a rate of 101 bpm but was otherwise unremarkable. She was referred for exercise echocardiography to assess for latent LVOT obstruction. Prior to commencing exercise, her LVOT gradient was re-assessed at rest. Her LVOT gradients were 30 mmHg at rest, 49 mmHg during Valsalva and 91 mmHg on standing. A diagnosis of significant latent LVOT obstruction was made and the patient was started on bisoprolol, a cardioselective beta-blocker. Bisoprolol was slowly uptitrated from 1.25 mg to 5 mg once daily, following which the patient reported a significant improvement in her symptoms with an improved exercise capacity. Follow-up echocardiography demonstrated a dramatic reduction in LVOT gradient, with a maximum of 11 mmHg assessed both with Valsalva and on standing. This case is a reminder that patients with a ‘common’ basal septal bulge can develop significant LVOT obstruction, the symptoms of which may respond to pharmacological therapy. Orthostatic assessment of LVOT gradient using echocardiography should be considered during standard LVOT obstruction provocation manoeuvres such as a Valsalva.

Learning points:

- Differentiation between basal septal hypertrophy (BSH) and hypertrophic cardiomyopathy (HCM) may be challenging. Key factors favouring HCM include a positive family history of HCM or sudden cardiac death, septal thickness >15 mm/posterior wall thickness >11 mm, systolic anterior motion of the anterior mitral valve (SAM), late gadolinium enhancement on cardiac MRI, a causative genetic mutation associated with HCM and an abnormal ECG.
- Significant LVOT obstruction may develop in patients with BSH and is potentially responsive to pharmacotherapy.

Key Words

- basal septal hypertrophy
- orthostatic LVOT assessment
- latent LVOT obstruction
- provokable gradient
Standing reduces venous return, resulting in decreased LV volume. Compensatory mechanisms to maintain cardiac output involve sympathetic nervous system activation leading to increased LV contractility and subsequent increased LVOT gradient. Significant LVOT obstruction may be unmasked by an orthostatic posture. Orthostatic LVOT gradient assessment should be part of the routine echocardiographic assessment of all patients with an increased LVOT gradient at rest. The post-prandial state has been associated with increased LVOT gradient due to splanchnic dilatation and the consequent increased cardiac output required to maintain blood pressure. Post-prandial status should therefore be considered when assessing LVOT gradient.

Background

The 2015 European Society of Cardiology (ESC) guidelines recommend provocation testing to assess for latent LVOT obstruction in all patients with exertional symptoms who have either hypertrophic cardiomyopathy (HCM) or isolated basal septal hypertrophy (BSH) (1). LVOT obstruction is a major cause of symptoms in HCM and has been associated with a worse prognosis (1). BSH, often referred to as a ‘sigmoid septum’ or a ‘ventricular septal bulge’, is common in elderly patients, particularly those with hypertension. The differentiation between this very common finding and genetically inherited HCM can be difficult. Canepa et al. proposed that a family history of HCM or sudden cardiac death, presence of symptoms, septal thickness >15 mm/posterior wall thickness >11 mm, systolic anterior motion of the anterior mitral valve (SAM) and LVOT obstruction, late gadolinium enhancement on cardiac MRI, a genetic mutation associated with HCM and an abnormal ECG all make HCM more likely versus BSH (2). Importantly patients with BSH can have latent LVOT obstruction and there is evidence demonstrating improvement in symptoms with pharmacological treatment (3). Aortoseptal angulation has also been shown to be predictive of latent outflow tract obstruction, with smaller angles in patients with provable LVOT obstruction (4).

In most echo departments, standard outpatient provocation testing involves measuring the LVOT gradient at rest and following a Valsalva manoeuvre. If no significant gradient is induced in a symptomatic patient, stress testing is usually implemented at this stage. Shah et al. demonstrated that 2/3rd (62.1%) of patients who had no previous documented LVOT obstruction (LVOT gradient ≤30mmHg) developed LVOT obstruction during exercise (5). Approximately 20% of these patients went on to have invasive treatment with subsequent improvement in symptoms (5).

