Infiltrative cardiomyopathy: A review of literature on clinical status and meta-analysis of diagnostic and clinical management methods

Aref Albakri*
St-Marien hospital Bonn Venusberg, Department of internal medicine, Bonn, Germany

Abstract

Infiltrative cardiomyopathy (ICM) refers to a heterogeneous group of myocardial disorders mainly cardiac amyloidosis, sarcoidosis, and hemochromatosis with characteristic deposition of abnormal substances within cellular and intercellular spaces, which leads to the development of ventricular diastolic dysfunction, systolic dysfunction or both. ICM results from a broader spectrum of inherited and acquired cardiac or systemic disorders. Despite being relatively rare, ICM is often misdiagnosed due to variable physiologic and morphologic characteristics. Early diagnosis of ICM can lead to potentially curative treatment. This paper reviews published studies on ICM, with a particular focus on its definitions, epidemiology, risk factors, clinical presentation, prognosis, etiopathogenesis, diagnosis, as well as clinical management. The ultimate aim of the review is to provide an in-depth understanding of ICM as well as improve its diagnostic and clinical management.

Introduction

History and clinical definition

Cardiomyopathies define abnormalities in the structure and functions of the ventricular myocardium in the absence of coronary artery diseases (CAD) or abnormal loading conditions [1-3]. The European Society of Cardiology (ESC) [4] classifies cardiomyopathies into primary or secondary myocardial disorders. Primary cardiomyopathies manifests as a cardiac or systemic disorder [3]. The ESC’s classification system defines cardiomyopathies using specific morphological and functional criteria and then classifying them into genetic and non-genetic subtypes, regardless of the existence of extra-cardiac disease [3]. In this regard, ICM has been defined as a form of cardiac muscle disease where the ventricular walls become progressively and excessively rigid, disrupting ventricular filling [2]. It is characterized by diastolic dysfunction (a reduction in diastolic volume of the ventricles) and systolic dysfunction in some cases. ICM can be idiopathic or associated with other diseases such as fibrosis, endomyocardial, or amyloidosis, resulting to interstitial fibrosis [2].

Forms of infiltrative cardiomyopathy

Infiltrative cardiomyopathies, generally refers to a group of diseases, which are characterized by restrictive cardiac physiology [5]. Sarcoidosis, haemochromatosis, and cardiac amyloidosis are the three most prevalent forms of ICM [6]. Other forms of ICM include Fabry disease, Friedrich ataxia, Danon disease and Mucopoly saccharidoses [1].

Epidemiology

Despite the preservation of EF, ICM patients exhibit severe limitations in exercise tolerance [5]. ICM are rare and thus often underdiagnosed [6]. In addition, their prevalence are incompletely documented and understood [2]. However, several clinical studies have been carried out to investigate the incidence and prevalence of various forms of ICM. Crooks [7] investigated 501 people with haemochromatosis and found the incidence of diagnosis of the condition increased by 2-fold between 1987 and 2002. Clinical evidence of heart-related sarcoidosis has been reported in at least 2-7% of patients diagnosed with sarcoidosis [8]. The prevalence of 2-7% have been reported in a number of studies [9-11]. Sarcoidosis can affect any part of the heart, with protein manifestations [8]. While most cardiac sarcoidosis-related deaths are because of arrhythmias or conduction defects, Lynch [8] contend that granulomatous infiltration of the myocardium can prove lethal. Antemortem recognition of cardiac sarcoidosis is rare due to high likelihood of sudden death as the presenting feature [12, 13]. Hamzeh [14] underscored the lack of a universally agreed diagnosis criteria of cardiac sarcoidosis. In their 2002 seminal paper, Deng [15] undertook an in-depth review of cardiac sarcoidosis, during which they discussed novel prevalence and epidemiological insights of cardiac sarcoidosis that have developed over the years.

