

Arrhythmogenic left ventricular cardiomyopathy

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Arrhythmogenic cardiomyopathy (ACM) is a

genetic heart muscle disease characterised by

substitution of the ventricular myocardium

by fibrofatty tissue.¹ The disease was origi-

nally termed 'arrhythmogenic right ventricular

(dysplasia/) cardiomyopathy' (ARVC) to define

a condition which distinctively affected the right

ventricle (RV) and predisposed to potentially fatal

ventricular arrhythmias, particularly in young indi-

viduals and athletes.²⁻⁴ New insights arising from

postmortem investigations, genotype-phenotype

correlation studies and myocardial tissue charac-

terisation by contrast-enhanced cardiac magnetic

resonance (CMR) led to increased awareness that

the disease often also involves the left ventricle

(LV).^{5–11} The current designation of 'arrhythmogenic cardiomyopathy' better reflects the evolving

concept of a heart muscle disease affecting both

ventricles, with some phenotypic variants char-

acterised by a parallel or predominant involve-

ment of the LV. The adjective 'arrhythmogenic' refers to the distinctive clinical propensity of the

disease to cause ventricular arrhythmias in relation

to the fibrofatty scar tissue, which is a recognised

arrhythmogenic myocardial substrate. Patients

with a long-standing disease may develop RV or

from that provided by the 2019 expert consensus

statement endorsed by the Heart Rhythm Society

(HRS),¹² that is, a non-specific entity which encom-

passes a heterogeneous group of heart muscle disor-

ders, including 'systemic' diseases such as sarcoidosis

and amyloidosis; 'inflammatory' diseases such as

myocarditis; 'infectious' diseases such as Chagas

disease; and 'genetic' diseases such as desmosomal and non-desmosomal forms; and 'ion channel

diseases'. According to the HRS document, the

vague common denominator of this miscellaneous

group of 'arrhythmogenic cardiomyopathies' was

the 'clinical presentation with symptoms or docu-

mentation of atrial fibrillation, conduction disease,

and/or RV and/or LV arrhythmia'. The HRS defi-

nition appears inappropriate because ACM is a

nosographically distinct condition characterised by

following major phenotypes: (1) the original ARVC

phenotype with predominant RV involvement and

no LV abnormalities ('dominant-right' variant); (2)

the phenotypic variant characterised by the equal

involvement of both ventricles ('biventricular');

and (3) the phenotypic variant characterised by

predominant LV involvement with no or minor RV

abnormalities ('dominant-left'), also referred to as

'arrhythmogenic left ventricular cardiomyopathy

The current classification of ACM includes the

typical cardiomyopathic features.

(ALVC)'.^{13 14}

This definition of ACM substantially differs

biventricular dysfunction.⁴

INTRODUCTION

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Learning objectives

- To review the genetic background and the phenotypic features including clinical, ECG, arrhythmic and imaging findings of arrhythmogenic left ventricular cardiomyopathy (ALVC).
- To address the 2020 international criteria for diagnosis of arrhythmogenic cardiomyopathy (the 'Padua criteria'), including the diagnostic criteria for ALVC and differential diagnosis with major phenocopies.
- To evaluate the current risk stratification and therapy for ALVC, with particular reference to indications for implantation of a defibrillator.

In 2000 we had the privilege to write a commissioned article for Education in Heart on 'Arrhythmogenic right ventricular cardiomyopathy: pathogenesis, diagnosis and therapy'.¹⁵ Twenty years later, we are proud to provide for the same educational section of *Heart* with an updated article on 'arrhythmogenic left ventricular cardiomyopathy (ALVC)', which represents the more recently characterised variant at the other end of the broad spectrum of the phenotypic expression of ACM.

The present article addresses the available knowledge on the genetic background, phenotypic features including clinical, ECG, arrhythmic and imaging findings, risk stratification and therapy of ALVC. The 2020 international diagnostic criteria for ACM (the 'Padua criteria'), which included criteria for diagnosis of left-sided phenotypic variants, will be also discussed.¹⁴

PHENOTYPE

The increasing availability of pathological and clinical studies on patients with left-sided phenotypic variants of ACM has provided significant insights into the prevalence and the clinical manifestations of ALVC.⁵⁻²³

In the seminal pathological study on the heart of sudden cardiac death (SCD) victims (or patients undergoing heart transplantation) with ACM, evidence of macroscopic and/or histological LV involvement was found in 76% of cases, a figure showing that the disease affects the LV in the majority of cases.⁶ LV lesions usually affected both the LV free wall and the septum, more often regionally, with a predilection for the posteroseptal and posterolateral areas. The LV lesion was most often located in the outer layer of the wall musculature and consisted of large areas of subepicardial (and less frequently mid-myocardial) fibrofatty myocardium replacement, similarly to the RV lesions. A more recent autopsy study confirmed that LV involvement was observed in majority (87%) of ACM-related SCD victims (20% athletes) and appeared as an isolated pathological finding (ALVC) in almost a fifth of cases.¹⁰ The most common areas of fibrofatty infiltration were the LV posterobasal (68%) and anterolateral (58%) walls. Postmortem genetic testing yielded pathogenic variants in ACMrelated genes in 25% decedents.

