Arrhythmogenic right ventricular cardiomyopathy/dysplasia (ARVC/D) in clinical practice

Ka Hou Christien Li1 | George Bazoukis MD2 | Tong Liu MD, PhD3 | Guangping Li MD, PhD3 | William K. K. Wu PhD, FRCPath4,5 | Sunny Hei Wong DPhil, MRCP, FHKCP5,6 | Wing Tak Wong PhD7 | Yat Sun Chan FRCP, FACC6 | Martin C. S. Wong MPH, MBA, MD, FRACGP8 | Katharina Wassilew MD, DScmed9 | Vassilios S. Vassiliou MA, MBBS, MRCP, PhD, FHEA, FESC10,11 | Gary Tse MA, MPH, PhD, FESC, FACC, FRCP5,613

1Faculty of Medicine, Newcastle University, Newcastle, UK
2Second Department of Cardiology, Laboratory of Cardiac Electrophysiology, “Evangelismos” General Hospital of Athens, Athens, Greece
3Tianjin Key Laboratory of Ionic-Molecular Function of Cardiovascular disease, Department of Cardiology, Tianjin Institute of Cardiology, Second Hospital of Tianjin Medical University, Tianjin, China
4Department of Anaesthesia and Intensive Care, Faculty of Medicine, Chinese University of Hong Kong, Hong Kong, China
5Li Ka Shing Institute of Health Sciences, Faculty of Medicine, Chinese University of Hong Kong, Hong Kong, China
6Department of Medicine and Therapeutics, Faculty of Medicine, Chinese University of Hong Kong, Hong Kong, China
7School of Life Sciences, Chinese University of Hong Kong, Hong Kong, China
8The Jockey Club School of Public Health and Primary Care, Faculty of Medicine, The Chinese University of Hong Kong, Hong Kong, China
9Department of Pathology, Rigshospitalet, University Hospital of Copenhagen, Copenhagen, Denmark
10Norwich Medical School, University of East Anglia, Norwich, UK
11Royal Brompton Hospital and Imperial College London, London, UK

Correspondence
Gary Tse, Department of Medicine and Therapeutics, The Chinese University of Hong Kong, Hong Kong, China. Email: tseg@cuhk.edu.hk

Funding information
Croucher Foundation of Hong Kong

Abstract
Arrhythmogenic right ventricular cardiomyopathy/dysplasia (ARVC/D) is an inherited myocardial disease characterized by fibro-fatty replacement of the right ventricular myocardium, and associated with paroxysmal ventricular arrhythmias and sudden cardiac death (SCD). It is currently the second most common cause of SCD after hypertrophic cardiomyopathy in young people <35 years of age, causing up to 20% of deaths in this patient population. This condition has a male preponderance and is more commonly found in individuals of Italian and Greek descent. To date, there is no single diagnostic test for ARVC/D and the diagnosis is made based on clinical, electrocardiographic, and radiological findings according to the Revised 2010 Task Force Criteria. In this review, we will discuss the mainstay treatment which includes pharmacotherapy, implantable cardioverter-defibrillator insertion for abortion of sudden cardiac death, and in the advanced stages of the disease cardiac transplantation.

KEYWORDS
arrhythmogenic right ventricular cardiomyopathy, arrhythmogenic right ventricular dysplasia
INTRODUCTION

Arrhythmogenic right ventricular cardiomyopathy/dysplasia (ARVC/D) is an uncommon inherited cardiac disease characterized by progressive right ventricular (RV) dysfunction due to fibro-fatty replacement of the myocardium and associated with high risk of ventricular arrhythmias and sudden cardiac death (SCD). ARVC/D has a predominantly autosomal dominant inheritance, although recessive forms associated with a cutaneous phenotype, such as Naxos disease and Carvajal syndrome, are also observed. Despite RV abnormalities being the predominant finding, it has been recently appreciated that patients with ARVC/D may also have some degree of left ventricular (LV) involvement and indeed severe LV impairment can sometimes be the initial manifestation of the disorder. An LV-predominant form of ARVC/D has recently been described. Independently of which ventricle is initially or predominantly affected, during the later stages, advanced disease can result in biventricular heart failure, which may closely resemble dilated cardiomyopathy (DCM). LV dysfunction is observed more frequently with greater RV dysfunction, worse functional class, and leads to an increased tendency to cardiovascular adverse events relating to heart failure. The same study showed no clear relationship between LV involvement and an increased rate of arrhythmic events. Clinical manifestations vary with the age of the patient and stage of disease. In this article, we will review the pathophysiology of ARVC/D, the main diagnostic modalities used clinically aiding the diagnosis and patient management.