Incidence of cardiac involvement has been reported in 19.5 to 28% of patients diagnosed with sarcoidosis in the U.S. and UK [12, 16, 17]. However, necropsy studies have reported a higher incidence of cardiac sarcoidosis (50-78%) among Japanese patients [18]. Cardiac involvement is the leading cause of death (accounting for 77-85% of total deaths) among patients diagnosed with sarcoidosis [19,20]. However, in the U.S., myocardial involvement is responsible for 13-50% of sarcoidosis-related deaths [12,21]. Cardiac sarcoidosis can manifest itself in various forms, including as a total heart block, congestive HF, pulmonary hypertension, ventricular arrhythmias, and ventricular aneurysms [22]. In Figure 1, cardiac amyloid is characterized by an increased LV wall thickness.

*Correspondence to: Aref Albakri, St-Marien hospital Bonn Venusberg, Department of internal medicine, Bonn, Germany, E-mail: arefalbakri@yahoo.com

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Risk factors

Given its underdeveloped diagnostic framework, the risk factors for the development of ICM is quite underreported. However, ICM is more associated with advanced age, female gender, and co-morbidities such as diabetes, hypertension, chronic kidney disease, and obesity [23,24]. ICM can result from a broader spectrum of both hereditary and acquired conditions with varying systemic manifestations [25]. Other forms of ICM like the Fabry disease caused by lack of lysosomal enzyme, α-galactosidase A, plays an essential role in breaking down neutral glycosphingolipids [26].

Clinical presentation

ICM presents itself with an occasional dynamic LV outflow, which appears in a similar manner to conditions with true myocyte hypertrophy such as hypertensive heart disease and hypertrophic cardiomyopathy [2]. In addition, some forms of ICM leads to increased wall thickness and small ventricular volume [2]. While some ICM lead to increased ventricular wall thickness, others result into chamber enlargement with secondary wall thinning [2].

There is no correlation between increased wall thickness and an increase in QRS complex amplitude. QRS complex amplitude may decrease with the increase in wall thickness that is often witnessed with accumulation within the interstitium as opposed to within myocytes [2]. The development of certain forms of ICM such as cardiac amyloid and Friedrich ataxia could lead to the absence of increased voltage on ECG, notwithstanding the appearance of hypertrophy [2]. However, research, as summarized in Table 1, has shown that low-voltage QRS complex phenomenon is not uniform across the various forms of ICM. For instance, the findings presented in Table 1 suggest that some forms of ICM such as Fabry disease and Fabry disease are characterized by increased-voltage QRS complex, which is often evident in circumstances of increased size of cardiac myocytes, normal intraventricular conditions, as well as synchronous activation within the myocardium.

Prognosis

Infiltrative cardiomyopathy generally refers to a group of diseases that are characterized by restrictive cardiac physiology [5]. ICM is among the severest forms of preserved ejection fraction. There is no standard prognosis for ICM, as a condition, given that each form of ICM has its own prognosis. This section therefore reviews the prognosis for various forms of ICM [5].

Cardiac amyloid

Cardiac amyloid characteristic echocardiographic appearance involves increased thickening of the LV and right ventricular walls, normal LV cavity size, as well as a nonspecific granular appearance of the myocardium, as shown in Figure 2. The cardiac amyloid is also characterized by atrial enlargement, thickened papillary muscles, small to moderate pericardial effusion, as well as valve leaflets [2].

Kyle [46] study to establish long-term survival (≥ 10 years) among 30 patients diagnosed with primary cardiac amyloid found patients who remain untreated have a median survival of less than six months after the onset of heart failure. Whereas post chemotherapy stem cell transplantation yields impressive results [47], the mortality rate associated with transplantation ranges between 11% and 40% [48].

Fabry disease

Cardiac involvement in Fabry disease, which is mostly associated with men in their thirties, may mimic the morphological and clinical features associated with HCM, including extremely thick walls. This condition accounts for 3% of cases that have been diagnosed as HCM [49]. A study conducted by Goldman [26] found that the thickening of the LV and LV mass increase as the patient’s age advances. Similarly, the severity of disease also increases with advancement in the patient’s age [26]. Progressive diastolic dysfunction is evident in patients diagnosed with Fabry disease as well as preservation of both EF and fractional shortening [26].