CMR studies in living patients fulfilling the 2010 International Task Force (ITF) criteria have consistently shown that LV involvement in terms of morphofunctional (LV global or regional systolic dysfunction) and/or structural (LV late gadolinium enhancement (LGE)) abnormalities is identified in more than half of patients.^{9 22 24}

According to the available findings of clinical studies, phenotypic features of left-sided ACM include the following (figure 1): (1) ECG abnormalities such as low-amplitude QRS complexes (peak to peak < 0.5 mV) in limb leads and T-wave inversion or flattening in the lateral (or inferolateral) leads. although the ECG is often normal; (2) ventricular arrhythmias with a right bundle branch block (RBBB) morphology of the ectopic QRS (denoting the origin from the LV); (3) normal or slightly depressed LV systolic function with no (or mild) dilatation; (4) large amount of myocardial fibrosis evidenced by contrast-enhanced CMR as LGE; and (5) 'non-ischemic' pattern of LGE, predominantly involving the subepicardial layers of the inferior and the inferolateral regions. This last finding is in agreement with the notion that the wave-front of myocardial loss and fibrofatty replacement in the LV wall proceeds from the epicardium to the endocardium, with scar tissue mostly confined to the epicardial layers of the wall. The subendocardial layer which mostly accounts for the regional LV wall contractility is usually conserved, a factor which explains the normal results of echocardiography, cine-CMR and left ventriculography in a sizeable proportion of patients, due to the preservation of global LV systolic function and the lack of wall motion abnormalities.^{22 25} As a corollary, an imaging approach limited to a mere evaluation of the LV function, either global or regional, by echocardiography, angiography or cine-CMR appears insufficient to detect LV involvement and characterise the LV phenotype in patients with ACM.

GENOTYPE

Most of the pathogenic mutations of ACM affect genes encoding structural proteins that are involved in the organisation of intercellular junctions.^{1 4 26 27} These include genes encoding cardiac desmosomal proteins most frequently, such as plakophilin (*PKP2*), desmoplakin (*DSP*), desmoglein (*DSG2*) and desmocollin (*DSC2*), and rarely (<1%) genes encoding for adherens junctional proteins of the 'area composita', such as α -T-catenin (*CTNNA3*) and N-cadherin (*CDH2*). ACM-causing mutations have also been found in non-desmosomal genes such as phospholamban (*PLN*), filamin C (*FLNC*), desmin (*DES*), titin (*TTN*) and lamin A/C (*LMNA*),

which are associated with other cardiomyopathies, such as dilated cardiomyopathy (DCM) and neuromuscular cardiomyopathies, and may lead to overlapping phenotypes. Causal mutations in nondesmosomal genes, such as transmembrane protein 43 (*TMEM43*) and transforming growth factor beta-3 (*TGFβ-3*) genes, have been uncommonly identified. Genotype–phenotype correlation studies have demonstrated that mutations in *DSP*, *PLN* and *FLNC* genes are the most common gene defects causing left-sided ACM (figure 2).^{1 11 17–23}

Desmoplakin is one of the major components of the desmosome that connects the cytoskeletal networks to the cellular membrane, specifically linking the desmocollins and desmogleins to the intermediate filament through interaction with plakoglobin and plakophilin. A number of human *DSP* gene mutations have been linked with ACM, which manifest characteristically with early LV involvement occurring in isolation or preceding RV disease.²⁸ Of note, in the initial report of the *DSP* gene mutation responsible for 'Carvajal syndrome', the cardiac phenotype resembled that of DCM as opposed to the classic ARVC phenotype.²⁹

Phospholamban normally inhibits the sarcoendoplasmic reticulum calcium transport ATPase, and *PLN* gene mutations cause dysregulated calcium flux, predisposing to prominent arrhythmia and ventricular dysfunction. Mutations in *PLN* gene are relatively common in the Netherlands, where a founder mutation of *PLN* (p.Arg14del) is found in up to 15% of probands with ACM. *PLN* mutation most frequently gives rise to an ACM predominantly affecting the LV, similar to DCM.²¹

Mutations of the *FLNC* gene were initially associated with a particular form of skeletal myofibrillar myopathy. The protein product, filamin C, is mainly expressed in skeletal and cardiac muscles and functions at Z-discs and in the subsarcolemmal regions. Truncating mutations of *FLNC* were subsequently shown to lead to cardiomyopathy, without skeletal muscle involvement. Isolated *FLNC*-related cardiomyopathy shows an overlapping phenotype with DCM and is characterised by LV systolic dysfunction with mild dilatation and marked fibrosis.¹⁹

DIAGNOSIS

In 2020, an international expert report critically reviewed the 2010 Task Force (TF) criteria for the diagnosis of ARVC with the aim to identify areas of potential improvement.¹³ The 2010 TF guidelines³⁰ exclusively targeted the original RV phenotype features, did not provide diagnostic criteria for the left-sided forms, and lacked tissue characterisation findings by gadolinium-enhanced CMR for detection of myocardial fibrosis, which are determinants of an accurate diagnosis of the LV phenotype.^{13 22-24}

A 2020 international expert consensus document provided upgraded criteria for the diagnosis of ACM, with the introduction of specific diagnostic criteria for the LV phenotype.¹⁴ The 2020 international criteria, also known as the 'Padua criteria', represents a modern approach to diagnosis

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Figure 1 Phenotypic features of arrhythmogenic left ventricular cardiomyopathy. ECG strips showing low QRS voltages (<0.5 mV) in the limb leads (arrow) (A), negative T-waves in leads V4–V6 (arrows) (B), premature ventricular beat with a right bundle branch block and superior axis (C), and ventricular tachycardia with a right bundle branch block and inferior axis morphology (D). Two-dimensional echocardiogram (diastolic frame in four-chamber view) showing a non-dilated and normokinetic (not shown) left ventricle (E). Cardiac magnetic resonance (diastolic frame of four-chamber view on postcontrast sequences) showing diffuse late gadolinium enhancement involving the subepicardial layer of the left ventricle free wall and the septum (F). Histological panoramic view of the posterolateral left ventricular wall showing replacement-type myocardial fibrosis mostly involving the subepicardial layer (trichrome stain) (G).

of ACM based on research achievements and the clinical and imaging progress over 30 years at the Medical School of the University of Padua. The Padua criteria were reviewed and shared by several international experts, resulting in an international consensus document.