It is recognised that following a meal mesenteric vasodilatation occurs, resulting in reduced peripheral vascular resistance. As a compensatory mechanism, heart rate and stroke volume increase to maintain blood pressure (6). Using radionuclide imaging, Kelbaek et al. demonstrated post-prandial increases in cardiac output of 62% due to increased heart rate and stroke volume (6). This post-prandial haemodynamic change is well recognised to increase LVOT gradient and exacerbate symptoms (7). Protocols used for the assessment of latent LVOT obstruction should therefore take into consideration post-prandial status.

LVOT gradient is sensitive to preload. Bedside manoeuvres that affect preload will therefore change the outflow gradient. Squatting increases preload and therefore reduces the gradient through the outflow tract, whereas standing up reduces preload and has the opposite effect. The reduction in venous return caused by standing up also results in decreased LV volume. Compensatory mechanisms to maintain cardiac output involve sympathetic nervous system activation leading to increased LV contractility, which also contributes to increased LVOT gradient (8). A number of studies have demonstrated increased LVOT gradient in the standing versus the supine position (7, 8, 9).

Crucially, most patients will be in a fully upright position when symptoms are usually experienced. Orthostatic assessment of LVOT gradient is therefore more representative of ‘real-life’ haemodynamics and is commonly employed during stress testing to look for outflow obstruction.

Case presentation

A 70-year-old female presented with an 18-month history of exertional breathlessness occurring on walking up inclines. She had no history of chest pain or syncope.
She had no relevant past medical history. She specifically had no history of hypertension. She had never smoked. There was no family history of HCM or sudden unexplained death. Regular medications included only prophylactic antibiotics. Cardiac examination revealed a resting regular tachycardia with a heart rate of 101 bpm (Fig. 1). Resting blood pressure was 134/80 mmHg. She was comfortable at rest and was clinically euvoletic. Pulse character was normal. There was a loud ejection murmur heard throughout the precordium with normal

**Figure 1**
12 lead electrocardiogram at initial presentation.

**Figure 2**
Continuous wave Doppler through the left ventricular outflow tract (A) at rest (B) on standing (C) with Valsalva and the (D) parasternal long axis view with measurements.
first and second heart sounds. Examination was otherwise unremarkable.

**Investigation**

Her resting ECG showed a sinus tachycardia with a rate of 101 bpm and no other significant abnormality (Fig. 1). Initial echocardiogram demonstrated a hyperdynamic left ventricle with BSH measuring 15 mm (Fig. 2 and Videos 1, 2). There was a peak outflow gradient of 30 mmHg measured in the left lateral position at rest. This increased to 49 mmHg with a Valsalva. There was no significant SAM; however, there was mild mitral regurgitation. No other pathology was demonstrated.

**Video 1**

**Video 2**

She had a cardiac MRI to investigate for any evidence of cardiomyopathy, specifically HCM. This reconfirmed isolated BSH and with a maximum thickness of 15 mm. There was no late gadolinium enhancement, and native T1 relaxation times were normal. Adenosine stress imaging was performed, which demonstrated no inducible ischaemia.

In view of the mild left ventricular outflow tract (LVOT) gradient of 30 mmHg detected at rest and ongoing unexplained breathlessness, she was referred for stress echocardiography to assess for provokable LVOT obstruction during exercise.

The patient was instructed to eat a small meal, such as a sandwich, 1 h prior to the test. This is the recommendation made to all patients in order to standardise the impact of splanchnic dilatation. Pre-exercise LVOT gradient was measured at rest, with Valsalva and on standing. The resting gradient on continuous wave Doppler was 30 mmHg, on Valsalva this increased to 49 mmHg and on standing further increased to 91 mmHg (Fig. 2).

Treadmill stress was not deemed appropriate as significant obstruction had been demonstrated.