Danon disease

Danon disease mostly affect men in their teens and women in their twenties [2]. Affected men present with HF, mental retardation, and skeletal myopathy [31]. The affected women on the other hand present with affected cardiac myocytes [50]. Whereas cardiac symptoms become evident during adolescence, patients diagnosed with the disease are likely to die of HF in their thirties [31]. The echocardiographic characteristics associated with the Danon disease include LV wall thickness (20 mm to 60 mm) [30] or a significant increase in right ventricular wall thickness (≥ 10 mm) in the absence of pulmonary disease [31]. A severely impaired LV systolic dysfunction has also been reported among patients diagnosed with this condition [30, 31]. Danon disease may also present as sub-endocardial LGE [32] contrary to HCM in which LGE appears mid-epicardial and patchy [30]. Short PR interval and delta waves (Wolf-Parkinson-White syndrome) is a common feature in Danon disease, and consequently resulting into syncope in most patients diagnosed with the disease [31].

Friedreich ataxia

This condition results from expanded Guanine-Adenine-Adenine repeats (120-1,700 times, instead of the usual 8-22) in the frataxin gene on chromosome 9 [51]. The symptoms of the condition begin to show at the age of between 2 and 51 years [51]. However, the cardiac manifestations are likely to occur at the age of between 4 and 5 years, following the onset of the neurologic disorders [2]. Eventually, the development of cardiac disease is expected in 90% to 100% of the patients [39]. As shown in Figure 2, studies have revealed that cardiac involvement is characterized by an increase in wall thickening of the interventricular septum or posterior wall [38,39,52]. Severe HF might develop in some patients with mild LV dilation and systolic dysfunction [38]. Morvan [39] found that if present, valvular insufficiency is often mild in patients diagnosed with this condition. Non-cardiac dyspnea and frequent respiratory infections associated with Friedrich ataxia
### Table 1: Infiltrative Cardiomyopathies with Thick Ventricular Walls and Increased LV Mass

<table>
<thead>
<tr>
<th>Condition</th>
<th>Age at Presentation (Years)</th>
<th>History &amp; Clinical Presentation</th>
<th>Echocardiography</th>
<th>ECG Profile</th>
<th>CMR LGE</th>
<th>Biopsy</th>
<th>Ref. #</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cardiac amyloid</td>
<td>&gt; 30</td>
<td>HF symptoms, nephrotic syndrome, hepatomegaly, idiopathic peripheral neuropathy.</td>
<td>Symmetrical increase in wall thickness of the LV &amp; RV, dilated RA &amp; LA, decreased EF in advanced cases.</td>
<td>Increased to normal QRS complex voltage, short PR interval.</td>
<td>Global, diffuse, pronounced in subendocardium (RV &amp; LV walls).</td>
<td>Myocyte atrophy, normal cardiac tissue is replaced by amyloid.</td>
<td>[27-28]</td>
</tr>
<tr>
<td>Danon Disease</td>
<td>&lt; 20</td>
<td>HF, skeletal myopathy, mental retardation.</td>
<td>A thicker LV (20-60 mm), thick RV in certain cases.</td>
<td>Increased to normal QRS complex voltage, short PR interval.</td>
<td>Sub-endocardial, lack of correspondence to perfusion territory.</td>
<td>Sarcolemmal vacuolization, focal storage of PAS-positive material, myofibrillar disarray.</td>
<td>[30-32]</td>
</tr>
<tr>
<td>Cardiac oxalosis</td>
<td>&gt; 20</td>
<td>Nephrocalcinosis and juvenile urolithiasis.</td>
<td>Symmetrical increase in wall thickness of LV &amp; RV, normal EF, and echo dense speckled reflection.</td>
<td>Increased to normal QRS complex voltage, total heart block.</td>
<td>Increased myocardium attenuation on CT.</td>
<td>Intra and extracellular deposition of oxalate crystals without concomitant inflammation and necrosis.</td>
<td>[33-35]</td>
</tr>
<tr>
<td>Fabry disease</td>
<td>Male: 11+/7, Female: 23+/16</td>
<td>Neuropathic pain, skin rashes, impaired sweating.</td>
<td>Symmetrical increase in wall thickness of LV &amp; RV, and normal EF.</td>
<td>Increased to normal QRS complex voltage, short/ prolonged PR interval.</td>
<td>Focal, midwall, inferolateral wall.</td>
<td>Enlarged myocytes with clusters of concentric glycolipid within lysosomes.</td>
<td>[26, 27, 36, 37]</td>
</tr>
<tr>
<td>Friedreich ataxia</td>
<td>25 (range 2-51)</td>
<td>Gait abnormality.</td>
<td>Increased wall thickness in LV and posterior, normal EF, patchy, echo dense speckled.</td>
<td>Normal QRS complex voltage, ventricular tachycardia.</td>
<td>-</td>
<td>-</td>
<td>[38-41]</td>
</tr>
<tr>
<td>Mucopolysaccharidoses</td>
<td>1-24 (10 is the median year)</td>
<td>Variable depending on condition subtype, delayed mental development, corneal clouding, coarse facial features, hepatosplenomegaly, skeletal deformities.</td>
<td>Asymmetrical septal hypertrophy, normal EF, mitral/aortic valve stenosis/insufficiency.</td>
<td>Increased/decreased QRS complex voltage, malignant arrhythmia.</td>
<td>-</td>
<td>Swollen myocytes with clear cytoplasm due to accumulation of mucopolysaccharides within lysosomes.</td>
<td>[42-45]</td>
</tr>
</tbody>
</table>