According to the 2020 international criteria, the diagnosis of biventricular and left-dominant variants



Figure 2 Disease-causing genetic defects in arrhythmogenic cardiomyopathy according to predominant ventricular involvement. Mutations in *DSP*, *PLN* and *FLNC* genes are the most common gene defects causing arrhythmogenic left ventricular cardiomyopathy. LV, left ventricle; RV, right ventricle.

of ACM is based on the same multiparametric approach recommended by the 2010 TF criteria for classic ARVC, with six categories of criteria which encompass morphofunctional ventricular abnormalities, structural myocardial tissue alterations, electrocardiographic changes of ventricular depolarisation and repolarisation, ventricular arrhythmias, and familial and genetic findings. The criteria considered *necessary* for the diagnosis of LV disease variants are classified as 'major', while those *not necessary* but contributing to the refinement of LV phenotype characterization as 'minor'.

The proposed 2020 international criteria by different categories for the diagnosis of RV and LV phenotypes are summarised in table 1.

While based on the 2010 TF criteria myocardial tissue characterisation relied exclusively on the histopathological study by endomyocardial biopsy samples, the 2020 international criteria recommend that structural myocardial abnormalities are investigated by the CMR LGE technique, which allows the identification of replacement of the LV myocardium by fibrofatty scar tissue.^{13 22-24 31-34}

The accuracy of the 2020 diagnostic scoring system for phenotypic diagnosis of left-sided ACM varies according to different disease forms. In case of biventricular ACM, the fulfilment of the diagnostic criteria for the RV phenotype ensures that the concomitant left-sided abnormalities are disease-specific. Hence, the diagnosis of biventricular ACM can be reliably made in the presence of phenotypic features such as morphofunctional and structural abnormalities of both ventricles.

In patients without clinically detectable RV abnormalities, LV phenotypic criteria in isolation do not allow a conclusive diagnosis of ALVC as the morphofunctional and structural LV abnormalities are not sufficiently disease-specific due to the overlap with the phenotypic features of other heart muscle diseases such as DCM, myocarditis or cardiac sarcoidosis.¹⁵ Hence, in the presence of consistent LV phenotypic features, demonstration of a pathogenic or likely pathogenic mutation of ACM related genes is mandatory for diagnosis of ALVC.

According to the 2020 international criteria, the diagnosis of each phenotypic variant of ACM requires that at least one criterion from category 1 (ie, morphofunctional ventricular abnormalities) or category 2 (ie, structural myocardial abnormalities), either major or minor, is met. The reason is that ACM is not a channelopathy whose diagnosis relies on demonstration of pathogenic mutations, ECG abnormalities or arrhythmias in the presence of a normal heart, but a cardiomyopathy with distinctive structural ventricular abnormalities, which are an integral part of the phenotype.

Although CMR has become the leading imaging modality for diagnosis of ACM due to its unique capability of identifying fibrofatty myocardial replacement by LGE tissue characterisation, novel and sensitive ultrasound techniques such as strain echocardiography can be useful for detection of early ventricular dysfunction.^{35 36}

In the 2020 international criteria, the scoring system for diagnosis of left-sided phenotypic variants of ACM is substantially unchanged compared with the 2010 TF criteria for diagnosis of ARVC.¹⁴ The biventricular variant is diagnosed in patients fulfilling morphofunctional and/or structural abnormalities of both the RV and the LV. The diagnosis is classified as *definite* when two major, one major and two minor, or four minor criteria of different categories from both ventricles are fulfilled; borderline, one major and one minor, or three minor criteria; and *possible*, one major or two minor criteria. Of note, ≥ 1 morphofunctional and/or structural criteria, either major or minor, are needed for each diagnosis. The diagnosis of ALVC, without clinical evidence of RV abnormalities, needs demonstration of ACM-causing gene mutations or familial ACM, in association with consistent LV structural abnormalities (figure 3).

DIFFERENTIAL DIAGNOSIS

The ALVC phenotype can overlap with that of other genetic cardiomyopathies, neuromuscular diseases, or non-genetic disorders such as acquired DCM, myocarditis and sarcoidosis.¹³ These latter heart muscle disorders represent the most challenging differential diagnoses with ALVC.

Dilated cardiomyopathy

DCM is the most common condition requiring a differential diagnosis with ALVC due to overlapping phenotypic features such as LV systolic dysfunction and myocardial fibrosis. The main discriminant features between the two cardiomyopathies are reported in table 2. Significant differences are identified by CMR imaging study, which consist of the specific LV remodelling pattern, the extent and regional distribution of myocardial fibrosis as evidenced by LGE, and the relation between the amount of LGE and LV systolic dysfunction.^{16 17 22} The typical ventricular remodelling pattern of ALVC consists of a hypokinetic (or normokinetic) LV, with no or mild cavity dilatation, in comparison with that of DCM characterised by significant LV ventricular dilatation and systolic dysfunction. Tissue characterisation findings by CMR offer the potential to identify and quantify the myocardial LGE/fibrosis. While LGE is detected in <50% of DCM cases, 100% of patients with ALVC with LV systolic dysfunction show the presence of LV LGE.²² The distribution of LGE differs between the two conditions, predominantly affecting the subepicardial inferolateral regions in ALVC versus mid-mural septal segments in DCM (figure 4). In ALVC, the coexistence of fatty myocardial infiltration is often observed on dedicated sequences in the same regions of LGE or in remote LV areas.

