**EHRA/EACVI expert consensus statement on arrhythmic mitral valve prolapse and mitral annular disjunction complex**

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**EHRA/EACVI 2022 Expert consensus statement: Arrhythmic Mitral Valve Prolapse and Mitral Annular Disjunction complex**

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**Abbreviations**

AMVP = arrhythmic mitral valve prolapse

CT = Computed-Tomography

CMR = cardiac magnetic resonance

DMR = degenerative mitral regurgitation

ECG = Electrocardiogram

ICD = implantable cardioverter defibrillator

ILR = implantable loop recorder

LA = left atrial

LGE = late gadolinium enhancement

LV = left ventricular

MAD = mitral annular disjunction

MMVP = myxomatous mitral valve prolapse

MR = mitral regurgitation

MRI = magnetic resonance imaging

MVP = mitral valve prolapse

NSVT = non-sustained ventricular tachycardia

PVC = premature ventricular contraction

PVS = programmed electrical stimulation

SCD = sudden cardiac death

TGF = transforming growth factor

TWI = T wave inversion

VA = ventricular arrhythmia

VF = ventricular fibrillation

VT = ventricular tachycardia

# **Introduction**

Mitral valve prolapse (MVP) is the most common valvular heart disease1 involving about 2-3% of the general population2-4 and is well characterized by echocardiography.5 While the outcome in MVP is mostly benign in the absence of mitral regurgitation (MR) and its left-ventricular (LV) consequences,6, 7 a small yet poorly defined subset of individuals remain at higher risk of malignant ventricular arrhythmias (VAs) and sudden cardiac death (SCD). This link between MVP and SCD is reported with an annual incidence <1% in unselected individuals with MVP.8, 9 However, at autopsy, the prevalence of MVP among young patients with sudden arrhythmic death is reported between 4%10 to up to 7%.11 Due to the low event-rate and the lack of very large cohorts, assessing the precise incidence of SCD in MVP in general and in specific subsets of patients remains challenging.

Mitral annular disjunction (MAD, discussed in detail in section 4) is often observed concomitantly with MVP12. MAD results in an abnormal motion of the mitral annulus, termed curling13. It is associated with increased risk of arrhythmias and is therefore and integral component of the AMVP complex14, 15.

Over the last decade, a multidisciplinary approach has been employed to identify specific MVP subsets with high arrhythmic risk by careful examination of Electrocardiograms (ECG), Doppler-Echocardiography, Cardiac Magnetic-Resonance-Imaging (CMR), cardiac computed-tomography-imaging (CT) and confirmed by autopsy findings16.

# **Scope of the document**

This consensus statement reviews and summarizes current literature regarding the AMVP complex based on an international multidisciplinary collaboration to collect evidence from experts in clinical cardiology, echocardiography, CMR, cardiac CT, electrophysiology and cardiothoracic surgery. It summarizes the main gaps of knowledge regarding identification, risk stratification and management of AMVP and provides practical suggestions for diagnosis and management of patients with AMVP. All recommendations were subjected to voting and required at least 75% agreement to reach consensus. The recommendations are ranked using the colored heart system (***Table 1: Categories of consensus statements***).

# **Aims of the document**

This document aims, using the most recent advances in knowledge regarding SCD and AMVP, to

1. Define the phenotype of the AMVP complex.
2. Propose approaches for screening and diagnosing the AMVP complex.
3. Propose approaches for risk stratification.
4. Propose comprehensive imaging protocols using transthoracic Doppler-Echocardiography and CMR to assess the AMVP complex.
5. Propose management approaches.
6. Highlight critical gaps of knowledge to guide future clinical trials and collaborative research.

# **Mitral valve prolapse (MVP)**

## **Definition and diagnostic criteria**

MVP is defined as a systolic displacement of one or both mitral leaflets ≥ 2mm above the plane of the mitral annulus in the sagittal view of the mitral valve (***Figure 1***).2, 17, 18 Two main etiologies of MVP are described19:

* Myxomatous MVP - also known as Barlow’s disease. Characterized by excess tissue including chordal thickening/elongation, annular dilation and calcification with low probability of chordal rupture.
* Fibroelastic Deficiency - characterized by chordal thinning, elongation, and/or high probability of rupture, with classic findings of prolapse and MR of varying severity. This is the most common form of MVP.20, 21

## **Epidemiology**

MVP is the most prevalent valve abnormality in Western countries2. Using the current definition, its prevalence varies from 0.6 to 3.1% depending on the age of the population examined (increasing with age) with slight predominance of female sex2-4 but consistent across ethnic groups.4 The prevalence of MVP among athletes is similar to that of the general population22.While the vast majority of MVP cases are sporadic, familial clusters have been reported.23 Parental MVP was associated with a ~5 fold increase in the risk of MVP, and first degree relatives have a 30-50% likelihood of being affected.24 Taken as a whole, long term survival of subjects with MVP is not different from the general population, with its prognosis determined mainly by MR severity.25

## **Mitral valve prolapse in the context of connective tissue diseases (syndromic MVP)**

Heritable connective tissue diseases with abnormal synthesis and processing of collagen proteins are often characterized by myxomatous MVP. The incidence of MVP in Marfan syndrome patients is linked with age, with up to 75% affected at >60 years.26-29 The reported prevalence of MVP is up to 45% in patients with Loeys-Dietz Syndrome. On the other hand, the prevalence of MVP is approximately 6% in patients with Ehlers-Danlos Syndrome29. Patients with Osteogenesis imperfecta and Adult polycystic kidney disease are also frequently reported to have MVP.26

## **Long term outcomes and association with sudden cardiac death**

Clinical outcome in MVP is primarily determined by both the presence of MR and its severity, as well as its consequences on LV function and size.6, 7 Indeed, in the absence of significant changes in MR or LV size/function, MVP is considered a benign condition with excellent prognosis.30, 31

However, reports of SCD25, 32 and VAs33 in MVP patients, with or without MR, raised the concern that a small subset may incur higher risk of VAs and SCD, independent of MR severity or LV dysfunction.34 SCD in patients with isolated MVP (with or without mitral annular disjunction) was initially described in case reports;11, 35-37. Later, the link between SCD and MVP was corroborated by a meta-analysis.38 Importantly, it appeared that MVP patients who suffered from SCD showed a characteristic MVP-phenotype.8, 37, 39

The prevalence of MVP in published juvenile SCD series ranges from 0% to 24%.8 In young SCD victims, prevalence of myxomatous MVP was between 4%10 and 7%, reaching 13% when considering only the female subgroup.11 The annual risk of SCD in populations with MVP is difficult to estimate due to low incidence. Some estimates range between 0.2% and 0.4% per year,9, 25, 40 but SCD rates were reported at 1.8% per year with severe MR due to a flail leaflet34, lower (0.8% per year) in asymptomatic patients with flail and normal ejection fraction, and as low as 0.14% in a community population.38 The excess risk of SCD is even more difficult to assess as the rates of SCD in the general population are highly dependent on age. 41 While some estimates place the risk of SCD in patients with MVP at 3-fold higher than in the general population,16 it is probably only slightly higher than normal in MVP without MR, doubled in MVP with severe DMR and markedly higher with signs of heart failure or LV dysfunction42.

Recently, large cohorts of consecutive patients with isolated MVP, comprehensively characterized with 24-hour-Holter-monitoring, clinical and echocardiographic assessment, demonstrated that VAs are frequent but are rarely severe.14 However, severe arrhythmias are independently linked to excess mortality accumulating over time, irrespective of the severity of degenerative MR.14 Thus it appears that there is a phenotype of MVP prone to developing ventricular arrhythmias over time with increased risk of mortality and SCD when VAs become severe.

# **Mitral annular disjunction (MAD)**

## **Definition**

MAD is characterized by a systolic separation between the ventricular myocardium and the mitral annulus supporting the posterior mitral leaflet,.43 Conversely, without MAD the mitral annulus remains attached to the atrial and ventricular myocardium/endocardium. Hence, MAD is associated with the loss of mechanical annular function linked to its normal ventricular myocardial attachment but with maintained electrical function, isolating the left atrium and ventricle electrophysiologically13, 40. The circumferential extension of MAD is limited anteriorly by the mitro-aortic fibrous continuity, between the aortic cusps and the anterior leaflet of the mitral valve. As a consequence, MAD has been observed only at the insertion of the posterior leaflet. It can extend laterally variably under all scallops of the posterior leaflet but preferentially at the central posterior scallop.

