Abstract

The aetiology of sudden cardiac arrest can often be identified to underlying cardiac pathology. Mitral valve prolapse is a relatively common valvular pathology with symptoms manifesting with increasing severity of mitral regurgitation (MR). It is unusual for severe MR to be present without symptoms, and there is growing evidence that this subset of patients may be at increased risk of sudden cardiac arrest or death. The difficulty lies in identifying those patients at risk and applying measures that are appropriate to halting progression to cardiac arrest. This article examines the association of mitral valve prolapse with cardiac arrests, the underlying pathophysiological process and the strategies for identifying those at risk.

Case

A 45-year-old male had a sudden collapse at home, witnessed by their partner who started bystander cardiopulmonary resuscitation (CPR). The initial observed cardiac rhythm was pulseless ventricular tachycardia on arrival of the emergency medical services. A direct current shock was delivered resulting in asystole. The patient underwent a further 10 min of CPR prior to the return of spontaneous circulation, during which time endotracheal intubation and positive pressure ventilation were commenced. The total low-flow time was between 20 and 30 min. The patient was transported by air ambulance to a tertiary cardiac arrest centre. The patient was transferred to the accident and emergency department where intravenous sedation was started and maintained. The body temperature was measured at 34.9°C. Initial blood investigations were as follows: troponin I 350 ng/mL, C-reactive protein <4 mg/mL, leucocyte count 18 × 10⁹/µL, sodium 140 mmol/L, potassium 3.1 mmol/L, urea 6.6 mmol/L, creatinine 66 µmol/L and glucose 12 mmol/L. Initial arterial blood gas sampling demonstrated a pH 7.31, PaCO₂ 6.41 kPa, PaO₂ 62.9 kPa, base excess −1.7 and lactate 3.0 mmol/L. The ECG showed sinus rhythm, there were no signs of ischaemia and it fulfilled electrical criteria for left ventricular hypertrophy (Fig. 1). A chest X-ray showed the presence of an endotracheal tube, but it was otherwise unremarkable. A CT scan of the head was obtained which was reported as normal. Collateral history determined that the patient had no significant co-morbidities and that the patient was healthy prior to the sudden cardiac event. A bedside transthoracic echocardiogram was obtained (Videos 1 and 2). This reported a thickened and prolapsing anterior mitral valve leaflet with associated severe mitral regurgitation (MR). The left ventricular ejection fraction was inappropriately normal, but not hyperdynamic; although impaired when the severe MR was taken into account. There was no evidence of a systolic regional wall motion abnormality.
On this evidence, the patient was treated empirically with vancomycin and gentamicin for suspected infective endocarditis and transferred to the intensive care unit.

**Video 1**

**Video 2**

In line with current protocols, the patient’s temperature was allowed to increase to 36°C and maintained at this level for the first 24h. An infusion of norepinephrine was commenced to maintain a mean arterial blood pressure greater than 65mmHg. A transoesophageal echocardiography (TOE) was performed 12h after admission (Videos 3 and 4). This confirmed a flail A2 segment secondary to chordal rupture with evidence of myxomatous degeneration. The mitral annulus was dilated at 5.2cm. No vegetations were observed on the mitral valve. Doppler interrogation confirmed the presence of severe MR. The left ventricle (LV) diastolic dimension was 7.2cm, the systolic dimension was 5.3cm and the ejection fraction (Simpson’s biplane) was 58%. A coronary angiogram was performed and showed normal unobstructed coronary arteries. Empirical antibiotic therapy for bacterial endocarditis was discontinued following TOE.

**Video 3**
Sudden cardiac death (SCD) is an unexpected natural death from a cardiac cause within a short time period (1). In most epidemiologic studies, this short period is defined within 1h from the onset of symptoms. UK data show that an incidence of SCD is 100,000 adults per year (2). In the USA, it accounts for about 300,000 cases annually representing about 50% of mortality from cardiac causes (3). The overall incidence is about 50–100:100,000 people per year. On average, the survival with good neurologic recovery after OOHCA is about 5–10% (4). Due to the short time period from the onset of symptoms to arrest, identification of the high-risk population and prevention is the most effective strategy. Causes of SCD include coronary artery disease, cardiomyopathies, structural heart disease and primary electrophysiologic abnormalities. In some patients, the cause remains unclear, and hence, the term ‘idiopathic ventricular fibrillation’ is used (4).