The amount of LV myocardial fibrosis/LGE is significantly greater in patients with ALVC than in patients with DCM (online supplemental figure $1)^{17}$ ²² and directly related to the severity of LV systolic dysfunction (online supplemental figure $2)^{22}$ because it acts as a pathological marker of

Table 1 The 'Padua criteria' for diagnosis of arrhythmogenic cardiomyopathy			
Category	Right ventricle	Left ventricle	
1. Morphofunctional ventricular abnormalities	 By echocardiography, CMR or angiography: Major Regional RV akinesia, dyskinesia or bulging <i>plus</i> one of the following: Global RV dilatation (increase in RV EDV according to the imaging test-specific nomograms). Global RV systolic dysfunction (reduction in RV EF according to the imaging test-specific nomograms). Minor Regional RV akinesia, dyskinesia or angurysm of RV free wall 	 By echocardiography, CMR or angiography: Minor Global LV systolic dysfunction (depression of LV EF or reduction of echocardiographic global longitudinal strain), with or without LV dilatation (increase in LV EDV according to the imaging test-specific nomograms for age, sex and BSA). Minor Regional LV hypokinesia or akinesia of LV free wall, septum or both. 	
2. Structural myocardial abnormalities	 By CE-CMR: Major Transmural LGE (stria pattern) of ≥1 RV region(s) (inlet, outlet and apex in two orthogonal views). By EMB (limited indications): Major Fibrous replacement of the myocardium in ≥1 sample, with or without fatty tissue. 	 By CE-CMR: Major LV LGE (stria pattern) of ≥1 Bull's eye segment(s) (in two orthogonal views) of the free wall (subepicardial or mid-myocardial), septum or both (excluding septal junctional LGE). 	
3. Repolarisation abnormalities	 Major Inverted T-waves in right precordial leads (V1, V2 and V3) or beyond in individuals with complete pubertal development (in the absence of complete RBBB). Minor Inverted T-waves in leads V1 and V2 in individuals with complete pubertal development (in the absence of complete RBBB). Inverted T-waves in V1, V2, V3 and V4 in individuals with complete pubertal development in the presence of complete RBBB. 	 Minor Inverted T-waves in left precordial leads (V4–V6) (in the absence of complete LBBB). 	
4. Depolarisation abnormalities	 Minor Epsilon wave (reproducible low-amplitude signals between end of QRS complex and onset of the T-wave) in the right precordial leads (V1–V3). Terminal activation duration of QRS ≥55 ms measured from the nadir of the S-wave to the end of the QRS, including R', in V1, V2 or V3 (in the absence of complete RBBB). 	 Minor Low QRS voltages (<0.5 mV peak to peak) in limb leads (in the absence of obesity, emphysema or pericardial effusion). 	
5. Ventricular arrhythmias	 Major Frequent ventricular extrasystoles (>500 per 24 hours), non-sustained or sustained ventricular tachycardia of LBBB morphology. Minor Frequent ventricular extrasystoles (>500 per 24 hours), non-sustained or sustained ventricular tachycardia of LBBB morphology with inferior axis ('RVOT pattern'). 	 Minor Frequent ventricular extrasystoles (>500 per 24 hours), non-sustained or sustained ventricular tachycardia with an RBBB morphology (excluding the 'fascicular pattern'). 	
6. Family history/genetics	 Major ACM confirmed in a first-degree relative who meets the diagnostic criteria. ACM confirmed pathologically at autopsy or surgery in a first-degree relative. Identification of a pathogenic or likely pathogenetic ACM mutation in the patient under evaluation. Minor History of ACM in a first-degree relative in whom it is not possible or practical to determine whether the family member meets the diagnostic criteria. Premature sudden death (<35 years of age) due to suspected ACM in a first-degree relative.ACM confirmed pathologically or by diagnostic criteria in a second-degree relative. diomyopathy: BSA, body surface area; CE-CMR, contrast enhanced-cardiac magnetic resonance; EDV, end diastolic volume; EF, ejection fraction; FMR. 		

ACM, arrhythmogenic cardiomyopathy; BSA, body surface area; CE-CMR, contrast enhanced-cardiac magnetic resonance; EDV, end diastolic volume; EF, ejection fraction; EMB, endomyocardial biopsy; LBBB, left bundle branch block; LGE, late gadolinium enhancement; LV, left ventricle; RBBB, right bundle branch block; RV, right ventricle; RVOT, right ventricular outflow tract.

myocardial contractile reserve. On the contrary, LV LGE in DCM represents an epiphenomenon and is unrelated to the reduction of the LV systolic function.²²

In the current guidelines of the European Society of Cardiology, the so-called 'hypokinetic, non-dilated LV cardiomyopathy' has been classified as part of the clinical spectrum of DCM and considered as a disease variant with a less expressed phenotype.³⁷ However, there is compelling evidence that this cardiomyopathy remodelling, distinctively characterised by extensive, non-transmural scarring of the LV which scarcely affects the LV global and regional systolic function, fits better with the phenotype of ALVC, suggesting that a sizeable proportion of cardiomyopathies previously diagnosed as



Figure 3 Morphofunctional and structural characteristics of phenotypic variants of arrhythmogenic cardiomyopathy. Note that demonstration of a pathogenic arrhythmogenic cardiomyopathy gene mutation (or familial arrhythmogenic cardiomyopathy) is mandatory for diagnosis of ALVC, in association with consistent structural myocardial abnormalities. ALVC, arrhythmogenic left ventricular cardiomyopathy; ARVC, arrhythmogenic right ventricular (dysplasia/) cardiomyopathy; LV, left ventricle; RV, right ventricle.

morphologically mild expression of DCM should be reclassified as ALVC. $^{16\,17\,22}$

The LV systolic dysfunction of ALVC may become more severe in advanced stages, as a result of a disease progression by increasing the extent of LGE which affects multiple segments of the LV free wall and septum, with a more transmural involvement.^{1 15} Demonstration of an ACM-causing gene mutation in association with a consistent phenotype is mandatory for diagnosis of ALVC.¹⁴

Myocarditis

Isolated LV LGE with subepicardial/midmyocardial distribution (ie, non-ischaemic) is traditionally interpreted as the consequence of a previous myocarditis, although there is increasing evidence that it may reflect a segmental ALVC (figure 5).^{25 38 39} Similar to ALVC, a postinflammatory scar may induce ECG changes, such as low-amplitude ORS complexes in limb leads. T-wave inversion and ventricular arrhythmias with RBBB. Moreover, because of its focal and non-transmural wall involvement, the myocardial scar may be undetectable by echocardiography.³⁸ ³⁹ Non-ischaemic myocardial scar has been reported to act as a myocardial substrate for life-threatening ventricular arrhythmias and cardiac arrest, mostly in young people and athletes. A previous study reported that during a mean 3-year follow-up 22% of athletes with LV scar-related ventricular arrhythmias experienced major arrhythmic events including SCD, compared with none of the control athletes with ventricular arrhythmias in the absence of LV scar.²⁵ The most important clinical implication of detection of a non-ischaemic LV scar is