**Diagnostic criteria**

Diagnostically, the MAD trench is relatively easy to recognize in long-axis views by transthoracic-Echocardiography44-47 or MRI.47-49 Cardiac Computed-Tomography (CT) Scan can diagnose MAD.50, 51 DMR and MVP are defined based on the mitral annulus position,52 yet with MAD, the annulus is particularly difficult to identify precisely throughout systole as it falls briskly in mid-systole, posteriorly to the basal ventricular myocardium. Conversely, the ventricular myocardium, having lost its basal attachment, bulges more apically than normal, forming the apical margin of the MAD trench. This posterior movement of the mitral annulus is also known as “curling” and refers more broadly to the abnormal/excessive systolic upward motion of the posterior mitral annulus and of the adjacent postero-basal myocardium, with prominent basal myocardial wall thickening and hypertrophy.13 To recognize the mitral annulus position, it is recommended in long-axis views zoomed in on the mitral valve with the highest frame rate possible, to locate the thin structure of the annulus image by image from early to late systole (***Figure 2***).40 In turn, the precise location of the mitral annular position allows measurement of the MAD trench width and of MVP depth. Accordingly, the upper limit of MAD is defined at the level of posterior leaflet insertion on the annulus/left-atrial-wall, whereas the lower limit is defined at the level of the LV myocardium.49 Without dynamic examination, excessive posterior leaflet tissue arising from a normally implanted annulus could falsely be interpreted as MAD. Such dynamic analysis is required both on Echocardiographic and Cardiac MRI examinations.

MAD length is measured in the parasternal long axis view (or equivalent sagittal views on CMR), from the insertion of the posterior leaflet on the detached mitral annulus to the border of the bulging LV myocardium46, 48, 49, 53. Repetitive 2D-Transthoracic Echocardiography are of interest. MAD is described as a continuum around the annular circumference and is interspersed with normal tissue.48 Annulus disjunction can also be present on the right side of the heart.54

Also, MAD can be present in early childhood in syndromic MVP but whether MAD develops over time remains to be further explored.48, 55 MAD could additionally be assessed on 2D/3D transesophageal echocardiography56 when indicated for another reason (e.g. pre-operative examination) or in cases of inconclusive Transthoracic Echocardiography (***Figure 3***). Finally, MAD lateral extension should be determined in additional apical 4-chamber and 2-chamber views, by 2D-Transthoracic Echocardiography and CMR (***Figure 3 and 4***).48 The latter may be more sensitive for diagnosis.

|  |
| --- |
| Box 1-Summary of diagnosis criteria for MAD   * MAD is the anomalous attachment of the posterior leaflet, directly on atrial wall. * The slippage of the mitral annulus with MAD in systole is known as curling. * Sagittal views are recommended for identifying MAD and assessing its length. * Transthoracic Echocardiography is preferentially used as the first line imaging test for MAD diagnosis due to its wide availability with Cardiac-MRI reserved for patients with poor echocardiographic windows or image quality. * MAD diagnosis requires dynamic evaluation through the entire cardiac cycle. * MAD width extent should be evaluated in apical views. * Repeat 2D-Transthoracic Echocardiography over the years may be of interest to explore MAD development over time. |

## **Epidemiology**

MAD was first described in autopsy studies to be present in approximately 6% of human hearts,46, 48, 57-60. The reported prevalence of MAD varies partly due to the different imaging modalities, various cut-offs, heterogeneous subpopulations, and various MR-severity grades.44, 45, 61-64 Amongst patients with MVP, the prevalence of MAD varies between 20-58% (reported pooled rate of 32.6% from 3 studies).14, 44, 50, 63, 65-67 In the specific subset of MVP phenotype with myxomatous MVP, MAD prevalence varies between 21.8% and 98% (pooled rate of 50.8% from 3 studies).14, 45, 46, 63, 67, 68 In patients with MAD, reported presence of MVP was 78%.48

In syndromic MVP, MAD prevalence is reported between 34%69 in patients with Marfan syndrome and 40% in carriers of Loeys-Dietz syndrome.70 In these cases, MAD is a marker of severe disease including more arrhythmic events, a higher need for mitral valve intervention, and among patients with extensive MAD, more arrhythmic events.69, 70The first large prospective cohort of isolated MVP with systematic MAD assessment reported a prevalence of 30% of MAD, generally in younger patients40. Advanced myxomatous-degeneration, denoted by marked leaflet-redundancy and bileaflet-MVP, was the strongest MAD-associated MVP feature. The association was independent of all baseline characteristics, whereas MR-severity was not40.

# **The arrhythmic mitral valve complex/phenotype**

## **Arrhythmic MVP phenotypes: Proposed consensus definition and classification**

### **Arrhythmic mitral valve – Definition**

The arrhythmic mitral valve complex is defined by presence of MVP (with or without MAD), combined with frequent and/or complex ventricular ectopy in the absence of any other well-defined arrhythmic substrate (e.g. active ischemia, ventricular scar due to another defined ethology, primary cardiomyopathy or channelopathy).

Box 2- The diagnosis of arrhythmic MVP requires:

* The presence of MVP (with or without MAD)
* The presence of ventricular ectopy that is
  + Frequent (≥5% total PVC burden)  
    or
  + Complex (NSVT, VT, VF)
* The absence of any other well-defined arrhythmic substrate

### **Arrhythmic mitral valve – Phenotypes**

The writing committee recognizes 2 main phenotypes within the spectrum of this complex:

### **Severe degenerative mitral regurgitation (DMR)**

Patients with MVP and severe MR irrespective of valvular morphology are at increased risk of excess-mortality71 including excess SCD, compared to the general population.34 The risk of SCD is higher in patients with severe symptoms, atrial arrhythmias and reduced LV-systolic function. However, even without these risk factors, MVP patients with at least moderate to severe MR incur excess-mortality72 and SCD, with double the incidence compared to the general population.34 Surgical MR correction tends to normalize overall mortality72 and SCD incidence.34

### **Severe Myxomatous MVP independent of MR severity**

As a group, the overall survival of subjects with MVP without severe MR or LV dysfunction, is equivalent to that of the general population30. However, several reports of SCD in patients with MVP and in the absence of another relevant pathology,11, 25, 32, 33 have indicated the existence of a small subgroup of patients with a malignant course. Morphologically, this phenotype most often involves MAD, severe myxomatous degeneration with marked leaflet redundancy, excess leaflet length and thickness, and most often but not always, bileaflet MVP. MR may be absent to severe.14 This phenotype is not limited to young patients14 and is linked to more frequent occurrence of VAs. MAD appears as an important component of this phenotype. Importantly, the arrhythmic outcome of these patients is independent of gender, MR severity, LVEF or bileaflet-MVP.14

### **Atrial arrhythmia phenotype**

Patients with MVP and left atrial (LA) dilatation, in excess of that expected from MR, incur a higher risk of atrial arrhythmias.73-75 The combination of LA enlargement 75 and atrial fibrillation76 is associated with excess mortality, independent of all baseline characteristics including MR severity and LV parameters. Importantly, surgery is followed by improved survival, 75, 76 suggesting that both severe LA enlargement and atrial arrhythmias, even when paroxysmal, should be accounted for in the clinical decision making process.

### **MAD in the absence of MVP**

While MAD associated with MVP is frequent and associated with the occurrence of ventricular arrhythmic events40, isolated MAD without MVP has been described on static Cardiac-MRI examination.48 However, due to the problems associated with precise determination of the mitral annular position, its exact nature and whether MVP may have been under-detected remains uncertain. Even more unsubstantiated is its association with outcome. Accordingly, large cohorts with long-term follow-up are required to better define MAD without MVP and its outcome and consequences.

## **Determinants of arrhythmia**

### **Clinical risk factors**

Data derived form case reports and autopsies of SCD victims with MVP suggest excess female prevalence 37, 59, 77. This is likely resulting from selection bias as a nation-wide autopsy study found a similar proportion of both sexes60 and a large cohort of consecutive MVP patients did not find any association between sex and arrhythmia 14.

Several non-specific symptoms such as chest pain, palpitation, dyspnea on exercise and syncope have been attributed to MVP. Yet, multiple large cohort studies have shown that the incidence of these symptoms is comparable in subjects with and without MVP2, 78, 79. Nevertheless, syncope is uncommon in non-selected subjects with MVP (2-3.6%)2, 78 and was reported in 35% of MVP patients with malignant arrhythmia or SCD 59. Furthermore, in a large cohort of MVP patients, syncope was more frequently reported in patients with documented severe ventricular arrhythmia14. Therefore, syncope and particularly unexplained syncope, may have a high discriminative value in identifying MVP patients at risk of malignant arrhythmias. Palpitations and chest pain are frequent but are reported in similar proportions in subjects with MVP with and without history of ventricular arrhythmia14.

