Arrhythmias Associated with Inflammatory Cardiomyopathies

Roshan Karki, MBBS
Chaitra Janga, MBBS
Abhishek J. Deshmukh, MBBS

Address
Division of Cardiovascular Disease, Mayo Clinic, 200 1st Street, Rochester, MN, 55905, USA
Email: Deshmukh.Abhishek@mayo.edu

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Keywords
Inflammatory cardiomyopathy · Arrhythmias · Cardiac sarcoidosis

Abbreviations
AIC Arrhythmogenic inflammatory cardiomyopathy · CS Cardiac sarcoidosis · GCM Giant-cell myocarditis

Abstract
Purpose of review To provide an approach to the diagnosis and treatment of arrhythmias associated with inflammatory cardiomyopathies.

Recent findings Inflammatory cardiomyopathies are increasingly recognized as the etiology of both ventricular and supraventricular arrhythmias. There have been recent studies providing novel insights into the pathogenesis of arrhythmias in inflammatory cardiomyopathies and exploring the role of various diagnostic tools and treatment strategies.

Summary Patients with inflammatory cardiomyopathies often present with one or more arrhythmias, including atrioventricular block, atrial and ventricular tachyarrhythmias, and occasionally sudden cardiac death. Given dynamic pathophysiology and heterogeneous presentation, the management of arrhythmias in these patients presents unique challenges. We review the current approach to the diagnosis and treatment of arrhythmias in this challenging cohort of patients with an emphasis on cardiac sarcoidosis.
Introduction

Inflammatory cardiomyopathies (ICMs) are a spectrum of disorders characterized by acute or chronic inflammatory processes affecting the myocardium (myocarditis) and associated with mechanical cardiac dysfunction with altered hemodynamics, conduction disturbances, or electrical instability. Inflammatory cardiomyopathies are mostly caused by an infection or autoimmune disease that has been summarized in Table 1. The patients with ICMs frequently present with arrhythmias. However, a subset of these patients may present first with arrhythmias and even sudden cardiac death, and are later diagnosed as ICMs [1]. Bauer and colleagues [2] proposed an operational diagnosis of “arrhythmogenic inflammatory cardiomyopathy” for these patients until a specific etiologic diagnosis is made. In the current review, we discuss an approach to the diagnosis and treatment of arrhythmias associated with ICMs with an emphasis on cardiac sarcoidosis (CS).

Epidemiology

Although the global annual incidence of reported myocarditis is 22 per 100,000 [3], the actual burden of arrhythmias associated with ICM is unknown. In an autopsy study (n = 753, age < 50 years) of sudden cardiac deaths (SCDs), the prevalence of myocarditis was 6% [4]. Tung et al. demonstrated a 49% prevalence of occult myocardial inflammation by fluorodeoxyglucose positron emission tomography (FDG-PET) in selected patients referred for ventricular

| Inflammatory cardiomyopathy | Associated arrhythmias |
|-----------------------------|------------------------|
| Autoimmune                  |                        |
| Cardiac sarcoidosis         | AV block (42%); PVCs, NSVTs, and VTs (28%); supraventricular arrhythmias (15–30%, AF most common) |
| Scleroderma                 | PVCs (20–67%), NSVT (7–13%), SCD (21%); supraventricular arrhythmias (21–61%) |
| Rheumatoid arthritis        | AF (0.8–18.3%)         |
| Ankylosing spondylitis      | PACs (2.2–94%); PVC (28.6–55%) |
| Psoriasis                   | AF (2.5–7.1%)          |
| Infectious                  |                        |
| Chagas disease              | LAFB (22.5%), RBBB (22.7%), AV block (2.6%); AF (5.4%); PVCs (5.4%), VT (40%) |
| Lyme carditis               | AV block               |
| Leptospirosis               | AV block               |
| Coxsackie B myocarditis     | Sinus arrest, AV block, PAC, PVC, VT |
| Influenza                   | AV block (12%); AF (5%), VT (20%) |
tachycardia (VT) ablation [5]. Clinically manifest CS occurs in 5% of patients with sarcoidosis. Studies have shown a prevalence of CS to be 6–28% in patients presenting with nonischemic VTs [6, 7], and about 17% of CS patients present as sudden cardiac arrest [8]. Although giant-cell myocarditis (GCM) is rare, it is an important cause of ICM that can be rapidly fatal. The GCM patients often present with AV block (5–27%) and ventricular arrhythmias (14–22%), and about 55% of the patients develop life-threatening ventricular arrhythmias at the end of 5 years [9, 10].