PATHOPHYSIOLOGY OF ARVC/D

The pathophysiological mechanisms in ARVC/D involve desmosomal abnormalities that can arise from mutations in cell adhesion proteins or intracellular signaling components. A number of genes have been implicated in the pathogenesis of ARVC/D, as illustrated in Table 1. Particularly, reduced cardiac desmoglein-2 and desmocollin-2 levels appear to be specifically associated with ARVC/D, independent of the gene mutations found. The desmosome normally maintains cell-to-cell adhesion and confers mechanical strength to tissues (Figures 1 and 2). In the extracellular space, desmosomal cadherins (desmocollin and desmoglein) bind strongly to each other. Cadherins span the plasma membrane and attach to linker proteins (plakoglobin, desmoplakin, and plakophilin-2) in the intracellular space. Plakoglobin and desmoplakin are intracellular proteins anchoring desmosomes to desmin intermediate filaments. Moreover, plakoglobin contributes to interlinking adherens junctions with the actin cytoskeleton and participates in cellular signaling to the nucleus and desmosome organization. Defects in linking sites of these proteins can interrupt cell adhesion, especially under conditions of increased mechanical stress or stretch, leading to cell death, progressive loss of myocardium, and fibro-fatty replacement. As such, surviving myocardial fibers within the fibro-fatty tissue from zones of slow conduction provide a medium for re-entry ventricular arrhythmias. The degeneration-inflammation model posits that the resulting cellular damage is found in tissues under high mechanical stress. Indeed, this notion is in keeping with the observations that exercise increases age-related penetrance and risk of arrhythmias in ARVC/D-associated mutation carriers. The potential role of calcium-sensitive pathways in the pro-arrhythmia mechanism of ARVC/D has been proposed. In a recent meta-analysis, however, the presence of desmosomal gene mutations was not associated with global or regional structural and functional alterations, epsilon wave, or VT of left bundle branch morphology.

CLINICAL PRESENTATION

Classically, ARVC/D usually presents between the second and fourth decades of life with syncope, symptomatic arrhythmias, or SCD. An example of monomorphic VT in a patient with ARVC/D is shown in Figure 3 (reproduced from with permission). Chest pain can be the presenting finding of the disorder. One-third of the patients become symptomatic before the 30th year of life. ARVC/D can lead to deleterious consequences, such as ventricular arrhythmias, pump failure, and death. Competitive sports have been associated with a twofold increased risk of ventricular arrhythmias and mortality, and earlier presentation of symptoms comparing with inactive patients and patients who participated in recreational sport. Another interesting finding is the relation between meteorological factors and outcomes in patients with ARVC/D. Particularly, higher temperature and larger variation in humidity within 3 days of events were independently associated with the development of ventricular arrhythmic...
and sudden mortality events. Intracardiac thrombosis may occur in certain patients with ARVC/D. Atrial arrhythmias are also common in ARVC/D and present at a younger age than in the general population. Atrial arrhythmias are associated with male gender, increasing age, and left atrial dilation and clinically important, and they are associated with inappropriate implantable cardioverter-defibrillator shocks and increased risk of both heart failure and death. In addition to tachy-arrhythmias, brady-arrhythmias are also observed in this condition.