**Treatment and outcome**

Given the demonstration of a significant outflow tract gradient on standing, the patient was commenced on 1.25 mg of bisoprolol. The dose was increased to 5 mg once daily, which resulted in an improvement in her symptoms. The echocardiogram was repeated, at which point the patient had a heart rate of 74 bpm. The ventricle no longer appeared hyperdynamic at rest and peak gradient through the LVOT with provocation manoeuvres fell to 11 mmHg (Fig. 3 and Videos 3, 4).

**Video 3**

**Video 4**

**Discussion**

Latent LVOT obstruction should be considered in all symptomatic patients with either HCM or BSH. In most
echo departments, standard provocation testing in
the outpatient setting involves measuring the LVOT
gradient at rest and following a Valsalva manoeuvre. If
no significant gradient is demonstrated in a symptomatic
patient, stress testing is usually implemented at this stage.
It is possible, however, that a significant gradient
could be induced simply by standing the patient up, as
shown in this case. If provokable outflow obstruction is
demonstrated on standing, there is no requirement for
further stress testing at this stage.

Post-prandial status is crucial as the degree of
splanchnic dilatation will have an impact on the LVOT
gradient. A large meal will induce splanchnic dilatation
with a resultant increase in LVOT gradient due to a
compensatory increase in cardiac output (7). Starving
patients prior to assessment of LVOT gradient will
result in a lower inducible gradient and therefore the
diagnosis of latent LVOT obstruction could be ‘missed’.
Outflow tract gradient is also sensitive to preload. Bedside
manoeuvres that affect preload will therefore change
the outflow gradient. Squatting increases preload and
therefore reduces the gradient through the outflow
tract, whereas standing up reduces preload and has the
opposite effect. Crucially, most patients will be in a fully
upright position when symptoms are usually experienced.
Orthostatic assessment of LVOT gradient is therefore
more representative of ‘real-life’ haemodynamics and
should be carried out during standard echocardiographic
assessment. The ultimate goal should be to replicate the
environment and haemodynamics during which a patient
experiences symptoms.

A significant gradient is generally considered to be
an LVOT gradient >50mmHg (1). Lifestyle advice such
as avoiding large meals and dehydration, which can
exacerbate the LVOT gradient, should be given to all
patients. First-line treatment is pharmacological and has
been shown to be beneficial in LVOT obstruction caused
by HCM and BSH (3). Bisoprolol is a cardioselective
b-blocker, which is specific for beta-1 receptors found in
the heart. It is negatively inotropic and therefore reduces
contractility and heart rate. This has the effect of reducing
the LVOT gradient and can improve symptoms in patients
with hypertrophic obstructive cardiomyopathy. The
maximum dose is 10mg daily and patients should be
up titrated to the maximum tolerated dose.

If non-hydropyridine beta-blockers are not tolerated
or contraindicated, non-hydropyridine calcium channel
blockers, such as verapamil, can be started at 40mg TDS
and up titrated to 480mg SR once daily if required (1).
ESC HCM guidelines recommend caution when using
verapamil in patients with an LVOT gradient >100mmHg
due to the risk of pulmonary oedema (1).

In patients with resistant symptoms, disopyramide
can be introduced in addition to either beta-blockers or
calcium channel blockers. Disopyramide is a class IA anti-
arrhythmic drug and works by blocking sodium channels.
It reduces LVOT gradient due to its negative inotropic
effect. It should be used with caution in patients with atrial
arrhythmia due to the risk of enhanced AV conduction and
therefore increased ventricular rates (1). It is usually
started at a dose of 100mg BD and can be increased to
400–600mg/day (1). Its use is generally limited by anti-
cholinergic side effects such as a dry mouth, urinary
retention and dry eyes. ESC guidelines state that the QTc
interval should be monitored during dose up titration and
the dose should be reduced if the QTc exceeds 480ms (1).

Patients with HCM who have persistent moderate-
severe symptoms and LVOT obstruction despite optimal
pharmacotherapy should be referred for consideration
of invasive therapy, such as myectomy or alcohol septal
ablation (1). The use of invasive therapies in patients with
LVOT obstruction due to BSH is not well documented.


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