Etiology of inflammatory cardiomyopathy

Inflammatory cardiomyopathies are caused mostly by (1) infections, (2) autoimmune diseases, and (3) drugs but may be idiopathic. The infectious agents include viruses (coxsackievirus, adenovirus, parvovirus B19, influenza, HIV, hepatitis B, or hepatitis C), bacteria (Borrelia burgdorferi, meningococci, streptococci, psittacosis, diphtheria), rickettsia (typhus, Rocky Mountain spotted fever), fungi (aspergillosis, candidiasis), and parasites (Chagas disease, toxoplasmosis). The Sars-Cov-2 infection responsible for the ongoing COVID-19 pandemic has also been implicated as a cause of acute myocarditis [11]. The rheumatological conditions associated with inflammatory cardiomyopathies include CS, GCM, systemic lupus erythematosus, rheumatoid arthritis, systemic sclerosis, celiac disease, and inflammatory bowel disease [12••].

Pathophysiology of arrhythmias in inflammatory cardiomyopathy

The mechanism of the arrhythmias in ICM depends upon the stages of the inflammation. The primary substrate for the genesis of reentrant ventricular arrhythmias in ICM is tissue heterogeneity resulting from patchy interstitial or replacement fibrosis in the healed phase. However, the slow zones that promote reentry may occur during the stage of active inflammation due to electrical remodeling. In the inflammation-fibrosis continuum, active inflammation results in arrhythmias due to automaticity or triggered activity attributed to various mechanisms, including direct myocardial cell membrane injury, microvascular ischemia (Parvovirus B19), and gap junction dysfunction [13]. Recently, the role of autoimmune channelopathies in the genesis of arrhythmias has also been recognized [14•]. Ongoing active inflammation can also cause polymorphic VT or ventricular fibrillation (VF) akin to acute ischemia. In CS patients, Purkinje-related ventricular arrhythmias have also been described, but their exact mechanisms remain unknown [15]. Interestingly, fibrotic remodeling can also lead to automatic tachycardias by differentiation of fibroblasts into myofibroblasts, which then cause depolarization of electrotonically coupled cardiomyocytes [16]. This process is primarily driven by transforming growth factor-β1 (TGF-β1) [16] and is being explored as a therapeutic target to reduce arrhythmias [17].

Although atrial arrhythmias are more common in CS patients who have late gadolinium enhancement and those with left atrial enlargement, these patients often lack echocardiographic evidence of left ventricular diastolic dysfunction
or increased left atrial filling pressures [18]. Therefore, direct atrial involvement by granulomatous inflammation and resulting scarring is proposed as a putative mechanism [18]. However, subclinical left ventricular diastolic dysfunction cannot be confidently ruled out, and scarring/inflammation of the atria is challenging to study with currently available imaging technologies. Therefore, the mechanisms of atrial arrhythmias in CS remain to be elucidated.

In ICMs associated with bradycardia, such as CS and GCM myocarditis, there is a predilection for involvement of the conduction system in the basal interventricular septum. The AV block may be due to inflammation, direct mechanical compression (with noncaseating granuloma or giant cells), or fibrosis, or both, resulting in variable response to steroids [19].

Clinical presentation

Inflammatory cardiomyopathies have a myriad of clinical presentations, from asymptomatic left ventricular systolic dysfunction or mild heart failure to fulminant myocarditis. The patients often present with arrhythmias incident on a previously diagnosed ICM or as the first manifestation. There may or may not be extra-cardiac manifestations at the time of presentation. In addition to symptoms typical of heart failure and arrhythmias, many patients may have subtle symptoms such as fatigue. The common arrhythmic presentations of various ICMs are summarized in Table 1 and discussed briefly in the following paragraphs. The clinical spectrum of CS is summarized in Fig. 1.

![Clinical spectrum of cardiac sarcoidosis from inflammatory to “burnt out” phase with diagnosis based on resting myocardial perfusion SPECT fused with cardiac MRI (left), FDG-PET fused with cardiac MRI (middle), and gadolinium-enhanced cardiac MRI. Adapted with permission from Kouranos et al. [20].](image-url)

**Fig. 1.** Clinical spectrum of cardiac sarcoidosis from inflammatory to “burnt out” phase with diagnosis based on resting myocardial perfusion SPECT fused with cardiac MRI (left), FDG-PET fused with cardiac MRI (middle), and gadolinium-enhanced cardiac MRI. Adapted with permission from Kouranos et al. [20].
Bradyarrhythmias

For CS, high-grade atrioventricular (AV) block is one of the most common initial presentations in about 42% of patients [8]. In patients < 60 years of age presenting with AV block, CS should be suspected [18, 21]. Other potential differentials include Lyme carditis and giant-cell myocarditis [21, 22]. The CS patients with high-degree AV block are at risk of sudden cardiac death (9–34% in 5 years) with or without either of left ventricular systolic dysfunction or ventricular arrhythmias [23].