### **Electrocardiography**

### **T Wave inversion**

Repolarization abnormalities have long been recognized as markers of arrhythmic risk in subjects with MVP. The most consistent feature is T wave inversion (TWI), most often in the inferior and lateral leads (***Figure 5***). TWI or biphasic T waves, in the presence of normal QRS, have been observed in the majority (65%) of patients with MVP who present with malignant arrhythmia38 and more importantly, have an independent association with VAs in a large unselected cohort of MVP patients.14 This is thought to be the result of abnormal stretch of the papillary muscles and adjacent myocardium80, and/or locally disturbed repolarization of the same regions.81 However, the specificity of this finding, as a marker for high arrhythmic risk, is questionable since the reported rates of TWI range as high as 40% among subjects with MVP.82

### **QT prolongation**

Subjects with MVP have been found to have longer QT interval than control populations in some83 but not all reports.11, 84 Furthermore, QT prolongation was found to correlate with more severe leaflet prolapse and thickening of anterior leaflet85 and to be independently associated with VAs among MVP patients.14 An animal model simulating papillary muscle traction found alteration in the repolarization properties of the involved regions providing an explanation for the observed QT prolongation and arrhythmic potential.86 Importantly, while prolonged QT has been consistently shown to be associated with SCD in several large population studies,87, 88 to date, its link to mortality in subjects with MVP or MAD remains uncertain.

### **Fragmented QRS**

Fragmented QRS (fQRS) is defined as the presence of additional R waves (R') or notching in the R wave or the S wave, or the presence of >1 R' in 2 contiguous leads 89. It has been shown to mark localized scar89 and to be associated with increased risk of malignant arrhythmic mortality and SCD 90 in a variety of populations91-93. A recent study has showed that fQRS was associated with complex arrhythmia in patients with MVP (***Table 2:*** ***Ventricular ectopy – definitions)*** 94.Further research is needed to determine sensitivity and specificity of this feature in MVP patients as well as its role in clinical practice.

### **Premature ventricular contractions (PVCs)**

The observation of PVCs on repeated standard ECG tracings, usually indicates a significant burden of ventricular ectopy95. While the total burden of ventricular ectopy may not be inferred, captured tracings of complex arrhythmia is a marker of higher risk96 (***Table 2)***. PVC morphologies observed are often compatible with papillary muscle or mitral annular/LV basal origins. In many cases, a fascicular origin may be inferred, but these are by no means consistent or exclusive findings.14, 97 Ectopy arising from the outflow tracts are also common98. A unique pattern of non-sustained ventricular tachycardia (NSVTs) with morphologies alternating between papillary and right ventricular outflow tract has been described in a small cohort of MVP patients that survived out-of-hospital SCD.97 The prognostic significance of the various morphologies remains speculative.

### **Holter monitoring**

VAs on Holter monitoring are common in MVP patients, reported 43-85% of cases during 24h monitoring.14, 99 Most patients present with mild to moderate VAs, while severe VAs are rare (<10%) but associated with a 3-fold increased risk of mortality (***Table 3: Arrhythmia severity classification***).14We acknowledge the limitation of using specific cutoff values derived from a single center, albeit from a large cohort. While these data are consistent with previous studies from other patient populations showing increased mortality with higher burden of ectopy100, 101 and with NSVT102, 103, additional validation by other cohorts is needed.

A 24h Holter monitoring is superior to standard ECG as a screening modality in all patients with MVP (including asymptomatic) to assess arrhythmia burden and disease prognosis. In most asymptomatic MVP patients, detection of simple and infrequent (total burden<5% of single PVCs) VAs on 24h Holter monitoring may not warrant further evaluation. Symptomatic patients with mild VAs detected on 24h Holter monitoring would require a longer period of rhythm monitoring (∼7 days) to accurately detect overall PVC burden.95 However, patients with complex ventricular ectopy, short coupled PVCs (coupling interval <350 ms)104 or fast NSVT are at risk of SCD and warrant further investigation (***Figure 6***).11, 105 Unexplained presyncope or syncope with frequent or complex VAs (***Table 2***) on Holter monitoring should lead to strong consideration of long-term rhythm monitoring (e.g .repeated Holter monitoring, external or implantable loop recorders)106, especially if other phenotypic risk features are present.

### **Echocardiographic** **Predictors of Arrhythmias**

### **Pathophysiological Predictors of Arrhythmias**

MR quantification is essential for evaluating the risk of ventricular arrhythmias,11 as several studies have demonstrated excess mortality associated with increasing MR severity,12, 13 as well as a specific increase in the risk of SCD in patients with severe degenerative MR42. MR quantification is performed by Doppler-Echocardiography based on stroke volumes measurement by Doppler or by a combination of Doppler and LV volume measurements. The most commonly used method is however the flow convergence (Proximal Isovelocity Surface Area) method based on shifting of color baseline to define the radius of the flow convergence107. By these methods, the variables measured are the regurgitant volume and the effective regurgitant orifice area. The thresholds for severe degenerative MR are ≥60 mL/beat for regurgitant volume and ≥40 mm2 for effective regurgitant orifice area while mortality risk starts at 20-30 mm2 with a linear increase with larger values108, 109.

DMR with reduced LV function in itself is also associated with higher risk of SCD34 and excess-mortality110, 111 although its causal link to VAs has not been formally proven. LV ejection fraction <60% and end-systolic diameters ≥40mm are the accepted thresholds for reduced LV function and accepted indication for mitral valve surgery.20

### **Morphologic Predictors of Arrhythmias**

Morphologic features associated with arrhythmia consist of MAD11, 13, 37 and severe myxomatous degeneration, defined by thick and redundant leaflets with multi-segment bileaflet MVP. Their combined presence characterizes the typical phenotype of the AMVP complex14. Patients presenting with the AMVP complex are at excess risk of developing VAs, independent of MR severity.14 Importantly, bileaflet prolapse is quite prevalent among patients with MVP, and while small series have suggested an association with higher mortality112, 113 this was not confirmed in larger cohorts or population based studies.30 Of note, MAD in subjects with MVP is in itself linked with VAs during follow-up.40 However, MAD within the first 10 years of follow-up is not associated with excess mortality40 because arrhythmia occurrence is often delayed long after MVP and MAD diagnosis. SCD and excess mortality are even more delayed after detection of severe arrhythmias.14 Whether MAD depth is a predictor with more frequent arrhythmias is unclear as is the mechanism of MAD-associated arrhythmia, although myocardial fibrosis is hypothesized as causative.

### **Strain Predictors of Arrhythmias**

Patients presenting with the AMVP complex exhibit more frequent myxomatous disease (40 vs. 60%)14 with left ventricular mechanical dispersion.48 While the global longitudinal strain value can be preserved (slightly altered as compared to patients without arrhythmia), some degree of dispersion in longitudinal strain peaks may be observed.114 Post-systolic deformations are often observed with marked MAD and MVP (***Figure 7***) attributed to increased interaction between wall deformation and mitral valve motion. Longitudinal strain is useful in assessing interplay between mitral leaflets and myocardium, and in demonstrating changes in temporal pattern of myocardial deformation.114 It has been suggested that providing these relevant automatically measured parameters could identify higher risk of VAs. However, in light of the current scarcity of data, further research is need to verify the clinical utility and predictive value of this.

### **Cardiac magnetic resonance imaging**

Cardiac MRI can also identify prolapsed scallops using a stack of cine images and is ideally suited for risk stratification of VAs in patients with MVP, due to its unique ability to non-invasively identify focal myocardial fibrosis using late gadolinium enhancement (LGE). Several studies suggested an association between LGE at the mid-wall of the papillary muscles or patchy myocardial fibrosis (non-ischemic pattern) in the LV infero-basal region, and complex VAs.115-117 Diffuse LV fibrosis determined by T1 mapping is also associated with increased risk of complex VAs among patients with MVP (significant shorter post-contrast T1 time).117 Overall, fibrosis is common in MVP (28-37%)118, 119, usually located close to the annulus in the basal left ventricular wall including papillary muscles and inferior wall.39 Importantly, only LGE within the mitral apparatus (papillary muscles and peri-annular region) has a clear pathophysiological association with arrhythmia. The significance of LGE in other regions remains unclear in this context.

Any cardiac MRI standard protocol can identify MAD,49 but, complete assessment of MAD extent over the mitral annulus circumference warrants 6 LV long-axis cine sequences with 30° interslice rotation.48 Comprehensive MAD assessment should include careful description of the mitral valve, MAD severity, LV remodeling and fibrosis. Important mitral characteristics include bileaflet MVP, myxomatous MVP, mitral annulus enlargement in systole, prolapse depth, increased ratio of basal to midventricular wall thickness laterally, and presence of curling.37, 49 Assessment of MAD severity comprises extent around the mitral annulus and width of MAD. The link between MAD extent and VAs remains ill-defined.46, 49

### **Genetics**

The evidence for a potential role of systematic genetic analysis in these patients is weak. Some data exist for familial clustering of non-syndromic MVP with X-linked inheritance pattern as first reported in 1969120. Later studies reported autosomal dominant transmission suggested MVP as the most common Mendelian cardiovascular abnormality with various phenotypic expressions.121 Filamin A (FLNA) mutations have been described recently in a large family. The FLNA*-MVP* phenotype is characterized by both congenital and degenerative alterations of the MV apparatus with more severe disease in men due to the X-linked inheritance.27 A truncating FLNA variant has been linked to the AMVP proposing a proarrhythmic effect of this variant.122

MVP has been related to the Dachsous1 gene (DCHS1), a member of the cadherin superfamily.123 In clinical practice, genetic testing is recommended in patients with syndromic disease (Marfan and Loeys Dietz syndrome, etc.) due to the implications on follow up for aortic disease. Routine genetic testing in non-syndromic MVP is currently not justified.