**Definition**

MVP is defined as displacement of mitral leaflet tissue into the left atrium past the mitral annular plane during systole (5). It was first described by Barlow in the 1960s as an auscultatory and cine-angiographic phenomenon, prior to the availability of diagnostic echocardiography (6, 7, 8). Advances in echocardiography (e.g. transoesophageal echocardiography (TOE) and three-dimensional imaging) have made it possible for accurate diagnosis and quantification of MR (9).

**The role of echocardiography in MVP**

Echocardiography can be used for diagnosis, surveillance and assessment of interventions in MVP. Carpentier's functional classification of MR described MVP (Type II classification) as an abnormality of leaflet motion, where one or several components of the valve protrude into the left atrium (LA) during ventricular systole (10). Two-dimensional (2D) echocardiography can be used to divide MVP into classical and non-classical criteria for diagnosis (11). Classical MVP describes >2mm displacement of the mitral valve leaflets into the LA in long-axis view during ventricular systole, with a leaflet thickness ≥ 5 mm. Non-classical MVP describes >2mm leaflet displacement with a leaflet thickness < 5 mm.

Classical MVP will have either a symmetrical or an asymmetrical point of coaptation. Both leaflet tips meet at the same point on the mitral valve annulus in symmetrical MVP. Asymmetrical coaptation results in one leaflet being displaced towards the LA in relation to the other leaflet. Asymmetrical coaptation is more likely to deteriorate and develop flail prolapse and result in greater severity of MR. Flail segment or prolapse describes the presence of leaflet tips that turn outwards and point into the LA. Flail prolapse can involve a single segment, multiple segments, one leaflet, or both leaflets (likely secondary to chordal rupture).

Both 2D TTE and TOE can be used to evaluate mitral valve morphology (Table 1). TOE will more reliably provide superior views across the LA window and should be considered in all cases of MVP assessment. The diagnosis of MVP using TTE should only be made in the parasternal long-axis view and/or the apical long-axis view as the hyperbolic paraboloid shape of the mitral valve annulus can give a false-positive diagnosis of MVP.
(12). In addition, a description of the leaflet thickness or redundancy, annular dilatation and chordal length should be included. Visual accuracy of mitral valve shape and deformity may be improved using 3D echocardiography techniques, especially for anterior leaflet or commissural involvement (13, 14).

Doppler imaging is essential to determine the severity of MR. This should involve quantitative measures to determine disease progression, predict outcome and assess suitability for intervention. It is recommended that the colour flow Doppler area should not be used to quantify the severity of MR. Where feasible, the vena contracta or proximal isovelocity surface area should be used as a measure of severity (12). Both the pulsed Doppler mitral-to-aortic time velocity–integral ratio and the systolic pulmonary flow reversal can be used as adjuncts to assist with quantifying the severity of MR (12).

The downstream effects of MR, including LA dilatation, LV dilatation, LV dysfunction, pulmonary vein flow reversal, pulmonary hypertension, RV dilatation and tricuspid regurgitation will also help determine the severity of MR. LV dilatation is a particularly important marker of progression in asymptomatic regurgitation, with monitoring of LV-end systolic diameter used to suggest when surgical intervention may be indicated.

Intraoperatively 2D TOE and/or 3D TOE is recommended to assist surgical repair or replacement of the valve (14). Three-dimensional TOE has been shown to be more reliable than surgical inspection at accurately diagnosing the cleft-like indentations of the posterior mitral valve leaflet of myxomatous mitral valve disease and aiding repair (15). It is also important in percutaneous methods of mitral valve repair such as with MitraClip (16).

Epidemiology

According to recent figures, the MVP prevalence is 1–2.4% (17, 18). This is down from previous estimations of up to 35%. This difference can be explained by better understanding the anatomy of the mitral valve, stricter diagnostic criteria and better diagnostic technology. Nevertheless, MVP remains the most common cause of MR in developed countries.

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Pathophysiology of MVP

Several pathological processes may cause prolapse of the mitral valve, including rheumatic heart disease, Marfan's syndrome, endocarditis and myocardial ischemia, but degenerative MVP refers specifically to a specific spectrum of primary lesions (5, 19). On the two extremes of the spectrum are fibroelastic deficiency (FED) and Barlow's disease. FED is a fibrillin deficiency leading to rupture of one of the chordae. In this case, the mitral valve leaflets are thinned and the annular size is normal (18, 20). At the other end of the spectrum is Barlow's disease affecting younger patients. The mitral annulus may become calcified and dilated with thickened leaflets secondary to myxomatous degeneration (19, 20).