Ventricular arrhythmias and sudden cardiac death

The patients with acute myocarditis often present with frequent premature ventricular complexes, irregular and polymorphic VT, or ventricular fibrillation [24•]. In contrast, the patients with prior or healed acute myocarditis tend to present with monomorphic and regular ventricular tachycardia [24•]. The patients with CS often have frequent premature ventricular complexes, nonsustained VTs, or sustained VTs. Monomorphic ventricular tachycardia may be an initial presentation of CS with one study showing a prevalence of 28% in this population [6]. However, the prevalence of CS was reported to be much lower at 5% in another study (n = 435) of patients with nonischemic ventricular tachycardia referred for ablation [7]. The CS patients were noted to have multiple VT circuits (resulting in pleomorphism) and preferential involvement of the interventricular septum and the right ventricle.

Inflammatory cardiomyopathies have been implicated as the cause of sudden cardiac death, especially of young and middle-aged patients [8, 25, 26]. In CS, sudden cardiac arrest may be an initial presentation in about one-seventh of the patients [8]. In an autopsy study of cardiac sarcoidosis patients who had SCD, the predilection for septal or subepicardial involvement was demonstrated [27].

Supraventricular arrhythmias

Supraventricular arrhythmias are often seen in CS patients (15–32%) and predominantly include atrial arrhythmias—atrial fibrillation (most common), atrial tachycardia, and atrial flutter—despite their relatively young age [28, 29]. Interestingly, atrial arrhythmias are more common than the ventricular arrhythmias and often occur in those with left atrial enlargement or presence of late gadolinium enhancement of ventricle(s) on cardiac MRI [28, 29].

Diagnosis

Focused history to look for an infectious or rheumatic etiology is valuable, but many patients may only have cardiac involvement, making the diagnosis challenging. Family history is paramount as myocarditis may be a red herring for, or arguably feature of, inherited arrhythmogenic cardiomyopathies. In patients with suspected ICM, a constellation of history, imaging studies, and invasive biopsy is required to make a diagnosis. In the 2014 Heart Rhythm Society consensus statement, criteria for the diagnosis of CS (summarized in Table 2) were proposed.
Laboratory studies

There are serologic tests specific to infections (e.g., Lyme carditis) or inflammatory conditions (e.g., lupus) that can be performed in suspected cases. Acute phase reactants such as sedimentation rate and C-reactive protein may be elevated but are nonspecific. Biomarkers may provide both diagnostic and prognostic information. For example, an elevated B-type natriuretic peptide identifies CS in patients with sarcoidosis with greater accuracy (AUC 0.85) compared to angiotensin-converting enzyme or soluble IL-2, while cardiac troponin I is the strongest predictor of fatal arrhythmias [30].

Echocardiography

Echocardiography remains an essential initial study for the assessment of cardiac chamber size and function. The patients tend to have global systolic dysfunction or regional wall motion abnormalities in noncoronary distribution. Although typically characterized by basal septal thinning, some CS patients may have septal hypertrophy in the inflammatory phase of the disease [31].

CMR and FDG-PET

Late gadolinium enhancement is seen in patients with inflammatory cardiomyopathies, including CS, but the pattern of distribution is nonspecific for diagnosis. CS patients tend to have subepicardial or mid-myocardial involvement. In the early inflammatory phase, late gadolinium enhancement may not be present. The 2009 Lake Louise criteria may be utilized to diagnose myocarditis. However, comprehensive CMR with the mapping of T1 and T2 relaxation times has superior accuracy in acute and chronic myocarditis, respectively [32].
In an adequately prepared patient, myocardial uptake of $^{18}$FDG on PET would indicate the presence of active inflammation. It is important to discuss with the radiologist or imaging cardiologist the sarcoid-specific protocol for preparation of the patients for FDG-PET to avoid false-positive results from physiological uptake (Supplemental Fig. 1). Although FDG-PET likely has utility in most inflammatory cardiomyopathies, it has been extensively studied in CS patients. The focal or focal-on-diffuse pattern would suggest CS as a diagnosis and would also serve as a guide to anti-inflammatory therapy. Therefore, FDG-PET and CMR have complementary roles in the diagnosis of CS patients in different stages (Fig. 1) and their downstream management (Fig. 2) [33]. In addition to cardiac PET, whole-body FDG-PET is useful for the diagnosis of extra-cardiac involvement that could potentially be an easily accessible site for biopsy.