4 | DIAGNOSING ARVC/D: THE CHALLENGES

There is no single diagnostic test for ARVC/D. The diagnosis is made based on major and minor clinical, electrical, and imaging criteria that have been devised by expert consensus of the Task Force Criteria (TFC) originally proposed in 1994 and further revised in 2010 (Table 2). In the original 1994 TFC, the clinical diagnosis was based strongly on symptomatic index cases and SCD victims—those with overt and severe phenotypes. Consequently, the 1994 criteria were highly specific, but they lacked sensitivity for early and familial disease. A systematic review and meta-analysis of five retrospective studies compared the diagnostic concordance between the 1994 and 2010 criteria. 8.6% and 3.6% satisfied the 1994 and 2010 major criteria, whereas 29.2% and 1.9% satisfied their minor criteria. Therefore, the 2010 revised TFC have resulted in a significant reduction in the number of patients that satisfy the cardiovascular magnetic resonance criteria and in a statistically significant increase in the number of patients diagnosed with definite ARVC/D compared to the 1994 criteria. Since then, modifications of the original criteria have been proposed to facilitate clinical diagnosis in

FIGURE 1 Desmosomes (arrows) connect cytoskeletons of adjacent cells (myofibrils, double arrows) and contain centrally dense lamellae composed of polypeptides, plakoglobin, and other proteins. Clusters of mitochondria (asterisks) can be seen

FIGURE 2 The desmosomes (ellipse) show dense cell borders and glycoproteins in the intercellular space. Plakoglobin is a plaque protein of intercellular junctions
first-degree relatives, who often have an incomplete disease phenotype. According to these recommendations, familial ARVC/D is said to occur when the following conditions are met: (i) T-wave inversion in the right precordial leads in individuals older than 14 years of age; (ii) late potentials by signal-averaged ECG (SAECG); and (iii) ventricular tachycardia with left bundle branch block morphology on the ECG or exercise testing or >200 premature ventricular contractions in 24 hours.

4.1 | Electrocardiography

In the ECG, epsilon waves, which are late potentials occurring between the end of the QRS complex and the onset of the T-wave, and T-wave inversion in the right precordial leads of V1 to V3 may be observed. Epsilon waves are specific for ARVC/D although it is only observed in 30% of patients and are best seen in the right precordial leads V1-V3 (Figure 4). Particularly, epsilon waves in lead aVR in patients with arrhythmogenic right ventricular cardiomyopathy are rare electrocardiographic findings with a specificity of 100%. The detection of epsilon waves on 12-lead ECG has been associated with higher episodes of sustained VT, but reassuringly, this did not lead to increased SCD incidence. A case of a child with extensive involvement of both right and left ventricular walls and epsilon waves in all precordial leads has been reported. However, interobserver variability in the assessment of epsilon waves is high. As a result, the assessment of the epsilon waves must be performed cautiously particularly in patients with who would not otherwise meet diagnostic criteria. The sensitivity of epsilon waves on the ECG is low between 25% and 38%, and therefore, a normal ECG does not exclude the diagnosis of ARVC/D. The use of Fontaine bipolar precordial lead electrocardiography (F-ECG) increased the sensitivity to 50%. Electrical abnormalities in ARVC/D are important as these changes precede structural changes.

A more specialized electrocardiographic technique, known as signal-averaged electrocardiography (SAECG), can also be utilized. It aims to filter interference, unmask any microvariations, and display late potentials (if any) within the QRS complex by averaging the multiple electrical signals generated by the heart. Late potentials are thought to represent electrical depolarization abnormalities and are defined within the minor depolarization criteria in the 2010 TFC (Table 2). The criteria include the following:

1. Filtered QRS duration ≥114 ms
2. Duration of terminal QRS <40 μV: ≥38 ms

FIGURE 3 An electrocardiogram (ECG) showing monomorphic ventricular tachycardia (VT) with a left bundle branch block pattern with superior axis from a patient with arrhythmogenic right ventricular cardiomyopathy/dysplasia (ARVC/D). Figure reproduced from ref. 23 with permission.
### TABLE 2 International Task Force Criteria modified in 2010 (reproduced from 31 with permission)