Arrhythmias

Both supraventricular (SVA) and ventricular arrhythmias (VA) are associated with complications of MVP. This high incidence rate has been recognised for more than 20 years. A study in 1994 reported an incidence of SVA and VA associated with MVP of 14 and 30%, respectively (21). When tested with continuous ECG monitoring, MVP patients had a prevalence of VA as high as 34% with premature ventricular contractions as the most common pattern (66% of cases). Moderate-to-severe MR has been shown to be an independent risk factor to developing arrhythmias (22). Ikeda studied patients with idiopathic VT and found a high prevalence of MVP (12 out of 35 patients). VT originated from the LV in most of these cases in contrast to non-MVP (91.7 vs 69.6%) (23). Abnormal ventricular repolarisation, including early repolarisation, has also been linked to the presence of MVP (24, 25). A cross-sectional study of 100 patients with MVP showed a high incidence of early repolarisation (represented by notch in the descending arm of QRS and J-point and/or ST segment changes) compared with healthy individuals without MVP (25).
Left ventricular remodelling

The presence of a dilated left ventricle in the context of severe MR may indicate a period of LV remodelling. In acute primary MR, afterload may initially decrease due to the alternate pathway for ejection. However, with volume loading of the LV over time, the relatively thin-walled LV may dilate and hypertrophy. Consequently, the afterload in chronic compensated MR will be normal and elevated in chronic decompensated MR (26). Remodelling of the LV may allow MR to be tolerated with mild or no symptoms by increasing the stroke volume. However, progression to heart failure and possibly cardiac arrest can occur rapidly often in the presence of myocyte dysfunction and sympathetic activation (27). LV remodelling has been associated with evidence of VAs. However, there is a paucity of evidence assessing this link in the context of MVP without coronary artery disease (28).

Incidence of SCD

Despite the general impression of a benign course, several case reports since the 1980s describe SCD in MVP patients, with a significant proportion of young and previously asymptomatic individuals (29, 30). A debate continues as to whether MVP is the cause of SCD or merely an association. Data from forensic autopsy examinations have reported floppy mitral valves in 5% of specimens (99 out of 2007 specimens). From those, MVP was considered directly responsible for the death of 17 patients (0.8% of cases) (31). This would rank MVP as the most common congenital and valvular cardiac cause of SCD. A consensus statement in 1997 from American and European societies does not attribute the cause of SCD to MVP, unless the prolapse is associated with valve redundancy, thickening and regurgitation, QT prolongation or ST-T wave changes (32).

It is not known if the mitral valve repair or replacement will have a preventive effect on VAs. Subsequently, the American Heart Association/European Society of Cardiology guidelines for VAs and SCD have no distinctive recommendations for the management of VA or SCD in mitral valvular heart disease (33). Moreover, neither the recent American nor the European valvular heart disease guidelines mention a criterion for predicting or assuming SCD secondary to MVP (33, 34).

The 10-year mortality of asymptomatic MVP patients is reported to be 19%, with greater than moderate MR and impaired LV function as the most important risk factors (35). This draws similar comparisons to the all-cause mortality from MR with a flail leaflet (MR-FL). Collective data from the all-cause mortality from MR-FL showed an annual mortality of 1.8%. Not surprisingly, left ventricular ejection fraction and NYHA class were the risk factors for mortality. However, in the subgroup of patients with SCD, 40% of patients were categorised as NYHA class I (36). This highlights a subgroup of patients with SCD predominantly due to cardiac arrhythmia and not related to the severity of MR or LV failure in MVP. These patients were mostly young and asymptomatic. The subgroup of patients with NYHA class I had a yearly risk of SCD of 1% (36). This is equal to the overall mortality in hypertrophic obstructive cardiomyopathy (HCM, a pathology considered as one of the most common causes of SCD in young people (37). Consequently, it is logical to identify a high-risk group of MVP similar to the recommendation for primary prevention in HCM.