## Endomyocardial biopsy

Because of patchy, preferentially mid-myocardial or subepicardial, and left ventricular involvement, an endomyocardial biopsy has limited sensitivity to justify its routine use for the diagnosis of CS or other inflammatory cardiomyopathies [34]. Therefore, an extra-cardiac biopsy is often performed to support the diagnosis in the presence of other findings suggesting cardiac involvement [35]. The diagnostic yield of the endomyocardial biopsy may be increased by left ventricular or biventricular biopsy [36]. There has been increasing experience with abnormal or low-voltage electrogram-guided biopsy (Fig. 3) to increase the diagnostic yield [37]. According to 2007 ACC/AHA/ESC scientific statement, endomyocardial biopsy is recommended (class I) in patients with

![Fig. 2. FDG-PET of patients with cardiac and extra-cardiac sarcoidosis before (a) and after (b) treatment with anti-inflammatory agents (courtesy: Lori A. Blauwet, MD).](image)
new-onset heart failure (2 weeks to 3 months) with new ventricular arrhythmias, 2nd or 3rd degree AV block, or refractory to standard treatment within 1–2 weeks [38]. The primary aim of this recommendation is to identify giant-cell myocarditis that is rare but can be rapidly fatal without prompt and often advanced therapies.

Management

The treatment strategy depends upon the patient’s clinical presentation and stage of the disease process. The principle of management of CS patients is summarized in Fig. 4 but applies to other ICMs as well.

Medical therapy

All patients who have left ventricular systolic dysfunction with or without clinical heart failure are treated with guideline-directed medical therapy. It is reasonable to consider antiarrhythmic therapy as the first-line management of atrial and ventricular arrhythmias in both the inflammatory and the “burnt out” phases of the disease. The data to guide the choice of an antiarrhythmic drug is limited. While class III drugs such as sotalol and amiodarone are often used initially, concurrent pulmonary involvement should be taken into consideration.

Fig. 3. Endomyocardial biopsy of cardiac sarcoidosis. a Traditional endomyocardial biopsy with a bioptome. b Noncaseating granuloma suggesting a diagnosis of cardiac sarcoidosis. c The abnormal electrogram (right) is lower in voltages and fractionated. d, e RAO and LAO views of EGM-guided biopsy (red arrow, mapping catheter; red triangle, bioptome; star, intracardiac echocardiography). f Voltage map of patient with sarcoidosis created during biopsy with blue points representing abnormal bipolar signals. d, e Reproduced with permission from Liang et al. [37].
It is intuitive to consider anti-inflammatory therapy in patients who demonstrate active inflammation by FDG-PET; its usefulness for the treatment of arrhythmias remains uncertain. In a meta-analysis by Sadek [40], AV conduction improved in 47% of patients who received corticosteroid but failed to improve in any patients that did not receive corticosteroid. However, the influence of corticosteroids on the outcome of left ventricular systolic function or ventricular arrhythmias could not be conclusively determined. Some patients treated with corticosteroids may develop electrical storms, mostly within the first year of therapy [41]. Once corticosteroid is started, the CS patients are followed clinically and with FDG-PET at 3–6 months. If there is an improvement of FDG uptake, steroid-sparing agents such as methotrexate should be considered, and corticosteroids are tapered off gradually over the next 6–9 months. The role of other steroid-sparing agents (e.g., mycophenolate mofetil or azathioprine) is less well-defined, and the utility of biologic agents is unknown.