![Figure 2](image-url): Increased LV Mass and Thickening of Ventricular Walls in Cardiac Amyloidosis

(A) Hypertrophic non-obstructive cardiomyopathy; (B) Hypertensive heart disease with secondary renal failure; (C) Cardiac amyloid; (D) Mucopolysaccharidosis; (E) Cardiac oxalosis; (F) Friedreich ataxia. Adopted From: Seward and Casaclang-Verzosa, (2010), p.1774, [2]
are because of severe scoliosis and neuromuscular impairment of respiratory muscles [38]. Leone [53] analyzed survival among Italian patients diagnosed with Friedreich's disease found the 10-year, 20-year, and 30-year survival rates among were 96%, 80%, and 61% respectively. Inter-current pulmonary infection and cardiac dysfunction are the causes of death among patients diagnosed with Friedreich's disease [38,42].

**Myocardial oxalosis**

This condition is characterized by an enhanced generation of oxalic acid, caused by the deposition oxalate crystals in various organs, including the kidney and the heart [33, 54]. Palka Schulze [33] and Palka [34] identified biventricular symmetrically thickened wall (Figure 2), considered the echocardiographic characteristic of myocardial oxalosis. The early stages of this disease may be characterized by a normal EF [34]. However, mild biventricular dilation in advanced cases of the disease [34]. Computed tomography may demonstrate oxalate deposits as increased attenuation of the myocardium during the very late stages of the disease (Figure 3). Diastolic function in patients diagnosed with the condition maybe severely impaired, with restrictive filling and elevated filling pattern [33,34].

**Mucopolysaccharidosis**

This condition is a representation of inborn errors of metabolism due to deficiencies in lysosomal enzymes that are responsible for the breaking down of glycosaminoglycan [55]. The proper functioning of the cell is impaired by the accumulation of partially degraded mucopolysaccharides resulting into a range of clinical manifestations. These disorders are hereditary in nature [2]. The inheritance of the disorders occur in an autosomal recessive manner, and have an equal effect on both men and women [56]. Patients diagnosed with the condition may exhibit normal intellect or mental retardation, as well as experience developmental delay or severe behavioral problems [57]. Whereas the occurrence of cardiac disease is nearly 100% in all subtypes of Mucopolysaccharidosis, Type I Mucopolysaccharidosis (Hurler-Scheie syndrome) presented the most severe cardiac involvement [58]. Common findings associated with Mucopolysaccharidosis include significant valvular thickening and secondary pulmonary hypertension [42, 58]. Rigante [58] reports that asymmetrical septal hypertrophy precedes thickening of the valves. According to Rigante [58], the mitral and the aortic valves are more often involved, and hence producing insufficiency and/or stenosis. LV systolic function is normal in most cases [58]. However, ECG evidence of LV hypertrophy is uncommon [55, 58]. The characteristic small QRS complex voltages to poor conductance of glycosaminoglycan and rare intraventricular conduction delay or malignant arrhythmias are common among patients diagnosed with Mucopolysaccharidosis [58].