<table>
<thead>
<tr>
<th>Domain</th>
<th>Major criteria</th>
<th>Minor criteria</th>
</tr>
</thead>
<tbody>
<tr>
<td>Family history</td>
<td>• ARVD confirmed in a first-degree relative</td>
<td>• History of ARVD in a first-degree relative in whom it was not possible</td>
</tr>
<tr>
<td></td>
<td>• ARVD confirmed at surgery or autopsy in a first-degree relative</td>
<td>to determine whether the current Task Force Criteria is met.</td>
</tr>
<tr>
<td></td>
<td>• Pathogenetic mutation in a gene associated with ARVD</td>
<td>• Premature death at &lt;35 years of age due to suspected ARVD.</td>
</tr>
<tr>
<td></td>
<td>• History of ARVD confirmed pathologically or by current Task Force criteria in a second-degree relative</td>
<td>• ARVD confirmed pathologically or by current Task Force criteria in a second-degree relative</td>
</tr>
<tr>
<td>ECG abnormalities</td>
<td>• Epsilon wave (reproducible low-amplitude signals between end of QRS complex and beginning of T-wave in leads V1 to V3)</td>
<td>• Late potentials by signal-averaged ECG in ( \geq 1 ) of 3 parameters in an absence of QRS ( \geq 110 ) ms</td>
</tr>
<tr>
<td></td>
<td>• Inverted T-waves in leads V1 to V3 in individuals &gt;14 years of age in the absence of RBBB and QRS &gt;120 ms)</td>
<td>• Filtered QRS duration ( \geq 114 ) ms</td>
</tr>
<tr>
<td></td>
<td>• Duration of terminal QRS ( &lt;40 ) ( \mu )V and ( &gt;38 ) ms</td>
<td>• Root-mean-square voltage of terminal QRS ( &lt;40 ) ( \mu )V and ( &gt;20 ) ( \mu )V</td>
</tr>
<tr>
<td></td>
<td>• Terminal activation duration of QRS ( \geq 55 ) ms measured between ( S ) v the end of the QRS complex, including ( R' ), in V1, V2, or V3, without RBBB</td>
<td>• Terminal activation duration of QRS ( \geq 55 ) ms measured between the nadir of the ( S ) wave and the end of the QRS complex, including ( R' ), in V1, V2, or V3, without RBBB</td>
</tr>
<tr>
<td></td>
<td>• T-wave inversion in V1 and V2 in individuals &gt;14 years of age in an absence of RBBB, or in V4 to V6</td>
<td>• T-wave inversion in V1 and V2 in individuals &gt;14 years of age in an absence of RBBB, or in V4 to V6</td>
</tr>
<tr>
<td></td>
<td>• T-wave inversion in leads V1 to V4 in individuals &gt;14 years of age in the presence of complete RBBB</td>
<td>• T-wave inversion in leads V1 to V4 in individuals &gt;14 years of age in the presence of complete RBBB</td>
</tr>
<tr>
<td>Arrhythmias</td>
<td>• Nonsustained or sustained VT with a LBBB morphology</td>
<td>• Nonsustained or sustained VT of RVOT configuration, LBBB morphology with inferior axis (positive QRS in leads II, III, and aVF and negative in aVL) or of unknown axis</td>
</tr>
<tr>
<td></td>
<td>with a superior axis (negative or indeterminate QRS in leads II, III, and aVF and positive in aVL)</td>
<td>• 500 ventricular extrasystoles within 24 h on Holter monitoring</td>
</tr>
<tr>
<td>Tissue characteristics</td>
<td>• Fibro-fatty replacement of the myocardium on endomyocardial biopsy</td>
<td>• Residual myocytes ( &lt;60% ) by morphometric analysis (or ( &lt;50% ) if estimated) with fibrous replacement of the RV free wall myocardium in &gt;1 sample, with or without fatty replacement of tissue on endomyocardial biopsy</td>
</tr>
<tr>
<td></td>
<td>• Residual myocytes 60%-75% morphometric analysis (or 50%-65% if estimated) with fibrous replacement of the RV free wall myocardium in &gt;1 sample, with or without fatty replacement of tissue on endomyocardial biopsy</td>
<td>• Residual myocytes 60%-75% morphometric analysis (or 50%-65% if estimated) with fibrous replacement of the RV free wall myocardium in &gt;1 sample, with or without fatty replacement of tissue on endomyocardial biopsy</td>
</tr>
<tr>
<td>Global or regional functional or structural abnormalities</td>
<td>• Fractional area change ( &gt;33% )</td>
<td>• Regional RV akinesia, dyskinesia, and one of the following at end diastole:</td>
</tr>
<tr>
<td></td>
<td>• Regional RV akinesia, dyskinesia, or aneurysm and one of the following at end diastole:</td>
<td>• Regional RV akinesia, dyskinesia, and one of the following at end diastole:</td>
</tr>
<tr>
<td></td>
<td>• PLAX RVOT ( \geq 32 ) mm (corrected for body size [PLAX/BSA] ( \geq 19 ) mm/m(^2))</td>
<td>• PLAX RVOT ( \geq 32 ) and ( &lt;36 ) mm (corrected for body size [PLAX/BSA] ( &gt;18 ) and ( &lt;21 ) mm/m(^2))</td>
</tr>
<tr>
<td></td>
<td>• PLAX RVOT ( \geq 36 ) mm (corrected for body size [PLAX/BSA] ( \geq 21 ) mm/m(^2))</td>
<td>• PLAX RVOT ( \geq 29 ) and ( &lt;32 ) mm (corrected for body size [PLAX/BSA] ( &gt;16 ) and ( &lt;19 ) mm/m(^2))</td>
</tr>
<tr>
<td>Echocardiography</td>
<td>• Fractional area change ( \leq 33% )</td>
<td>• Fractional area change ( &gt;33% ) and ( \leq 40% )</td>
</tr>
<tr>
<td>MRI</td>
<td>• Regional RV akinesia or dyskinesia or dysynchronous RV contraction and one of the following:</td>
<td>• Regional RV akinesia or dyskinesia and one of the following:</td>
</tr>
<tr>
<td></td>
<td>• Ratio of RV end-diastolic volume to BSA ( \geq 100 ) mL/m(^2) (male) ( \geq 90 ) mL/m(^2) (female)</td>
<td>• Ratio of RV end-diastolic volume to BSA ( \geq 100 ) mL/m(^2) and ( &lt;110 ) mL/m(^2) (male) or ( &lt;90 ) mL/m(^2) and ( &lt;100 ) mL/m(^2) (female)</td>
</tr>
<tr>
<td></td>
<td>• RV ejection fraction ( \leq 40% )</td>
<td>• RV ejection fraction ( &gt;40% ) and ( \leq 45% )</td>
</tr>
<tr>
<td>RV angiography</td>
<td>• Regional RV akinesia, dyskinesia, or aneurysm</td>
<td>• Regional RV akinesia, dyskinesia, or aneurysm</td>
</tr>
</tbody>
</table>