**Device therapy and risk stratification for SCD**

Although SCD remains the most feared outcome of inflammatory cardiomyopathies, the tools to risk-stratify these patients are limited. Similar to other cardiomyopathies, ICD is implanted in CS patients who have sustained ventricular arrhythmias (secondary prevention) or those who have LVEF ≤ 35% despite optimal medical therapy, including immunosuppressive therapy if FDG positive (primary prevention). In patients with myocarditis, the presence of late gadolinium enhancement on CMR or endocardial bipolar low-voltage areas on

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**Fig. 4.** The 2017 guidelines for ICD for prevention of SCD in patients with cardiac sarcoidosis. Reprinted with permission from Al-Khatib et al. [39].
electro-anatomical mapping has been associated with increased arrhythmic risk [42–44]. Cardiac pacing is indicated in CS patients with high-degree AV block, as it does not improve with corticosteroid therapy in about half of the patients [40]. Irrespective of left ventricular systolic dysfunction or ventricular arrhythmias, these patients are at increased risk of SCD. Programmed ventricular stimulation has been demonstrated to predict arrhythmic events in CS patients [45, 46]. Therefore, it is also reasonable to implant ICD in the presence of (1) syncope or near-syncope concerning for arrhythmic etiology on CMR, (2) sustained ventricular arrhythmia inducible with programmed electrical stimulation, or (3) indication for cardiac pacing [35]. Given a high risk of SCD in CS patients with mildly reduced ejection fraction (LVEF > 35% and < 50% or RVEF < 40%), ICD may be considered especially in the presence of LGE or sustained inducible VT on programmed ventricular stimulation (Fig. 3). In the presence of LGE and inducible VT, ICD may be useful in patients with preserved systolic function.

There is limited evidence of the utility of cardiac resynchronization therapy in CS patients. One study (n = 18) showed poor reverse remodeling and increased ventricular arrhythmias [47], and a second study (n = 19) showed improvement of LVEF, but there was no improvement of LV dimensions, mitral regurgitation, or RV systolic function when compared with other nonischemic cardiomyopathies [48]. Until we have more evidence, it is reasonable to implant a biventricular device or upgrade a previously implanted device in line with the current heart failure guidelines.

**Catheter ablation**

Although CS confers a substrate for reentrant VT, catheter ablation can be challenging due to the complexity of substrates with multiple circuits and subepicardial locations likely requiring an epicardial ablation [49–52]. Besides, catheter ablation has been shown to have a lower likelihood for acute success and a higher recurrence rate if performed in the inflammatory phase compared to the healed phase [50•]. Therefore, catheter ablation is reserved for patients with ventricular arrhythmias refractory to immunosuppressive and antiarrhythmic therapies, sometimes early in the course for management of incessant ventricular tachycardia or electrical storm. For CS patients with atrial arrhythmias, catheter ablation has been demonstrated to be feasible with reasonable efficacy and safety [53].

**Treatment of refractory ventricular tachycardia**

The patients with ventricular tachycardia refractory to antiarrhythmic therapy and catheter ablation may require cardiac sympathectomy as adjunctive therapy, failing which a mechanical circulatory device such as a left ventricular assist device and cardiac transplantation will need to be considered [7].

**Multidisciplinary team approach**

Given the heterogeneity and complexity of inflammatory cardiomyopathies and associated arrhythmias, the key to an effective acute and long-term management is a multidisciplinary team including cardiac electrophysiologists, heart failure cardiologists, imaging cardiologists, pulmonologists, and rheumatologists (Fig. 5).
**Future directions**

Much remains to be known about pathophysiology, diagnosis, and treatment of arrhythmias associated with ICMs, including CS. For the more streamlined treatment of CS patients, there is a need for randomized controlled trials to determine the outcomes of immune suppression, antiarrhythmics, catheter ablation, and device therapy. The Cardiac Sarcoidosis Multi-Center Randomized Controlled Trial (CHASM CS-RCT, NCT03593759) is seeking to determine the noninferiority in the efficacy of low-dose prednisone and methotrexate when compared with standard-dose prednisone, in addition to the superiority in quality of life measures [54]. The Interleukin-1 Blockade for Treatment of Cardiac Sarcoidosis Trial (MAGiC-ART, NCT04017936) will study the effect of anakinra on cardiac inflammation and fibrosis.

**Conclusions**

Inflammatory cardiomyopathies often have an arrhythmic presentation, with AV block, atrial arrhythmias, or ventricular arrhythmias, including SCD. Given their heterogeneous nature, accurate diagnosis and risk stratification can be challenging. A multidisciplinary approach is important to guide effective treatment.
Supplementary Information

The online version of this article (https://doi.org/10.1007/s11936-020-00871-5) contains supplementary material, which is available to authorized users.

Compliance with Ethical Standards

Conflict of interest
The authors has no conflicts of interest to declare.

Animal rights and informed consent statement
This article does not contain any studies with human or animal subjects performed by any of the authors.

Code availability
Not applicable

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