**Etiopathogenesis**

The etiopathogenesis for ICM is incompletely understood. However, a number of studies have reported etiologic or pathologic processes for the different etiopathogenic mechanisms of ICM. These mechanisms includes cardiac amyloidosis and cardiac sarcoidosis as discussed in the subsequent sections.

**Cardiac amyloidosis**

The extracellular deposition of amyloid fibrils within the heart result in abnormalities in three vital processes for the normal cardiac function namely; conduction, contractility, and coronary blood flow [1]. The thickening of the myocardial wall, and its consequent firm, rubbery, and noncompliant state is due to amyloid accumulation [59]. The thickening of the myocardial wall leads to a rise in intra-cardiac pressures, impeding progressive biventricular diastolic dysfunction [60,61]. Cumulative myocyte damage and local fibrosis could lead to systolic dysfunction as the disease progresses [1]. However, systolic dysfunction is typically evident in extremely advanced disease [2]. Ridolfi [62] analyzed the conduction system in cardiac amyloid and established the involvement of the sinoatrial node, bundle branches, atrioventricular (AV) note could manifest at various degrees of heart block, and complex ventricular arrhythmias can be evident.

**Cardiac sarcoidosis**

Sarcoidosis needs at least three key events to develop: exposure to antigen; acquired cellular immunity directed against the antigen mediated through antigen presenting cell and antigen specific T lymphocytes; as well as the appearance of immune effector cells necessary for promoting a more no-specific inflammatory response [22]. Granulomatous infiltration leading to the disruption of normal organ function is the major pathologic feature of sarcoidosis [1]. Figure 4 illustrates the pathogenesis of cardiac sarcoidosis.

In Figure 4, cytokine profile shifts to that of T helper type 2 response at a later stage of lesion evolution [22]. This shift exerts anti-inflammatory effects, and leads to tissue scarring. Cardiac sarcoidosis is characterized by three main successive stages namely; edema; granuloma formation; and fibrosis, which results in scar formation [22]. Sarcoid can involve any part of the heart, including the myocardium, endocardium, and pericardium [1]. However, Roberts [63] established that the disease's involvement is typical of the LV free wall and the basal aspect of the interventricular septum. In Figure 5, iannuzzi [9] illustrates the presence of CD4+ T cells, interacting with antigen-presenting cells to initiate the formation and maintenance of granulomas, is a cardinal feature of the sarcoidosis.

The CD4+ T cells interacting with antigen-presenting cells causes the myocardium to thicken and predominate leading to diastolic dysfunction in the initial stages of the disease [1]. However, in the later stages of the disease dilation of the ventricles, global or segmental hypokinesia, and predominance of systolic dysfunction occurs. The
patchy nature of myocardial involvement can lead to uneven wall motion abnormalities that fail to match any given coronary distribution [64]. Despite the various genetic or acquired etiologies, the hallmark etiopathogenesis process of the ICM is the infiltration of the heart by abnormal substances [1]. Given the systemic nature of the underlying disease, ICM occurs in various age groups presenting as extra-cardiac clinical manifestations. Whereas conduction abnormalities and diastolic HF with restrictive physiology predominate the initial stages of the disease, the advanced stages of the disease on the other hand are characterized by systolic dysfunction.

**Diagnosis**

The diagnosis for ICM requires a high degree of clinical suspicion [1]. The 2016 Scientific Statement from American Heart Association on current diagnostic and treatment strategies for specific dilated cardiomyopathies recommends the basis of ICM diagnosis is the detection of its two primary underlying infiltrative disorders – amyloidosis or sarcoidosis [64]. The most commonly adopted diagnostic methods for obtaining confirmatory evidence for ICM are cardiac imaging methods – endomyocardial biopsy, echocardiography, and CMR [1]. Other methods providing important clues for differential diagnosis include electrocardiography [66], tissue diagnosis [67-69], nuclear imaging [70, 71]

**Cardiac amyloidosis/sarcoidosis**

The AHA recommends the diagnosis of ICM require multiple diagnostic methods to detect cardiac amyloidosis and cardiac sarcoidosis as summarized in Table 2 [64].