### **Biomarkers**

Besides electrocardiographic and imaging modalities, there is no established circulating biomarker associated with the AMVP. Short, noncoding micro RNAs (miRNA)124 and proteomics have been employed to identify patients with MVP.125, 126 A recent study explored stretch-related and fibrosis-related circulating biomarkers in 72 patients with MVP/MAD.127 Circulating soluble suppression of tumourigenicity-2 (sST2) was associated with VAs, supporting the theory of stretch induced arrhythmia mechanisms. Although tissue TGF-β1 signaling contributes to fibrosis and matrix remodeling,128, 129 circulating TGF-β1 level was not related to VAs.

Currently, the assessment of biomarkers in these patients is not justified.

### **Pathological evidence**

MVP has been an underappreciated cause of SCD for a long time. MVP has been frequently overlooked in pathological examinations due to the absence of a uniform criteria and lack of awareness of its potential rule as a cause of SCD. This resulted in uncertainty regarding the true burden of MVP.130 Careful mitral valve structural analysis of SCD patients with MVP suggested an association with increased annular circumference, leaflet length and thickness, and presence/extent of endocardial fibrous plaques (friction lesions) on the LV myocardium.11, 131 Among victims of SCD with MVP, bileaflet MVP was present in 70% and endocardial fibrous plaques in the posterolateral wall in >50%. Moreover, histological analysis revealed elongated MAD and LV fibrosis in SCD cases with MVP versus normal control hearts.13 In particular, patchy fibrosis was localized at the level of PMs with adjacent free wall in all and of the inferobasal wall in 88% of young SCD victims 13.

## **Mechanism of arrhythmia**

The uneven distribution of major arrhythmias in MVP remains unexplained but may be linked to the anatomical substrates and triggers.

### **Substrate**

Histological evidence of myocardial substrate is consistent with patchy fibrosis between the mitral valve, papillary muscle (PM) and adjacent infero-basal and LV myocardium in MVP patients who sustained SCD.11, 13 These findings correlate with cardiac MRI LGE distribution in the papillary muscles and infero-basal LV wall. These regionalized fibrotic, molecular and cellular changes suggest a reactive response to increased mechanical traction from a prolapsing mitral valve.

The areas of patchy myocardial fibrosis in the sub-valvular apparatus leading to conduction delays serve as the substrate132.

### **Triggers**

Mechanical stretch of papillary muscles may also shorten action potential duration and decrease resting diastolic potential leading to stretch-activated early afterdepolarizations resulting in triggered activity.133 Furthermore, endocardial and mid-myocardial fibrotic changes on the papillary muscles and adjacent LV can create abnormal repolarization resulting in inverted T-waves frequently observed on the inferolateral leads on 12-lead ECG. These abnormalities in ventricular repolarization with QT prolongation, and ST-T changes may give rise to polymorphic VT.16, 84

Detailed invasive voltage mapping correlates PVC sites of origin with the papillary muscles, fascicular system, LV outflow tract and mitral annulus with predominant right bundle-branch block morphology (***Figure 8***). Subtle PVC morphology variations are highly suggestive of multiple exits from a single source.134

Ventricular ectopy arising from Purkinje tissue may trigger ventricular fibrillation in patients with bileaflet MVP and SCD (8,9). Invasive voltage mapping may discover fractionated, split and delayed Purkinje potentials suggesting the presence of abnormal tissue. The Purkinje system is a critical component in arrhythmogenesis and risk of SCD.98, 135 It has been suggested that mechanical snap of redundant leaflets on ventricular myocardium sets this cascade into motion,81 but this concept is not measurable and its link to the heterogeneous occurrence of VAs and SCD remains putative.

## **Risk stratification**

## **Recommended screening**

Risk stratification aiming at assessing the risk of VAs and SCD is challenging even for the most common and well-studied conditions.136, 137 Approaches for screening for AMVP are limited by the lack of prospective outcome data and absent recommendations for rhythm monitoring in current valvular guidelines.7, 20

In this novel endeavor, the present expert group was guided by 3 pillars: a) all conventional indications for ICDs based on current guidelines apply to MVP,106, 138, 139 b) AMVP, particularly in the presence of LGE on MRI should not be regarded as a “structurally normal heart” and any VAs complicating MVP should not be considered idiopathic or benign, c) several clinical, electrocardiographic, echocardiographic and other imaging parameters (detailed in section 6.1) have been shown to be associated with increased risk of death and/or VA in AMVP patients and should be screened for in patients with MVP.

All patients with MVP should be stratified for SCD risk (*Green Heart,* ***Table 4: Risk stratification***). Risk stratification of patients with MVP includes focused history, 12-lead ECG, extended ECG monitoring and detailed echocardiography (*Green heart,* ***see recommendation tables***). The use of Cardiac MRI and implantation of a loop recorder are more selective depending on the probability of ventricular arrhythmia. Risk stratification involves two phases based on the clinical and imaging context and the detected arrhythmia (***Central Illustration: Risk stratification***).

Clinical and Imaging context

In patients with MVP, the arrhythmia’s risk is not uniform and is higher in certain contexts. While patients who have recovered from sudden cardiac arrest have *overt* severe arrhythmia, the other patients are at risk of serious arrhythmias. The risk is higher when they present with unexplained syncope or even presyncope than when they present with isolated palpitations and lower when asymptomatic. Similarly features such as negative T waves, severe myxomatous degeneration with redundant leaflets, MAD and generally bileaflet prolapse, or LGE on CMR represent a higher risk of VAs.

Therefore, the clinical and imaging context strongly influences the intensity of the search for the presence of serious VAs. For example, patients presenting with unexplained syncope, even without VT on Holter monitoring are prime candidates to undergo extended ECG monitoring by longer ECG monitoring or ILR, if necessary, to uncover with the highest sensitivity the existence of VAs. Conversely, asymptomatic patients without complex ventricular arrhythmia on the initial screening Holter would only require episodic Holter assessment, more frequently if they present with phenotypic risk features. Because VAs may develop secondarily, a plan of reassessment over time is necessary, the frequency and intensity of which depends on the presence and number of phenotypic risk features. The presence of multiple phenotypic risk features suggests a higher arrhythmic risk and justifies more intense monitoring.

Observed arrhythmia

Few studies have addressed the potential for SCD or even for excess-mortality based on the arrhythmia detected, prospectively or retrospectively and the grading of arrhythmia below is mostly based on extension of the general grading of VAs.

**High-risk**: The committee considered the following VAs as high-risk, detected at rest, by ECG, Holter or ILR. Whether their detection during exercise carries a similar high-risk is probable but uncertain.

-Sustained VT not originating from the right or left ventricular outflow tract (Box 3).

-Spontaneous polymorphic NSVT.

-Rapid NSVT monomorphic (>180 bpm) has been associated with subsequent excess-mortality.

**Intermediate risk**: This category is even less well defined. The following VAs were considered by the committee as probable intermediate risk:

-Polymorphic PVCs.

-NSVT monomorphic, of lower rate (<180 bpm).

-Highly frequent or complex PVCs.

**Low risk**: Patients with frequent PVCs but not complex VAs (and no morphological higher risk features) are considered low risk.

The frequency of repetitive evaluation for risk reclassification is not well defined but regular Holter monitoring may prove helpful for this purpose.

In the absence of structural cardiac abnormalities, outflow-tract VAs are considered benign. Their clinical significance in patients with AMVP remains unclear but concerns have been raised regarding a possible mechanism where outflow-tract ectopy triggers a more malignant arrhythmia arising from the abnormal tissue in papillary muscle.81, 135

**Box 3- localization of origin of ventricular ectopy**

The likely origin of ventricular arrhythmias may be ascertained with accuracy from the QRS morphology on a standard ECG 140-142. Several sets of criteria and algorithms, focusing on the mitral annulus and the papillary muscles have been published and validated141, 143-145. ***Figure 8*** summarizes the most relevant ECG characteristics of ventricular arrhythmias arising from the mitral annulus or the papillary muscles and shows typical examples of PVCs originating from these structures.