3. Root-mean-square voltage of terminal 40 ms: \( \leq 20 \) \( \mu \)V

As long as any of the above is satisfied, the minor criterion in the 2010 TFC is fulfilled. Even though SAECG can be useful, it must be noted that an abnormal SAECG is not specific to ARVC as it can also be seen in other conditions such as myocarditis and scarring in ischemic heart disease.

4.2 Imaging: echocardiography and cardiac magnetic resonance (CMR)

Noninvasive imaging modalities such as echocardiography and CMR can be used to identify structural and functional abnormalities described in the revised TFC. It is also possible to investigate these invasively using RV angiography. Echocardiographic features of
ARVC/D include global ventricular dilatation (Figure 5), reduced RV ejection fraction with normal LV, or mild segmental dilatation of the RV or regional RV hypokinesis. CMR similarly demonstrates these abnormalities but at a higher resolution (Figure 6). There are several advantages of CMR over echocardiography. Firstly, it provides gold standard values for both LV and RV volumes. Secondly, it provides clear delineation of RV anatomy for micro-aneurysms and trabeculations. Thirdly, it can clearly visualize RV hypokinesis and akinesis. All of these features are part of the 2010 criteria, making CMR an excellent modality for diagnosis. Moreover, CMR can help to identify conditions that can mimic ARVC/D. For example, Khaji et al presented a patient with mega-epsilon wave, right ventricular dilatation, and inducible VT that was initially diagnosed as ARVC/D by the Task Force Criteria. Further examination with endomyocardial biopsy revealed the diagnosis of sarcoidosis. A study which compared clinical and electrophysiological parameters between patients with ARVC/D and cardiac sarcoidosis showed that patients with sarcoidosis had reduced left ventricular ejection fraction, a significantly wider QRS, right-sided apical VT, and more inducible forms of monomorphic VT. Furthermore, cardiovascular magnetic resonance and specifically the cardiac volume, in addition to the degree and location of cardiac involvement, can be used to distinguish between these two disease entities. The combined use of signal and wall motion parameters of cardiovascular magnetic resonance has been proposed to be contemporaneously considered for achieving a better diagnostic accuracy for the diagnosis of ARVC/D.