Table 2 Differential diagnosis between ALVC and DCM

	ALVC	DCM
Predominant genetic background	Mutations of genes encoding for desmosomal proteins, phospholamban and filamin C.	Mutations of genes encoding for cytoskeleton, muscular sarcomere and nuclear envelope proteins.
Main clinical manifestations	Palpitations, syncope and cardiac arrest.	Heart failure and cardiac arrest.
ECG abnormalities	Low QRS voltages in limb leads; negative T-waves in (infero-)lateral leads.	Left ventricular hypertrophy with a strain pattern of ST segment; LBBB.
Echocardiography and CMR imaging features	Non-dilated and hypokinetic LV with a large amount of non-ischaemic (subepicardial) LGE (inferolateral LV wall). Regional wall motion abnormalities (common). Systolic LV dysfunction related to the global extent of LGE.	Dilated and hypokinetic LV with no or patchy non-ischaemic (mid-myocardial) LGE (septum). Regional wall motion abnormalities (uncommon). Systolic LV dysfunction unrelated to the global extent of LGE.
Histopathological features	Fibrofatty myocardial replacement.	Non-specific myocardial abnormalities±fibrosis.
Types of ventricular arrhythmias	PVBs, NSVT and monomorphic sustained VT (RBBB pattern; both LBBB and RBBB patterns in biventricular form); VF.	PVBs and NSVT (RBBB pattern); uncommon sustained VT; VF.
Mechanism of VT	Scar-related re-entry.	Scar-related or functional re-entry (branch to branch re-entry).
Most common site of VT origin	Subepicardial inferolateral LV free wall.	Intramural septum.

ALVC, arrhythmogenic left ventricular cardiomyopathy; CMR, cardiac magnetic resonance; DCM, dilated cardiomyopathy; LBBB, left bundle branch block; LGE, late gadolinium enhancement; LV, left ventricle; NSVT, non sustained ventricular tachycardia; PVBs, premature ventricular beats; RBBB, right bundle branch block; VF, ventricular fibrillation; VT, ventricular tachycardia.



Figure 4 Left ventricular remodelling and regional distribution of LGE in ACM-LV phenotype versus DCM-LV phenotype. Comparison between CMR findings in the subgroup of patients with ACM (left) and patients with DCM (right) with overlapping LV phenotypes (ie, LV systolic dysfunction and LV LGE) shows that (1) ACM has lower LV volume and mass, less depressed LVEF (not shown) and greater amount of LV LGE (ie, 'hypokinetic, non-dilated and fibrotic left ventricle') compared with patients with DCM); (2) LV LGE predominately affects the inferolateral segments in ACM and septal segments in DCM; and (3) in ACM the LV LGE appears as a stria pattern that more frequently affects the subepicardial layers, compared with the spot/patchy pattern and the mid-myocardial location of DCM . Modified from Cipriana *et al.*²² ACM, arrhythmogenic cardiomyopathy; ARVC, arrhythmogenic right ventricular (dysplasia/) cardiomyopathy; CMR, cardiac magnetic resonance; DCM, dilated cardiomyopathy; LGE, late gadolinium enhancement; LV, left ventricular; LVEF, left ventricular ejection fraction.



Figure 5 Cardiac magnetic resonance features and histopathological findings of structural changes in myocarditis and ALVC. Myocarditis (A and B): postcontrast T1 inversion recovery sequence in short-axis view showing subepicardial LGE of the inferolateral LV wall (arrows) (A); corresponding panoramic histopathological view of the inferolateral LV wall showing extensive fibrous tissue replacement in the subepicardial layer of the myocardium (B). Desmosomal gene-related ALVC in a sudden cardiac death victim carrying a *DSP* mutation (C and D): postcontrast T1 inversion recovery sequence in short-axis view showing subepicardial LGE of the inferolateral LV wall (arrows) in a *DSP* gene mutation carrier (C). Panoramic histopathological view showing fibrofatty myocardial replacement of the outer layer of the inferolateral LV wall (D). Modified from Corrado *et al.*¹³ ALVC, arrhythmogenic left ventricular cardiomyopathy; LGE, late gadolinium enhancement; LV, left ventricle.

that it cannot be dismissed as a benign sign of a healed remote inflammatory process but deserves proper clinical attention. Affected patients should be evaluated for a family history of cardiomyopathy, clinical history of acute myocarditis, presence of symptoms and the arrhythmogenicity of the myocardial fibrosis. Molecular genetic testing with demonstration of ACM gene mutation is needed in selected non-familial cases for a conclusive differential diagnosis between ALVC and myocarditis.

Cardiac sarcoidosis

Clinical manifestations common to cardiac sarcoidosis and ALVC include myocardial scarring with dilatation/dysfunction leading to lifethreatening ventricular arrhythmias and heart failure. The most important feature distinguishing cardiac sarcoidosis from ALVC is the presence of infrahisian conduction abnormalities such as bundle branch block and atrioventricular block as a consequence of the predilection of granulomatous infiltration for the basal interventricular septum, which is usually absent in ALVC.⁴⁰ Although definitive diagnosis of sarcoidosis is made by demonstration of granulomatous infiltration on endomyocardial or extracardiac biopsy, non-invasive imaging modalities such as CMR⁴¹ and positron emission tomography⁴² are increasingly used in the clinical setting.

Suggestive of cardiac sarcoidosis are findings of LGE localised in the basal septum and regional fluorodeoxyglucose uptake which indicates active inflammatory lesions (figure 6).