A likely mitral apparatus origin (papillary muscles, and mitral annular region) may even be suspected based on a 3-lead Holter provided that those include 2 precordial leads (usually V1/2 and V5/6) and an inferior lead (usually aVF) thus allowing the estimation of the maximal QRS width, axis, precordial transition and type of bundle branch like morphology. Nevertheless, 12-lead Holter monitoring is more accurate for ventricular arrhythmia localization and should be preferentially used when available.

The likely origin of ventricular arrhythmia may also be inferred by the distribution of LGE in CMR.

## **Clinical evaluation**

A comprehensive clinical evaluation is mandatory to identify MVP patients with higher risk of arrhythmias. Family history of SCD is relevant to suggest the possibility of inherited arrhythmia syndromes (long-QT syndrome, arrhythmogenic right ventricular cardiomyopathy) as alternative diagnoses. Family history of MVP should call attention to heritability, as either sporadic or familial forms.23, 27 While autopsy studies have focused on patients without any other cause of SCD (young, often female) no demographic characteristic (age or sex) is particularly associated with SCD.30, 59 However, with the current state of knowledge it is not yet possible to affirm whether a particular sex or age range is associated with excess risk of sudden death compared with that occurring in the community. Patients may be completely asymptomatic or experience mild nonspecific symptoms (atypical chest pain, anxiety). Common symptoms associated with AMVP are palpitations, chest pain and dyspnea.37 However, presentation with syncope or presyncope may be a harbinger of SCD and warrants prompt thorough investigation.37 History of ventricular tachycardia (VT) or ventricular fibrillation (VF) should be retrieved from all available ECG tracings. The typical MVP auscultatory feature is an apical high-pitched, late systolic murmur with or without mid-systolic click,146, 147 but does not allow inference of SCD risk. Holosystolic apical murmurs generally suggest more significant MR. Although no isolated clinical characteristic is independently predictive of malignant arrhythmias or SCD, clinical presentation with MVP and unexplained syncope, or palpitations should alert the clinician to further risk stratification. (***Table 5 – Clinical evaluation***).

## **ECG monitoring**

## **Holter monitoring**

Holter monitoring is a crucial tool to establish the diagnosis of AMVP even in the absence of palpitations. The yield of Holter monitoring depending on symptomatic versus asymptomatic status remains to be defined. Frequent and sometimes polymorphic PVCs may be detected.48 Less frequently, complex VAs may be detected and when ‘severe’ (NSVT>180 bpm or more severe VAs) are associated with subsequent excess mortality.14 Data regarding the prognostic significance of PVC is not as robust67, yet several studies demonstrated an increased risk of mortality14, 100 and heart failure101, 148.

In addition, Holter monitoring provides information on the total burden of ventricular ectopy, its morphology, coupling interval and may capture complex ectopy149. The clinical importance of these markers remain to be established. (***Table 6 – Extended ECG monitoring***).

## **Implantable loop recorder (ILR)**

The short duration of a Holter monitor favors implantable loop recorders (ILRs) as an option to improve arrhythmia detection and risk stratification in patients with AMVP.106, 150, 151 While there are no systematic studies on ILR utility in patients with MVP (only case reports), 152, 153 it appears to be an useful option to improve sensitivity in detecting AMVP. Thus, ILR use appears reasonable in patients with high risk features such as unexplained syncope with inconclusive Holter monitoring, or intermediate risk cases such as patients with multiple phenotypical risk features including positive LGE on CMR. (***Table 6 – Extended ECG monitoring***).

## **Echocardiographic protocol**

In addition to screening echocardiography, comprehensive Doppler-echocardiography following guideline-based standard imaging protocol is required.154

Mitral morphology should be analyzed and MVP phenotype noted (myxomatous MVP or Fibroelastic Deficiency with/without flail leaflet) as well as which leaflet(s) is involved.

Regarding LV, accounting for the volume load of the prolapsing volume (e.g. the volume above the LV muscle but below the prolapsing mitral leaflets in the ventricularized portion of the left atrium) reconciles the disproportionate LV enlargement that is noted in some MVP patients.155 In the presence of MAD, the prolapsing volume should be measured between the mitral annulus and the prolapsing leaflets in end-systole.

The protocol should include:

* Degenerative MR integrative grading using specific, supportive and quantitative measures to classify degenerative MR from absent to severe according to guidelines.6, 7
* Leaflets length and thickness measurements, performed in the parasternal long axis view, in mid-diastole. Leaflet-redundancy can be assessed using M-mode over the mitral valve in the parasternal short-axis view and can be graded by evaluating excess valve tissue thickening, according to guidelines25, 156. The use of 3D and global longitudinal strain are of added value to better characterize the valve and the consequences at the level of the LV. Additionally, mechanical dispersion based on longitudinal strain in the apical 4C view has shown potential added value to predict the risk of VAs in MVP patients.
* MAD length, measured in parasternal long-axis view at end-systole and defined as the distance between mitral-annulus and systolic bulge of the ventricular myocardium. A range of 5-10 mm has been measured in long axis views.46 MAD can be observed at different locations on the mitral annulus. However, MAD is associated with increased risk of VAs when observed at the posterior LV wall.
* Additionally, distinctive spike during mid-systole to late-systole of the lateral mitral annulus using Doppler has been associated with MAD. 157 (***Table 7 – Detailed Echocardiography***).

## **CMR protocol**

CMR can refine the echocardiographic findings and provide additional unique information on myocardial structure, both important to improve risk-stratification of AMVP patients *118, 119*. Particularly, CMR should be performed in all patients who survived a SCD or experienced sustained VAs before implanting an ICD.158 CMR has been shown to increase the rate of etiological diagnosis in patients with resuscitated SCD or sustained VT159. Conversely, ruling out alternative etiologies will increase the certainty in the diagnosis of AMVP. Due to associated risk of SCD, we also suggest performing a CMR in patients with a history of unexplained syncope or documented NSVT14. It may also be useful in patients with AMVP and at least one phenotypical risk feature.

Furthermore, performing CMR in patients in whom echocardiography does not provide accurate assessment of left and right ventricular function, of structural changes or of mitral characteristics may be of value.158 CMR allows assessment of the presence and severity/extent of MAD or LV dilatation and dysfunction. Most importantly, CMR identifies myocardial fibrosis/scar, considered strongly associated with severe VAs.

CMR should be performed in centers with sufficient expertise and on a 1.5 or 3T scanner. The MRI protocol should be comprehensive, including the following morphofunctional parameters according to definitions previously mentioned: 13, 48, 49, 64, 66, 67, 119, 160 2-8

* Assessment of LV volumes and function (ejection fraction and regional wall motion abnormalities); measurement of LV mass and end-diastolic thickness, including the ratio of basal-to-midventricular wall thickness.
* Assessment of left atrium and right ventricular size and function.
* Identification of MAD (presence or absence) and its quantification, as longitudinal length at least from the long axis view, and possibly as circumferential extent expressed in degrees.
* Identification of the systolic curling (presence or absence) and possibly its quantification.
* Measurement of mitral valve annulus diameter (at end-systole and end-diastole in both anteroposterior and inter-commissural aspects), leaflet diastolic thickness, leaflet length and prolapsed distance.
* Assessment of myocardial fibrosis/scar with description of the localization (particularly at the level of the papillary muscles or adjacent myocardium, and next to the mitral valve annulus) and its quantification.
* In case of doubt from echocardiography on the severity of mitral regurgitation, regurgitant volume and fraction are calculated using the indirect calculation of the regurgitant volume. Such is obtained from the difference between the total LV volume (using planimetry of short-axis cine images) and the forward stroke volume across the aortic valve (using phase-contrast velocity mapping).
* In centers with the expertise, assessment of interstitial/reactive fibrosis by T1 mapping.

(***Table 8 – Evaluation by CMR***).

## **Electrophysiological study**

As there are no robust data specific to this population, we consider the standard programmed stimulation protocol that includes up to 3 extra-stimuli protocol from at least 2 sites (RV apex and RVOT) down to ERP or 200 ms. Induction of polymorphic VT and VF by delivery of 3 extra-stimuli is a nonspecific finding and does not predict future risk of SCD.161-163 However, induction of monomorphic VT may be more specific and linked to MVP if localized to the mitral apparatus. A recent systematic review reported on the outcome of EPS with programmed ventricular stimulation (PVS) in 22 patients with MVP and documented SCD.59 The findings were disappointing with sustained monomorphic VT (5%), NSVT (23%), VF (18%), and non-inducibility of VAs (55%). The fact that sustained monomorphic VT is only rarely induced may imply, that in most cases, the arrhythmia leading to SCD is non-reentrant. Thus, the role of PVS in identifying high-risk MVP is limited. This scarcity of data does not provide conclusive evidence regarding positive or negative predictive value of the test. While the committee does not endorse the use of EPS with PVS, if EPS is used, the induction of sustained monomorphic VT should be considered more specific than polymorphic VT or VF.