**Imaging methods**

**Endomyocardial biopsy**

Endomyocardial biopsy is the gold standard for the diagnosis cardiac amyloidosis [22]. Endomyocardial biopsy has a high diagnostic accuracy for cardiac sarcoidosis. However, an evaluation of endomyocardial biopsies by found that the sensitivity of endomyocardial biopsy for non-caseating granulomas was low (< 20%) [65]. The low sensitivity associated with endomyocardial biopsy is due to patchy myocardial

Table 2: The AHA Diagnostic Strategies for ICM

<table>
<thead>
<tr>
<th>No.</th>
<th>AHA Recommendations for Cardiac Amyloidosis</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>In patients suspected with cardiac and systematic amyloidosis, detection of amyloid proteins in tissues such as abdominal fat pad, gingiva or rectum or affected organs such as heart, liver and kidney are recommended for amyloidosis diagnosis.</td>
</tr>
<tr>
<td>2</td>
<td>Upon detection of transthyretin (TTR) from biopsy specimen, DNA mutational analysis should distinguish between senile and hereditary amyloidosis.</td>
</tr>
<tr>
<td>3</td>
<td>After the confirmation of cardiac amyloidosis, cardiac imaging and laboratory examination should be considered to identify cardiac involvement.</td>
</tr>
<tr>
<td>4</td>
<td>Echocardiography or cardiac MRI is recommended to patients with suspected cardiac amyloidosis to determine cardiac restrictive filling and heart failure.</td>
</tr>
<tr>
<td>5</td>
<td>Endomyocardial diagnosis is recommended to diagnose cardiac amyloidosis in suspected patients when there is no evidence of non-cardiac tissue amyloidosis</td>
</tr>
<tr>
<td>6</td>
<td>Natriuretic peptide (BNP of NT-pro BNP) is recommended to detect pre-clinical cardiac involvement in patients diagnosed with amyloidosis.</td>
</tr>
<tr>
<td>7</td>
<td>ECG abnormalities such as low QRS voltage in the presence of ventricular thickening may suggest cardiac amyloidosis.</td>
</tr>
<tr>
<td>8</td>
<td>Nuclear imaging may assist in differentiating AL from TTR amyloidosis.</td>
</tr>
</tbody>
</table>

**Endomyocardial biopsies** is useful to confirm cardiac sarcoidosis when pathology reveals non-caseating granulomas.
involvement, which frequently involves the intraventricular septum and the LV [22]. Despite the low sensitivity, endomyocardial biopsy remains the gold standard for diagnosing cardiac sarcoidosis [1].

Echocardiography

Echocardiography has been adopted for most forms of ICM, including cardiac amyloid [61, 72-76], cardiac sarcoidosis [77], Fabry disease [78], Danon disease [79], as well as Friedreich's disease [39, 80]. Echocardiography is useful for the assessment of thickening of the LV wall, while Doppler echocardiography for the assessment of abnormal alterations in cardiac hemodynamics such as basal strain rate, peak early velocity, peak late velocity, and isovolumic relaxation time [72-76].

Cardiac magnetic resonance

The use of CMR as a diagnostic method has for ICM has been investigated in various studies involving various forms of ICM such as cardiac amyloid [81], cardiac sarcoidosis [82, 83], Fabry disease [78], and Danon disease [84]. However, whereas the accuracy of cardiac MRI has not been established in assessment of myocardium in ICM patients and is not routinely employed for diagnosing the Friedreich's disease Weidemann [85], the modality is helpful in determining the extent of cardiac involvement.