RISK STRATIFICATION AND THERAPY

The first observational study on the clinical impact of implantable cardioverted defibrillator (ICD) therapy on the natural history of 134 patients with ARVC showed that nearly half of patients



Figure 6 Magnetic resonance and cardiac positron emission tomography features of cardiac sarcoidosis. Postcontrast T1 inversion recovery sequence in four-chamber view showing LGE of the interventricular septum (black arrows) and subepicardial lateral LV wall (white arrows) (A). Topographic concordance between the epicardial spot of LGE involving the anterior (B, single white arrow) and inferior (B, white arrows) LV regions and the areas of fludeoxyglucose uptake on positron emission tomography (C, white arrows pointing to yellow–red areas). Modified from Corrado *et al.*¹³ LGE, late gadolinium enhancement; LV, left ventricle.

had at least one episode of ventricular tachyarrhythmia that required ICD intervention over a mean follow-up period of 3.3 years and 24% experienced ventricular fibrillation/flutter that in all likelihood would have been fatal without termination by the device. On multivariate analysis, independent predictors of ventricular fibrillation/flutter included history of cardiac arrest or ventricular tachycardia with haemodynamic compromise, younger age and left ventricular involvement (evaluated as a reduction of the LV ejection fraction).⁴³ Subsequently, a study aimed to identify the risk factors for long-term prognosis among 130 patients with ARVC showed that during a mean follow-up of 8 years, 24 patients died (21 from cardiovascular death), with an annual mortality rate of 2.3%. Clinical signs of

Key messages

- Arrhythmogenic cardiomyopathy (ACM) is a biventricular muscle disease with left ventricle (LV) involvement which may parallel (biventricular) or exceed (left-dominant/arrhythmogenic left ventricular cardiomyopathy (ALVC)) the severity of the right ventricle (RV) disease.
- 'Functional' imaging evaluation of the LV is insufficient for diagnosis of ALVC, which requires myocardial tissue characterisation by the late gadolinium enhancement technique.
- In patients with an ALVC phenotype (and no RV abnormalities), demonstration of a pathogenic mutation in ACM-related genes is mandatory for diagnosis.
- Implantable cardioverted defibrillator implantation for primary prevention should be considered in the presence of a large arrhythmogenic ventricular scar, even if the LV systolic function is not severely depressed.

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The appropriate management strategy for ALVC is not completely established.⁴⁷ Restriction of sports activity and prophylactic β -blocker therapy are indicated to prevent 'adrenergic dependent' ventricular arrhythmias. Successful mapping/catheter ablation of scar-related sustained ventricular tachycardia (VT)is feasible using an epicardial approach through the pericardial space to reach the re-entry circuit which is located in the outer layer of LV wall.⁴⁸ Only the implantation of an ICD has been proven to successfully prevent SCD by interrupting potentially lethal ventricular arrhythmias.^{43 47} According to the 2015 TF consensus documents, three categories of arrhythmic risk ('high', 'moderate' and 'low' risk) are considered for indication to ICD implantation.⁴⁷ It is noteworthy that the LV systolic function was a key factor for risk stratification. The 'high-risk' category (class I recommendation to ICD implantation) comprised patients with a history of cardiac arrest or haemodynamically unstable VT or those with severe ventricular dysfunction, either right (RV fractional area change ≤17% or RV ejection fraction ≤ 35 %) or left (LV ejection fraction \leq 35%). ICD implantation was also indicated (class IIa recommendation) in patients from the 'intermediate risk' category with ≥ 1 'major' risk factors, such as syncope, non-sustained VT or moderate right (RV fractional area change 17%-24% or RV ejection fraction 36%-40%) and/or left (LV ejection fraction 36%-45%) ventricular dysfunction. This reflects the concept that, at variance with DCM, the pathogenesis of life-threatening ventricular arrhythmias in ALVC extends beyond the severe depression of LV systolic function, being strongly related to the large amount of myocardial fibrosis, which is an independent arrhythmogenic risk factor.^{22 23} Thus, ICD therapy for primary prevention of SCD should be considered earlier in the disease course, when the degree of LV systolic dysfunction is still mild or moderate.¹³ Subcutaneous ICD has been proven to substantially reduce lead-related complications but maintaining efficacy and should be considered for primary prevention in young patients.⁴⁹

Patients with advanced ALVC and severe systolic dysfunction are treated with the traditional heart failure therapy.⁴⁷ Heart transplantation is the final therapeutic option for those patients with either end-stage and unresponsive congestive heart failure or VT/ventricular fibrillation storms refractory to catheter (and surgical) ablation and/or ICD therapy.⁵⁰

SUMMARY/CONCLUSIONS

The spectrum of ACM phenotypes is broader than previously believed and besides ARVC includes biventricular variants and ALVC. The 2010 TF criteria lacked specific criteria for diagnosis of leftsided phenotypes, which resulted in underdiagnosis of ACM. The 2020 international diagnostic criteria were provided to fill the gaps of previous TF guidelines, mostly by introduction of a new criteria for the diagnosis of the LV phenotype, mostly based on CMR myocardial tissue characterisation findings which are determinants for detection of ALVC. Clinical studies on large patient cohorts are needed to validate the accuracy and quantitate the impact of the 2020 international criteria in diagnosing the entire spectrum of ACM variants, with particular reference to ALVC.

Future directions of research include comprehensive understanding of the pathogenesis, increased awareness of clinical manifestations and outcome, as well as more accurate risk stratification and effective therapy for ALVC.