## **Exercise stress testing**

Exercise stress testing may be an important tool to assess suspected exercise-induced VAs.158 Although there are no data on the significance of these arrhythmias in patients with AMVP, several reports have established exercise-induced VAs as a predictor of cardiovascular mortality even in asymptomatic patients164, 165. Furthermore, exercise stress testing may be useful in assessing exercise tolerance, particularly in AMVP patients who want to engage in sports166. The value of exercise stress testing for the identification of obstructive coronary artery disease is limited compared to other modalities167 and may be even less reliable in the presence of repolarization abnormalities at baseline. Nevertheless it may be reasonably included, alongside cardiovascular risk factors, in the preliminary assessment of the likelihood of coronary artery disease in patients with AMVP. (***Table 9 – Exercise stress testing***).

## **Management**

## **Medical therapy**

Current literature regarding medical management of AMVP patients is sparse. However, some general concepts may be inferred from associated better studied case scenarios. The medical management should be tailored to the specific presentation and aimed at symptomatic relief and improved survival.

## **Multiple PVCs**

Presence and burden of ventricular ectopy is associated with increased mortality100 even in the absence of structural heart disease.101 However there is no evidence that prophylactic treatment aimed at suppressing PVCs in asymptomatic patients is beneficial.

While impact on long term outcomes remains unclear,168 suspected PVC-induced cardiomyopathy warrants treatment aiming at reducing PVC frequency. Treatment is also indicated in symptomatic patients regardless of LV function.169 Beta blockers and verapamil improve symptoms and result in modest reduction of PVC burden 170, 171. Flecainide, Propafenone and Amiodarone yield more potent PVC burden reduction with frequent improvement in LV function.172-174The potential benefit from amiodarone must be balanced against risk of significant adverse effects associated with long term treatment. Sotalol can reduce the PVC burden effectively but failed to improve LV function 175.

## **PVC induced Polymorphic VT/VF**

Cases of patients with PVC-induced VF have been described,134, 176, 177 with triggers arising from Purkinje fibers within papillary muscles or the fascicles. A few cases have been described as drug refractory176, including in patients with MVP.37, 178 Quinidine may have an important role in the treatment of short-coupled PVCs triggering polymorphic VTs due its effect on the Purkinje system although this was not tested prospectively nor specifically in patients with MVP.179

## **Prevention of SCD**

Despite the link between PVC and mortality, there is no evidence that medical therapy alone may lower the risk of SCD.173 Therefore, medical treatment should focus on the improvement of symptoms. The need for an ICD should be assessed in all cases. (***Table 10 – Medical Therapy***).

## **Implantable cardioverter defibrillator**

Established primary and secondary prevention ICD indications also apply to patients with MVP.

Primary prevention ICD is indicated in symptomatic heart failure with EF≤ 35% despite three-months of optimal medical therapy.20 It is uncertain whether patients with MVP/DMR fulfill this type of indication as EF in that range is rarely observed in this context.111

Secondary prevention ICD is indicated in patients with MVP and documented history of sudden cardiac arrest with ventricular fibrillation (VF) or VT without reversible causes.138

Primary prevention implantation of ICD is more complex as no randomized trial has demonstrated benefit in any MVP subset. In the absence of conclusive data, the present committee concludes that ICD should be a strong consideration in patients presenting with unexplained syncope and high-risk VAs detected by ECG, Holter or ILR and possibly by exercise testing.

In the absence of any published data specific to AMVP, the choice of type of defibrillator (subcutaneous vs. trans-venous) should follow the same considerations used in other arrhythmic etiologies 138. A careful evaluation of the presenting arrhythmia, likelihood of response to anti-tachycardia pacing or a need for bradycardia pacing are warranted. The potential risk of inappropriate shocks in the specific patients should be balanced against the risk of endovascular infection and lead failure. (***Table 11 – SCD prevention***).

## **PVC/VT ablation**

VAs in MVP most commonly manifest as focal PVCs but may also involve PVC-triggered VT/VF or scar-related reentrant VT*.* All of these can be effectively treated by catheter ablation.98, 134, 178, 180-183 Indication for ablation of VAs is independent of MVP phenotype and may include PVC ablation in patients with symptoms refractory to ICD therapy,158, 184 to drug treatment, when drug treatment is not tolerated, or when ablation is preferred over long-term drug therapy by patients.

In addition, ablation of PVC/VT may be recommended in patients with PVC-induced cardiomyopathy, PVC-induced VF or in patients with frequent ICD therapies.158, 184 Ablation at the papillary muscle should be performed with caution as this may rarely lead to LV dysfunction and may impede mitral valve function..185

PVCs in MVP are usually related to the papillary muscles or the Purkinje system,98, 176 but ablation of foci within the papillary muscles is challenging, often requiring intracardiac echocardiography, contact force sensing technology and cryoablation to improve catheter contact and effective energy delivery.181, 182 Scar-related monomorphic reentrant VTs are uncommon and typically located in the inferobasal or inferolateral left ventricle98, 132.

Long-term success rates for PVC ablation in MVP range between 60 and 84%.98, 134, 176, 180, 181 The outcome of ablation does not appear to be adversely impacted by presence of MVP.178 Inducibility of sustained VAs and multifocal ectopy may indicate higher risk of VA recurrence during follow-up and may relate to a more diffuse myopathic process (higher degree of LGE on MRI).98

Systolic function and MR severity may improve after effective PVC elimination.134 Ablation of VAs in MVP patients should only be performed in centers with appropriate experience in ablation of left-sided VAs and in interventional and surgical treatment of mitral valve disease. (***Table 12 – Catheter ablation of VAs in AMVP***).

## **Role of mitral valve surgery**

Surgical intervention for AMVP remains controversial and limited to small case series and reports.186-193 While surgical correction of severe MR tends to reduce VA burden in patients with MVP and degenerative MR,14,194 results are inconsistent.176, 195 The respective role of various surgical procedures (repair/replacement, valve tissue resection etc.) remains undefined. Recently surgical cryoablation of VAs during mitral valve surgery has been reported,196, 197 but the impact on medium to long-term outcomes remains to be defined. There is uncertainty whether even with successful mitral surgery for the treatment of severe MR in patients with AMVP and high-risk VAs, that it is sufficient to prevent further arrhythmia. The need for an ICD may remain. Surgical indications in patients with MR in the moderate range are yet to be determined. Early surgery in patients with MVP and severe MR without VAs restores life-expectancy, by eliminating excess mortality. Thus, mitral surgery, through suppressing the progression of valve prolapse and its consequences, may have a role in preventing SCD, giving credence to the mechanical theory for VAs in MVP.

Well-designed clinical trials are warranted to better define the role of surgery in the treatment of AMVP. (***Table 13 – MV surgery in AMVP***).

## **Evidence Gaps**

The association between MVP and SCD has long been debated, suggested by case reports but questioned based on the generally good prognosis of MVP without significant DMR or LV dysfunction. Only recently has the entity of AMVP been better described and its presence recognized based on a prominent phenotype and the increased incidence of VAs detected in patients with the AMVP phenotype. However, while we are starting to modify our clinical algorithms for detection and treatment of AMVP based on current knowledge, much more needs to be learned:

1-The incidences and mechanisms of the various VAs in patients with MVP in general and in various subsets remains ill defined. The endeavor of defining incidence rates of VAs and SCD will require large samples with systematic rhythm monitoring using both Holter and ILR detection of VA and long-term follow-up. While prospective studies may offer definite conclusions, a more pragmatic approach is needed in order to produce timely results. Therefore, it is crucial that a consortia sharing retrospective data be formed in order to provide initial estimates that may guide management in the current patient population.

2-Determinants of VAs and SCD are only partially defined. This warrants the formation of large well designed study cohorts that will collect and assess clinical characteristics, imaging data and other biological features such as proteomics, genomics and metabolomics as potential predictors. Here again both prospective cohorts and retrospective consortia are indispensable to resolve this conundrum.

3-Progression of VAs in parallel with progression of MVP, DMR and their consequences is unknown and will require cohorts with repeat rhythm monitoring.

4- Intense arrhythmia detention may potentially lead to excess, unnecessary and even harmful interventions. This risk should not be underestimated, mandating a careful design of clinical trials involving patients with MVP.

5-Clinical outcome associated with VAs is based on relatively limited cohorts and the refined prognostic implications of various VAs should be assessed in large cohorts with long-term follow-up to redefine high, medium and low risk subsets.

6-Benefits of therapeutic interventions in MVP with VAs can only be affirmed in clinical trials. These may clarify the potential benefit of ICDs in patients with high-risk VAs, of mitral surgery in MVP with DMR and VAs, of surgical cryoablation versus transcatheter ablation and of antiarrhythmic drugs versus ablation for frequent PVCs, among the many therapeutic considerations.