4.3 Genetic studies, biopsy, and histology

Genetic testing can identify desmosomal mutations in approximately 30%-60% of ARVC/D cases. Of these, PKP2 mutations are most frequently observed, with an estimated prevalence of 10%-47% among unrelated probands and 70%-82% among familiar ARVC/D cases. The second commonest mutated gene is desmoplakin (DSP). Endomyocardial biopsy guided by voltage mapping may provide ARVC/D diagnosis confirmation.

The main macroscopical features of ARVC/D are shown in a short-axis section of an explanted heart at the time of heart transplantation for ARVC/D (Figure 7). This demonstrates a grossly dilated right ventricle. The ventricular wall is widely replaced by fat tissue. There are prominent right ventricular trabeculations, and the muscular tissue of the numerous right ventricular trabecules appears to be preserved.

In terms of histology, the hallmark of the disease is fibro-fatty replacement of the right ventricular myocardium and right ventricular part of the interventricular septum. Figure 8 shows full-thickness sections of anterior right ventricular wall and interventricular septum submitted from the explanted heart shown in Figure 7. On hematoxylin and eosin stain (left), there is an abrupt transition between small areas of preserved myocardium and areas of transmural fibro-fatty replacement in the anterior right ventricular wall and right ventricular myocardium of interventricular septum. Collagen stains, such
as Sirius red (right), are useful to visualize areas of fibrosis in the RV wall.

Immunohistochemistry for plakoglobin is often equivocal and shows, as in this confirmed case of ARVC/D, a variegated staining pattern with focal normal staining reaction in section submitted from interventricular septum (IVS) (left), and a decreased staining reaction in a transmural tissue section from RV, which corresponds to an abrupt transition to an area of fibro-fatty replacement (right) (Figure 9).

4.4 | Differential diagnosis

It is important to differentiate ARVC/D from other right ventricular disorders, such as Brugada syndrome, as overlapping features may be found. Moreover, cardiovascular conditions such as peripartum cardiomyopathy or athlete’s heart can present with similar clinical and imaging findings. A correct diagnosis is important because unlike ARVC/D, athlete’s heart would not justify disqualification from competitive sports. Distinguishing between ARVC/D and athlete’s heart remains a diagnostic challenge. High-level endurance training is associated with RV elongation, dilation, and hence enlargement compared to isometric physical activities. As such, an enlarged RV dimension alone is not a reliable diagnostic criterion for ARVC/D in elite athletes. A large proportion of athletes also express echocardiographic morphological findings often evident in documented ARVC/D, including rounded RV apex, and prominent RV trabeculations and moderator band. By contrast, impaired RV systolic function is associated only with ARVC/D and not with athlete’s heart. The use of both RV dilation and systolic dysfunction might serve as a useful diagnostic tool to separate between the two. Despite all the imaging parameters, and new normal specific ranges for athletes, reaching a diagnosis of ARVC/D in an athlete can sometimes remain challenging and a short period of detraining with subsequent assessment usually with cardiac MRI can be helpful in resolving this ambiguity.

5 | MANAGEMENT

In ARVC/D, the main goal is to avoid the high-risk events of malignant arrhythmias and SCD and slow the progression of heart failure. Competitive sports are discouraged. While patients with ARVC are allowed to perform exercise including sports as part of a healthy lifestyle, they should not exercise to maximal capacity and be vigilant to any symptoms of palpitations. Frequent endurance exercise increases the risk for VT/VF and heart failure. The management of patients with ARVC/D in specific situations such as pregnancy is beyond the scope of this review, but is directed to the following reference. Anti-arrhythmic medications, such as
beta-blockers and class-III agents, are advised. Sotalol and amiodarone with or without the need for conventional beta-blockers are potent (effective is used in 3 sentences in a row). Calcium channel blockers may be effective in selected patients. The addition of flecainide in combination with sotalol/metoprolol may be an adequate strategy for the control of ventricular arrhythmias in patients with ARVC/D refractory to single-agent therapy and/or catheter ablation. In general, the most sufficient combinations appear to be sotalol or flecainide and amiodarone/beta-blockers.