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REFERENCES

- *1 Corrado D, Basso C, Judge DP. Arrhythmogenic cardiomyopathy. *Circ Res* 2017;121:784–802.
- 2 Marcus FI, Fontaine GH, Guiraudon G, et al. Right ventricular dysplasia: a report of 24 adult cases. Circulation 1982;65:384–98.
- 3 Thiene G, Nava A, Corrado D, *et al*. Right ventricular cardiomyopathy and sudden death in young people. *N Engl J Med* 1988;318:129–33.
- 4 Corrado D, Link MS, Calkins H. Arrhythmogenic right ventricular cardiomyopathy. *N Engl J Med* 2017;376:61–72.
- 5 Basso C, Thiene G, Corrado D, et al. Arrhythmogenic right ventricular cardiomyopathy. *Circulation* 1996;94:983–91.
- 6 Corrado D, Basso C, Thiene G, *et al*. Spectrum of clinicopathologic manifestations of arrhythmogenic right ventricular cardiomyopathy/dysplasia: a multicenter study. *J Am Coll Cardiol* 1997;30:1512–20.
- *7 Sen-Chowdhry S, Syrris P, Prasad SK, et al. Left-dominant arrhythmogenic cardiomyopathy: an under-recognized clinical entity. JAm Coll Cardiol 2008;52:2175–87.
- 8 Sen-Chowdhry S, Syrris P, Ward D, et al. Clinical and genetic characterization of families with arrhythmogenic right ventricular dysplasia/cardiomyopathy provides novel insights into patterns of disease expression. *Circulation* 2007;115:1710–20.
- 9 Marra MP, Leoni L, Bauce B, et al. Imaging study of ventricular scar in arrhythmogenic right ventricular cardiomyopathy: comparison of 3D standard electroanatomical voltage mapping and contrast-enhanced cardiac magnetic resonance. *Circ Arrhythm Electrophysiol* 2012;5:91–100.
- Miles C, Finocchiaro G, Papadakis M, et al. Sudden death and left ventricular involvement in arrhythmogenic cardiomyopathy. *Circulation* 2019;139:1786–97.
- 11 Thiene G, Corrado D, Nava A, *et al*. Right ventricular cardiomyopathy: is there evidence of an inflammatory aetiology? *Eur Heart J* 1991;12 Suppl D:22–5.
- 12 Towbin JA, McKenna WJ, Abrams DJ. Hrs expert consensus statement on evaluation, risk stratification, and management of arrhythmogenic cardiomyopathy. *Heart Rhythm* 2019;2019:e373–407.
- *13 Corrado D, van Tintelen PJ, McKenna WJ, et al. Arrhythmogenic right ventricular cardiomyopathy: evaluation of the current diagnostic criteria and differential diagnosis. Eur Heart J 2020;41:1414–29.
- *14 Corrado D, Perazzolo Marra M, Zorzi A, *et al*. Diagnosis of arrhythmogenic cardiomyopathy: the Padua criteria. *Int J Cardiol* 2020;319:106–14.
- 15 Corrado D, Basso C, Thiene G. Arrhythmogenic right ventricular cardiomyopathy: diagnosis, prognosis, and treatment. *Heart* 2000;83:588–95.
- 16 Segura-Rodríguez D, Bermúdez-Jiménez FJ, Carriel V, et al. Myocardial fibrosis in arrhythmogenic cardiomyopathy: a genotype-phenotype correlation study. *Eur Heart J Cardiovasc Imaging* 2020;21:378–86.
- 17 Augusto JB, Eiros R, Nakou E, *et al*. Dilated cardiomyopathy and arrhythmogenic left ventricular cardiomyopathy: a comprehensive genotype-imaging phenotype study. *Eur Heart J Cardiovasc Imaging* 2020;21:326–36.
- 18 Hall CL, Akhtar MM, Sabater-Molina M, et al. Filamin C variants are associated with a distinctive clinical and immunohistochemical arrhythmogenic cardiomyopathy phenotype. Int J Cardiol 2020;307:101–8.
- 19 Ortiz-Genga MF, Cuenca S, Dal Ferro M, et al. Truncating FLNc mutations are associated with high-risk dilated and arrhythmogenic cardiomyopathies. J Am Coll Cardiol 2016;68:2440–51.
- 20 Groeneweg JA, van der Zwaag PA, Olde Nordkamp LRA, et al. Arrhythmogenic right ventricular dysplasia/cardiomyopathy according to revised 2010 Task force criteria with inclusion of non-desmosomal phospholamban mutation carriers. Am J Cardiol 2013;112:1197–206.
- 21 Te Rijdt WP, Ten Sande JN, Gorter TM, *et al*. Myocardial fibrosis as an early feature in phospholamban p.Arg14del mutation carriers:

Education in Heart

phenotypic insights from cardiovascular magnetic resonance imaging. *Eur Heart J Cardiovasc Imaging* 2019;20:92–100.