Therefore, one of the main goals of this document is to highlight the important and more urgent questions that require our collective attention. By suggesting clearer diagnostic criteria, we aim to enable standardized approaches to patient inclusion in research protocols and pave the way for multicenter international collaborations that are required to tackle these clinical conundrums.

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**Recommendation Tables**

***Table 1: Categories of consensus statements***

|  |  |  |
| --- | --- | --- |
| **Consensus statement** | **Definition** | **Symbol** |
| Recommended/indicated or ‘should do this’ | Scientific evidence that a treatment or procedure is beneficial and effective or, is strongly supported by authors’ consensus. |  |
| May be used or recommended | General agreement and/or scientific evidence favor the usefulness/efficacy of a treatment or procedure |  |
| Should NOT be used and is NOT recommended | Scientific evidence or general agreement not to use or recommend a treatment of procedure |  |

***Table 2:*** ***Ventricular ectopy – definitions***

|  |  |
| --- | --- |
| **Ectopy Type** | **Definition** |
| Polymorphic Premature ventricular contractions (PVCs) | 3 or more distinct PVC morphologies (not slight variations of the same morphology) suggestive of multiple origins rather than multiple exits of a single site |
| Short coupled PVC | A PVC with coupling interval less than 350 ms198 |
| Non-sustained Ventricular Tachycardia (NSVT) | Three or more consecutive ventricular beats at a rate of ≥100 beats per minute lasting 30 seconds or less |
| Monomorphic NSVT | NSVT with a single QRS morphology |
| Polymorphic NSVT | NSVT with a constant beat to beat change in QRS morphology |
| Sustained Ventricular Tachycardia | VT lasting more the 30 seconds or requiring immediate termination by external means |
| Frequent ventricular ectopy | Premature ventricular beat constituting at least 5% of the total complexes |
| Complex ventricular ectopy | NSVT, VT and VF |

***Table 3: Arrhythmia severity classification***

|  |  |  |  |
| --- | --- | --- | --- |
| **Severity** | **Arrhythmia burden/rate** | **Risk of Mortality HR [95% CI]** | **Ref** |
| ***Mild*** ventricular arrhythmia | PVC≥5% and/or  VT runs <120 bpm | 1.20 [0.68-2.14], *P*=0.5 | 14 |
| ***Moderate*** ventricular arrhythmia | VT runs 120-179 bpm |
| ***Severe*** ventricular arrhythmia | VT runs ≥180bpm and/or history of sustained VT/VF | 2.94 [1.36-6.36] *P*=0.006 |

***Table 4: Risk stratification***

|  |  |  |
| --- | --- | --- |
| **Consensus statement on Risk stratification** | **Symbol** | **Ref** |
| Patients with MVP should undergo a directed and structured risk stratification process aimed at identifying AMVP and assessing the risk of SCD. |  | Expert  consensus |

***Table 5: Clinical evaluation***

|  |  |  |
| --- | --- | --- |
| **Consensus statement on Clinical Evaluation** | **Symbol** | **Ref** |
| A careful clinical evaluation, including family history of SCD, previous syncope, comorbidities and physical examination, should be performed in patients with MVP. |  | Expert  consensus |
| Due to possibility of disease progression, clinical re-evaluation should be considered periodically or when clinical circumstances have changed (e.g. occurrence of syncope or palpitations). |  | Expert  consensus |

***Table 6: Extended ECG monitoring***

|  |  |  |
| --- | --- | --- |
| **Consensus statement on Extended ECG monitoring** | **Symbol** | **Ref** |
| A standard 24 hour Holter monitoring is warranted in all patients with MVP. |  | 48, 149 |
| Longer ECG recording (up to 7 days) may be beneficial in selected cases to allow more accurate quantification of PVC burden and/or correlation with symptoms. |  | 95 |
| MVP patients with complex ventricular ectopy (e.g. fast NSVTs) should be followed closely. |  | Expert  consensus |
| Due to the possibility of disease progression, periodic Holter may be considered as part of the routine follow up of AMVP! Patients. |  | Expert  consensus |
| Periodic Holter monitoring of PVC burden and periodic echocardiographic evaluation of LV function may be particularly helpful in patients with frequent PVC, even when asymptomatic and with normal LV function (to allow for a timely diagnosis of PVC induced cardiomyopathy). |  | Expert  Consensus  168, 199 |
| Longer ECG recording may be considered in patients with MVP and doubtful symptoms in whom 24 hour Holter monitoring was not revealing. |  | Expert  consensus |
| ILR may be considered in patients with MVP and unexplained syncope in whom non-invasive ECG monitoring was not revealing or inconclusive. |  | 106, 150 |
| ILR may be considered in patients with AMVP, high risk features& and negative CMR. |  | Expert  consensus |
| ILR may be considered in patients with AMVP, at least 1 phenotypic risk feature# and positive LGE on CMR@. |  | Expert  consensus |

*!AMVP- The presence of MVP with or without MAD, frequent ventricular ectopy (≥5% of total beats), complex ectopy or sustained VAs in the absence of any other well-defined arrhythmic substrate (e.g. active ischemia, ventricular scar due to another defined ethology, primary cardiomyopathy or channelopathy; ventricular scar due to other defined etiology refers to ischemic cardiomyopathy, DCM, post myocarditis, cardiac sarcoidosis etc.)*

*& High risk features – sustained VT (hemodynamically tolerated), NSVT, unexplained syncope*

*# phenotypic risk features - palpitations, T-wave inversion in the inferior leads, repetitive documented polymorphic PVCs, mitral annular disjunction (MAD) phenotype, redundant MV leaflets, enlarged left atrium or ejection fraction ≤ 50%.*

@*For this purpose, only LGE within the mitral apparatus (papillary muscles and peri-annular region) has a clear pathophysiological relevance. The significance of LGE in other regions remains unclear in this context.*

***Table 7: Detailed Echocardiography***

|  |  |  |
| --- | --- | --- |
| **Consensus statement on Detailed Echocardiography** | **Symbol** | **Ref** |
| In patients with suspected AMVP, comprehensive echocardiography assessment should include evaluation of leaflet length and thickness measurement, annular dimension, MAD characterization, degenerative MR grading and possibly advanced assessment# of LV function. |  | 14 |
| Due to the possibility of disease progression, periodic complete transthoracic echocardiography may be considered as part of the routine follow up of AMVP! patients. |  | Expert  Consensus |
| In the work up of patients with frequent PVCs, syncope or aborted cardiac arrest with no other obvious etiology, the comprehensive echocardiographic study should include the careful assessment of the mitral valve and the mitral annulus to diagnose AMVP. |  | Expert  Consensus |

*#Advanced” assessment of LV function may be based on the combination of Simpson’s bi-plane methods, assessed by 3D echo and Global Longitudinal Strain by speckle tracking imaging.*

!*AMVP - The presence of MVP with or without MAD, frequent ventricular ectopy (≥5% of total beats), complex ectopy or sustained VAs in the absence of any other well-defined arrhythmic substrate (e.g. active ischemia, ventricular scar due to another defined ethology, primary cardiomyopathy or channelopathy).*

***Table 8: Evaluation by CMR***

|  |  |  |
| --- | --- | --- |
| **Consensus statement on CMR** | **Symbol** | **Ref** |
| CMR should be performed in all AMVP patients who survived a cardiac arrest or experienced sustained VA, and before$ implanting an ICD for secondary prevention. |  | 159, 200 |
| CMR should be performed in all patients when echocardiography does not provide accurate assessment of LV and RV function and/or evaluation of structural changes. |  | Expert  consensus |
| CMR should be performed in all MVP patients with a history of unexplained syncope and/or NSVT. |  | Expert  consensus |
| CMR should include assessment of LV size and function, assessment of MR severity, leaflet length/thickness measurement, MAD characterization and curling, and LGE assessment. |  | Expert  consensus |
| CMR may be useful in patients with AMVP and at least 1 phenotypic risk feature#. |  |  |

*$CMR should not unduly delay the implantation of a defibrillator.*

*# phenotypic risk features - palpitations, T-wave inversion in the inferior leads, repetitive documented polymorphic PVCs, mitral annular disjunction (MAD) phenotype, redundant MV leaflets, enlarged left atrium or ejection fraction ≤ 50%.*

***Table 9: Exercise stress testing***

|  |  |  |
| --- | --- | --- |
| **Consensus statement on Exercise stress testing** | **Symbol** | **Ref** |
| In patients with AMVP, exercise stress testing should be used to assess for adrenergic-dependent rhythm disturbances. |  | Expert  consensus |
| The use of exercise stress testing to assess exercise tolerance in AMVP patients who want to engage in sports  is reasonable. |  | 166 |
| Exercise stress testing (plus echocardiography), alongside other risk factors may be used to assess the clinical likelihood of obstructive coronary artery disease |  | Expert  consensus |