The American College of Cardiology, the American Heart Association and the European Society and Cardiology recommended ICD implantation for the prevention of SCD events. Risk stratification and indication to ICD implantation in ARVC/D has been proposed by an international task force consensus statement. One study reported that the annual cardiac mortality in patients with ARVC/D who were implanted with an ICD was 0.9%. Finally, VT ablation targeting late potentials abolition seems to be effective in preventing VT recurrence in patients with or without RV structural abnormalities. In the multicenter registry, clinical response (freedom from SCD, VT requiring hospitalization, or heart transplantation) after the last ablation (predominantly endocardial) was 86% at 1 year, 69% at 5 years, and 60% at 10 years. On the other hand, the combined endocardial and epicardial approach resulted in better procedural success and long-term VT-free survival compared with the endocardial approach in ARVC/D patients with recurrent VTs. In fact, the combined endocardial and epicardial VT ablation eliminated all clinical and induced VTs, and the addition of scar dechanneling resulted in noninducibility in all cases. Identification of conducting channels (CCs) inside or between the scars can be achieved via endocardial high-density substrate mapping.

However, another single-center study showed that the vast majority of critical VT circuits were epicardial while epicardial ablation of VT appeared to be both safe and effective in achieving arrhythmia control in ARVC/D. A recent meta-analysis showed better outcomes with the combined endocardial and epicardial ablation approach compared with the endocardial approach only. Furthermore, an inducibility-guided catheter ablation strategy of VT in patients with ARVC/D has been proposed to prevent unnecessary epicardial ablation procedures. Electrical regression of SAECG after catheter ablation in ARVC/D has been found to be associated with fewer ventricular arrhythmia recurrences. Other treatment options like bilateral cardiac sympathectomy need to be studied further in order to investigate its optimal timing and use in ARVC/D management.
Individuals who present with congestive heart failure are managed with diuretics and angiotensin-converting enzyme (ACE) inhibitors or aldosterone inhibitors, with heart transplantation considered in terminal stages of the disease. Anticoagulation may be used in ARVC/D patients with large, hypokinetic RV and slow blood flow because of the risk of thrombosis. Management of family members of patients with ARVC/D is complex due to the incomplete penetrance and variable expressivity nature of the disease. This is beyond the scope of this review, and the reader is referred to the article here.

6 | PROGNOSTICATION

There are many factors that have been associated with adverse outcomes in patients with ARVC/D. These include clinical characteristics such as male gender, heart rate variability, and atrial arrhythmias. Other important factors include electrocardiographic characteristics (atrioventricular block, T-wave inversions), echocardiographic parameters (RV diameter), and genetic factors (PKP2 carriers), all of which are associated with higher likelihood of ventricular arrhythmias, heart failure, and death. Biomarkers such as soluble ST2 have been associated with RV and LV functions in patients with ARVC/D and may aid in the determination of disease severity in this disease.

7 | CONCLUSION

ARVC/D is an inherited disease characterized by fibro-fatty replacement of the right ventricular myocardium, which significantly increases the risk of paroxysmal ventricular arrhythmias and SCD. Diagnosis is based on the 2010 modified Task Force criteria, requiring clinical and family history, electrocardiography, and imaging. Diagnosis may be confirmed by endomyocardial biopsy, if the tissue is harvested from right ventricular wall or right interventricular septum and not from prominent right ventricular trabeculae. A further personalized approach in the management of the patients should be undertaken by specialists in high-volume centers to enable practice of evidence-based medicine in this condition with high morbidity and mortality.

ACKNOWLEDGEMENTS

Figures 1 and 2 provided courtesy of Dr. med. Georgi Wassilew of the Department of Pathology, Humboldt University, Berlin, Germany. GT and SW were supported by the Croucher Foundation of Hong Kong.

CONFLICT OF INTERESTS

Authors declare no conflict of interests for this article.

ORCID

George Bazoukis http://orcid.org/0000-0003-1009-9772
Gary Tse http://orcid.org/0000-0001-5510-1253

REFERENCES