- *22 Cipriani A, Bauce B, De Lazzari M, *et al.* Arrhythmogenic right ventricular cardiomyopathy: characterization of left ventricular phenotype and differential diagnosis with dilated cardiomyopathy. *J Am Heart Assoc* 2020;9:e014628.
- 23 Smith ED, Lakdawala NK, Papoutsidakis N, et al. Desmoplakin cardiomyopathy, a fibrotic and inflammatory form of cardiomyopathy distinct from typical dilated or arrhythmogenic right ventricular cardiomyopathy. *Circulation* 2020;141:1872–84.
- 24 De Lazzari M, Zorzi A, Cipriani A, et al. Relationship between electrocardiographic findings and cardiac magnetic resonance phenotypes in arrhythmogenic cardiomyopathy. J Am Heart Assoc 2018;7:e009855.
- 25 Zorzi A, Perazzolo Marra M, Rigato I, *et al.* Nonischemic left ventricular scar as a substrate of life-threatening ventricular arrhythmias and sudden cardiac death in competitive athletes. *Circ Arrhythm Electrophysiol* 2016;9:e004229.
- 26 Hoorntje ET, Te Rijdt WP, James CA, et al. Arrhythmogenic cardiomyopathy: pathology, genetics, and concepts in pathogenesis. *Cardiovasc Res* 2017;113:1521–31.
- 27 Rigato I, Bauce B, Rampazzo A, et al. Compound and digenic heterozygosity predicts lifetime arrhythmic outcome and sudden cardiac death in desmosomal gene-related arrhythmogenic right ventricular cardiomyopathy. *Circ Cardiovasc Genet* 2013;6:533–42.
- 28 Rampazzo A, Nava A, Malacrida S, *et al*. Mutation in human desmoplakin domain binding to plakoglobin causes a dominant form of arrhythmogenic right ventricular cardiomyopathy. *Am J Hum Genet* 2002;71:1200–6.
- 29 Norgett EE, Hatsell SJ, Carvajal-Huerta L, et al. Recessive mutation in desmoplakin disrupts desmoplakin-intermediate filament interactions and causes dilated cardiomyopathy, woolly hair and keratoderma. Hum Mol Genet 2000;9:2761–6.
- 30 Marcus FI, McKenna WJ, Sherrill D, *et al*. Diagnosis of arrhythmogenic right ventricular cardiomyopathy/dysplasia: proposed modification of the task force criteria. *Circulation* 2010;121:1533–41.
- 31 Sen-Chowdhry S, Prasad SK, Syrris P, et al. Cardiovascular magnetic resonance in arrhythmogenic right ventricular cardiomyopathy revisited: comparison with Task force criteria and genotype. J Am Coll Cardiol 2006;48:2132–40.
- 32 Haugaa KH, Basso C, Badano LP, et al. Comprehensive multimodality imaging approach in arrhythmogenic cardiomyopathyan expert consensus document of the European association of cardiovascular imaging. Eur Heart J Cardiovasc Imaging 2017;18:237–53.
- 33 Tandri H, Saranathan M, Rodriguez ER, et al. Noninvasive detection of myocardial fibrosis in arrhythmogenic right ventricular cardiomyopathy using delayed-enhancement magnetic resonance imaging. J Am Coll Cardiol 2005;45:98–103.
- 34 Marra MP, Cipriani A, Rizzo S. Myocardial tissue characterization in arrhythmogenic cardiomyopathy: comparison between cardiac magnetic resonance and endomyocardial biopsy. JACC Cardiovasc Imaging. In Press 2021.
- 35 Sarvari SI, Haugaa KH, Anfinsen O-G, et al. Right ventricular mechanical dispersion is related to malignant arrhythmias: a study of patients with arrhythmogenic right ventricular cardiomyopathy

and subclinical right ventricular dysfunction. *Eur Heart J* 2011;32:1089–96.

- 36 Leren IS, Saberniak J, Haland TF, et al. Combination of ECG and echocardiography for identification of arrhythmic events in early ARVC. JACC Cardiovasc Imaging 2017;10:503–13.
- 37 Pinto YM, Elliott PM, Arbustini E, et al. Proposal for a revised definition of dilated cardiomyopathy, hypokinetic non-dilated cardiomyopathy, and its implications for clinical practice: a position statement of the ESC Working group on myocardial and pericardial diseases. Eur Heart J 2016;37:1850–8.
- 38 Zorzi A, Rigato I, Pilichou K, et al. Phenotypic expression is a prerequisite for malignant arrhythmic events and sudden cardiac death in arrhythmogenic right ventricular cardiomyopathy. *Europace* 2016;18:1086–94.
- 39 Vio R, Zorzi A, Corrado D. Myocarditis in the athlete: arrhythmogenic substrates, clinical manifestations, management, and eligibility decisions. J Cardiovasc Transl Res 2020;13:284–95.
- 40 Philips B, Madhavan S, James CA, *et al*. Arrhythmogenic right ventricular dysplasia/cardiomyopathy and cardiac sarcoidosis: distinguishing features when the diagnosis is unclear. *Circ Arrhythm Electrophysiol* 2014;7:230–6.
- 41 Steckman DA, Schneider PM, Schuller JL, et al. Utility of cardiac magnetic resonance imaging to differentiate cardiac sarcoidosis from arrhythmogenic right ventricular cardiomyopathy. Am J Cardiol 2012;110:575–9.
- 42 Choo WK, Denison AR, Miller DR, *et al*. Cardiac sarcoid or arrhythmogenic right ventricular cardiomyopathy: a role for positron emission tomography (PET)? *J Nucl Cardiol* 2013;20:479–80.
- 43 Corrado D, Leoni L, Link MS, et al. Implantable cardioverterdefibrillator therapy for prevention of sudden death in patients with arrhythmogenic right ventricular cardiomyopathy/dysplasia. *Circulation* 2003;108:3084–91.
- 44 Hulot J-S, Jouven X, Empana J-P, et al. Natural history and risk stratification of arrhythmogenic right ventricular dysplasia/ cardiomyopathy. Circulation 2004;110:1879–84.
- 45 Aquaro GD, De Luca A, Cappelletto C, et al. Prognostic value of magnetic resonance phenotype in patients with arrhythmogenic right ventricular cardiomyopathy. J Am Coll Cardiol 2020;75:2753–65.
- 46 Lie Øyvind H, Rootwelt-Norberg C, Dejgaard LA, et al. Prediction of life-threatening ventricular arrhythmia in patients with arrhythmogenic cardiomyopathy: a primary prevention cohort study. JACC Cardiovasc Imaging 2018;11:1377–86.
- *47 Corrado D, Wichter T, Link MS, et al. Treatment of arrhythmogenic right ventricular cardiomyopathy/dysplasia: an international Task force consensus statement. Eur Heart J 2015;36:3227–37.
- 48 Oloriz T, Silberbauer J, Maccabelli G, et al. Catheter ablation of ventricular arrhythmia in nonischemic cardiomyopathy: anteroseptal versus inferolateral scar sub-types. Circ Arrhythm Electrophysiol 2014;7:414–23.
- 49 Migliore F, Viani S, Bongiorni MG, et al. Subcutaneous implantable cardioverter defibrillator in patients with arrhythmogenic right ventricular cardiomyopathy: results from an Italian multicenter registry. Int J Cardiol 2019;280:74–9.
- 50 Gilljam T, Haugaa KH, Jensen HK, et al. Heart transplantation in arrhythmogenic right ventricular cardiomyopathy - Experience from the Nordic ARVC Registry. Int J Cardiol 2018;250:201–6.