***Table 10: Medical Therapy***

|  |  |  |
| --- | --- | --- |
| **Consensus statement on Medical Therapy** | **Symbol** | **Ref** |
| AMVP with symptomatic severe MR not eligible for surgery should be treated with optimal HF medications. |  | 139 |
| Suspected PVC induced cardiomyopathy in the presence of MVP should be treated. Reasonable options include beta blockers, sotalol and amiodarone. |  | 139, 170, 171, 173, 175 |

HF-heart failure

***Table 11: SCD prevention***

|  |  |  |
| --- | --- | --- |
| **Consensus statement on SCD Prevention** | **Symbol** | **Ref** |
| ***Secondary prevention*** |  |  |
| Patients with AMVP and a documented history of VF or hemodynamically not tolerant VT, in the absence of reversible causes should receive an ICD. |  | 139, 158 |
| ***Primary prevention*** |  |  |
| MVP with LVEF <35% and symptomatic HF despite ≥3 months of OMT should receive an ICD. |  | 139 |
| In patients with AMVP, history of unexplained syncope and sustained ventricular tachycardia, likely arising from the mitral apparatus, the implantation of a defibrillator is reasonable. |  | Expert  Consensus |
| In patients with AMVP, history of unexplained syncope and NSVT, likely arising from the mitral apparatus, an implantation of a defibrillator may be reasonable. |  | Expert  Consensus |
| In patients with AMVP, 1 high risk feature& and 2 or more phenotypic risk features#, the option of an ICD maybe be reasonable. |  | Expert  Consensus |

*& High risk features – sustained VT (hemodynamically tolerated), NSVT, unexplained syncope*

***#*** *phenotypic risk features - palpitations, T-wave inversion in the inferior leads, repetitive documented polymorphic PVCs, mitral annular disjunction (MAD) phenotype, redundant MV leaflets, enlarged left atrium or ejection fraction ≤ 50%, LGE. For this purpose, only LGE within the mitral apparatus (papillary muscles and peri-annular region) has a clear pathophysiological relevance. The significance of LGE in other regions remains unclear in this context.*

OMT – optimal medical therapy

***Table 12: Catheter ablation of VAs in AMVP***

|  |  |  |
| --- | --- | --- |
| **Consensus statement on catheter ablation of VAs** | **Symbol** | **References** |
| Ablation of PVCs in patients with frequent PVCs who are symptomatic or have decreased LV function is recommended. |  | 98, 134, 158, 178, 180-184 |
| Ablation of VA in MVP patients should be performed in experienced centers with expertise in VA ablation and interventional and surgical treatment of MV regurgitation. |  | Expert  consensus |
| Ablation of papillary muscle PVCs/VA is challenging and use of intracardiac echocardiography, contact force sensing catheters or cryoablation may be helpful to improve catheter contact and effective manipulation. |  | 181, 182 |
| Ablation of PVCs is reasonable if triggering VF, particularly if not controlled by medications. |  | 134, 135 |
| Ablation of sustained monomorphic VT despite antiarrhythmic treatment or if antiarrhythmic treatment is not desired, or contraindicated should be performed in MVP patients with recurrent ICD therapies. |  | 158, 184 |

***Table 13: MV surgery in AMVP***

|  |  |  |
| --- | --- | --- |
| **Consensus statement on mitral surgery** | **Symbol** | **Ref** |
| Mitral valve surgery may reduce the burden of malignant VAs# in MVP patients and severe MR. |  | Expert  consensus |
| Surgical cryoablation during mitral valve surgery with history of complex VA may be considered. |  | Expert  Consensus |

# *Malignant VA – sustained VA, highly symptomatic NSVT.*

1. **Figure legends**

**Central Illustration: Risk stratification scheme.**

Risk stratification aiming at assessing the risk of VAs and SCD in patients with MVP, involving two phases based on the clinical and imaging context and the uncovered arrhythmia. In the absence of ventricular tachycardia, phenotypic risk features will trigger the intensity of screening for arrhythmia. Green boxes indicate green heart recommendations and yellow boxes indicate yellow heart recommendations.

High risk - sustained VT, polymorphic NSVT, fast (>180 bpm) NSVT, VT/NSVT resulting in syncope.

ICD = implantable cardioverter defibrillator; LA = left atrium; LGE = late gadolinium enhancement; LV-EF = left ventricular ejection fraction; MAD = mitral annular disjunction; MV = mitral valve; PVCs= premature ventricular contractions; TWI = T-wave inversion; VT = ventricular tachycardia.

**Figure 1: The mitral valve with and without mitral valve prolapse.**

Transthoracic echocardiographic long-axis view in end-systole zoomed in on the mitral valve with (A) the location of both mitral leaflets within <2mm under the plane of the mitral annulus (red line) refutes the diagnosis of mitral valve prolapse. (B) A displacement of both leaflets >2mm (red arrow) above the plane of the annulus (red line) defines mitral valve prolapse. Note the absence of detachment of the posterior leaflet from the left ventricular myocardium (no mitral annular disjunction).

**Figure 2: Mitral valve prolapse with and without MAD on TTE.**

Transthoracic echocardiographic long-axis view in end-systole displaying bileaflet mitral valve prolapse with (A) MAD (yellow line) of 11mm length vs. (B) without MAD. The red line indicates the plane of the mitral annulus.MAD = mitral annular disjunction; TTE= Transthoracic echocardiography.

**Figure 3: Mitral valve prolapse and MAD on TTE and TEE.**

(A) Transesophageal echocardiographic 3D view, displaying a large prolapse of P2 precisely assessed by “photo-realistic” tools now available. Severe mitral regurgitation is associated as shown on 120° TEE (B) and TTE apical 4-chamber views, with severe left atrial dilatation. A 10mm MAD is diagnosed both on these TEE (D) and TTE (E) views.

MAD = mitral annular disjunction; TEE = Transoesophageal echocardiographic; TTE = Transthoracic echocardiography.

**Figure 4: Mitral valve prolapse and MAD on Cardiac-MRI.**

CMR long-axis view in end-systole displaying mitral valve prolapse with MAD (yellow arrow). The red line indicates the plane of the mitral annulus. MAD = mitral annulus disjunction; MRI = Magnetic Resonance Imaging.

**Figure 5: 12-lead ECG showing T wave inversion in infero-lateral leads.**

An ECG of a 44 year old female who presented with ventricular fibrillation and was found to have mitral valve prolapse. The tracing is remarkable for T wave inversion in leads II, III, AVF, V4-V6.

**Figure 6: 24h Holter monitor showing complex ventricular ectopy and non-sustained ventricular tachycardia (NSVT).**

24h Holter monitoring of two patients with mitral valve prolapse, mitral annular disjunction, and palpitations. It shows a right bundle branch block with non-sustained ventricular tachycardia, likely from the posteromedial papillary muscle.

**Figure 7: Left ventricular mechanical dispersion associated with MAD.**

*Figure 7-1st panel:* Transthoracic echocardiographic long-axis view in end-systole (A) displaying mitral valve prolapse with MAD (yellow arrow). Transthoracic echocardiographic apical 4-chamber (B) 3-chamber (C) and 2-chamber (D) views for the measurement of global longitudinal strain (E). Note the disjunction inducing mechanical dispersion and post-systolic shortening of the posterior-basal segment of the left ventricle (in red). The patient has been seen for ventricular premature complexes. (F) The disjunction is associated with fibrosis in the basal inferolateral segments of the left ventricle and mild mitral regurgitation.

*Figure 7-2nd panel:* Transthoracic echocardiographic long-axis view in end-systole (G) displaying mitral valve prolapse with MAD (yellow arrow). Note here the disjunction involving all the insertion of the posterior leaflet on the annulus on transthoracic echocardiographic apical 3-chamber (H) and 4-chamber views (I, J K). The depth of the disjunction is only mild yet the circumferential extension is extensive. (L) It leads to mechanical dispersion, post-systolic shortening of the posterior-basal segment of the left ventricle. The patient has been seen for palpitations and > 10% ventricular premature complexes.

MAD = mitral annular disjunction.

**Figure 8: Typical examples of premature ventricular contractions arising from the mitral apparatus**.

PVCs arising from the **posterior papillary muscle** (PPM) are characterized by RBBB morphology, a predominantly negative QRS in V5, QRS >130 ms and superior axis with both leads II and III negative. Foci in the **anterior papillary muscle** (APM) also yield a RBBB morphology, typically with a late R/S transition (V3 to V5), QRS >130 ms and lead II discordantly negative while lead III is positive.

Ectopy originating from the mitral annulus result in RBBB morphology with positive precordial concordance. PVCs form the **anterior** **mitral annulus** (AMA) are defined by an inferior QRS axis while those originating from the **posterior** **mitral annulus** (PMA) result in a superior axis**.**

PVC = premature ventricular complex, RBBB = right bundle branch block.

Image courtesy of Jérôme Hourdain, MD, Hôpital la Timone, Marseilles, France.