Risk of SCD

As the incidence of SCD is very low in patients with MVP, studies have been directed towards identifying a high-risk subgroup that could benefit from primary or secondary prevention. Autopsy evaluation of SCD patients associated with MVP identified a subgroup of patients with isolated MVP. These patients were younger, mostly females and with less prevalence of MR (38). The high prevalence of MVP makes a strategy based on primary prevention feasible if the high-risk group could be strictly defined. Advances in echocardiography, cardiac magnetic resonance (CMR) and electrophysiology are promising in the identification of this group. Currently, no clear strategy is recommended for this small subgroup. Being young and asymptomatic makes it difficult to detect patients for primary prevention. However, we know that the incidence of SCD is much lower in young adults than in the general population (i.e. about 1:100,000). This incidence doubles if athletic (39). It is not known whether exercise increases the risk of SCD in MVP patients. Till now, MVP-associated SCD has been categorised as idiopathic VF. The real question is whether it is possible to prospectively identify risk factors for SCD in individuals with MVP.

A retrospective American study identified 12 out of 50 SCD patients with idiopathic VF as having MVP. A quarter of MVP patients had a family history of SCD (40). Another study by Vohra and coworkers in 1993 prospectively studied seven patients with MVP associated with mild MR and normal LV function. All patients initially presented
with syncope or OOHCA and the mean follow-up period was 2.5 years. They had proven VAs on Holter or electrophysiological study (EPS). Patients were treated with anti-arrhythmic pharmacotherapy, although with limited efficacy as there were two who suffered sudden death. Two patients underwent reparative surgery, but the arrhythmias were re-inducible on repeat EPS (41). This is in contrast to another study in which surgical correction had a protective effect (36).

**Prevention of SCD**

A cohort study that reviewed 24 patients who had idiopathic VF/VT OOHCA found echocardiographic evidence of bileaflet MVP in 10 patients (42%). The author named it malignant bileaflet MVP. Those patients with MVP had higher incidence of ECG abnormalities and VAs (42). Electrolyte disturbances can be an aggravating factor contributing to the occurrence of VAs and subsequently SCD (43). Screening programmes, technical availability (ECG, echocardiography) and cost remain obstacles for a wide application of primary prevention. Nevertheless, if MVP had been detected accidentally or in a screening program (e.g. athletes or relatives of patients with SCD), it may be easier to reconsider recommending a strategy for those patients depending on clear risk criteria. However, an implantable cardiac defibrillator (ICD) is not without risk, especially for young people. EPS, MRI and echocardiography may be helpful in identifying the right group.

Secondary prevention is an easier question. In patients diagnosed with idiopathic VF, ICD is a class I recommendation for survivors (44). A follow-up of 24 patients with OOHCA with VF/VT showed patients with bileaflet MVP had a significantly better ICD appropriateness compared with non-MVP patients (80 vs 36%, P=0.04). However, this may have been affected by a longer follow-up period in the MVP group (42).

**Cardiac magnetic resonance imaging**

The advent of CMR has shed further light on this condition. CMR has shown an association between MVP and papillary muscle fibrosis. In a study of 16 patients, 8 had a history of VA on previous Holter monitoring (couples of VPCs or non-sustained VT) and 63% of the MVP patients had papillary muscle fibrosis on late gadolinium enhancement (45). Basal LV hypertrophy is another abnormality, which can be better detected by CMR in such patients. However, its significance in cases of arrhythmias and SCD remains unknown (46).

**Other causes of SCD with MVP**

It should be noted that other possible mechanisms of cardiac arrest should not be ignored. Primary spontaneous chordal rupture is one of the recognised complications of MVP. This may cause acute MR and cardiogenic pulmonary oedema (47). Intramyocardial small vessel disease has a known association with SCD (48). One of its variants, fibromuscular dysplasia, had been observed by pathologic examination more frequently in MVP than in the controls (75 vs 25%). This variant of small vessel disease in MVP cases was associated with fibrosis of the basal interventricular septum (49).

**Conclusion**

In this case, the patient was previously asymptomatic and presented with an aborted SCD. Severe MR was demonstrated with echocardiography. The presence of a dilated LV at presentation and later on during CMR suggests chronic severe MR with extensive LV remodelling in the absence of symptoms. It is possible that the haemodynamic effects of the acute regurgitation may have caused syncope, which progressed to dysrhythmia and cardiac arrest due to reduced coronary flow. It is equally possible that there was a primary dysrhythmia associated with MVP or left ventricular remodelling. Early TOE and CMR are important, as is a greater understanding of the possible electrophysiological mechanisms of primary arrhythmogenesis.

**Declaration of interest**

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

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