Differential diagnosis

Differenitizing various forms of ICM such as cardiac amyloidosis and Fabry disease from hypertrophic cardiomyopathy (HCM) is challenging since these condition share a number of clinical and hemodynamic features such as LV hypertrophy and diastolic dysfunction [75,78,118,119]. Accurate diagnosis is therefore critical when handing patients suspected of ICM [120]. A number of studies have attempted to differentiate various forms of ICM from HCM. Kubo [117], through a case control study comprising 46 consecutive patients with ICM (11 patients) or HCM (35 patients), sought to investigate the potential of high-sensitivity cardiac troponin T (hs-cTnT) in differentiating ICM from HCM. The study found that serum hs-cTnT was significantly higher in ICM patients than in their HCM counterparts (0.083 ± 0.057 ng/ml versus 0.027 ± 0.034 ng/ml, p < 0.001). The findings led Kubo [117] to conclude that serum hs-cTnT is an effective and accurate diagnostic indicator for differentiating ICM from HCM.

Baccouche [75] differentiated cardiac amyloidosis (CA) from HCM using 3-dimensional (3D) speckle tracking-derived functional parameters. Using a cohort of 12 patients with CA and another cohort of 12 patients with HCM, diagnosed using CMR, Baccouche [75] performed a 3D speckle tracking echocardiographic with wall motion analysis at radial strain (RS) longitudinal, and circumferential. The results indicated that 3D speckle-tracking function parameters were relatively lower in HCM patients and minimal in CA patients. In addition, basal RS was significantly reduced in CA patients (7.5 ± 19.7 vs. 22.3 ± 22.7; P < 0.0001) [75]. The study concluded that the 3D speckle tracking echocardiography was a useful predictor for the accurate differentiation of CA from HCM. De Cobelli [78] performed a comparison of the myocardial location and distribution patterns of delayed enhancement between patients with Fabry disease and those with HCM. The study found that patients with Fabry disease exhibited LV delayed enhancement with a typical and consistently established pattern, characterized by the involvement of the inferolateral basal or mid basal segments and a mesocardial distribution, spreading to the subendocardium, and thus suggestive of Fabry disease.

Meta-analysis of diagnostic methods

Search strategy

The present meta-analysis sought to combine current research findings on studies investigating the diagnosis of ICM to advance the knowledge and understanding of diagnostic methods and features. Studies were included if they (a) recruited patients either prospectively and retrospectively; (b) investigated at least one type of ICM diagnostic method; (c) provided clinical outcomes of the diagnostic methods. The relevant studies were searched from online databases, both indexed and non-indexed, as well as from screening of bibliographies of included studies and review studies. Broad-based search terms were used including diagnosis” OR “diagnostic methods” AND “cardiac amyloidosis” OR “Friedreich's disease” OR “Fabry disease” OR “cardiac amyloidosis”. Titles and/or abstracts of the retrieved studies believed to satisfy the inclusion criteria were included for further review pending inclusion. Additional studies were obtained from screening bibliographies of the included studies and review articles. There was no restriction on publication date or language. Case reports (investigate few patients) or studies with data that was not readily extracted were excluded. Each included study was reviewed against the inclusion criteria, data extracted and recorded on Microsoft Excel Worksheet. The extracted data was name of the first author, period of study; type of ICM investigated; sample size of patients recruited; diagnostic method used; diagnostic feature targeted and a summary of the main findings. Table 3 summarizes the results of the meta-analysis of these different diagnostic methods for various forms of ICM.

Study characteristics

Nineteen (19) studies, 10 prospective and 9 retrospective meeting the inclusion criteria were included in the present meta-analysis. The 19 studies covered a period of thirty years, published between 1982 and 2012. The long period allows the assessment of the development and accuracy of diagnostic methods for ICM over time. The 19 studies recruited a combined population of 104 patients diagnosed with ICM or its main underlying cardiac etiologies. Cardiac amyloidosis was the most frequently investigated and diagnosed etiologic cause of ICM by 53% of the included studies [61,66,67,70,71,73,74,75,76,81] followed by cardiac sarcoidosis by 21% of the included studies [77,82,113,114]. The other less common cardiac conditions investigated were Fabry disease (FD) [78,115] and Friedreich ataxia (FA) [39,80] at 11% each, and finally Danon disease at 5% [79]. The most studied diagnostic method was echocardiography imaging studied by 47% followed cardiac MRI by 16%, electrocardiography by 11%, radionuclide imaging by 11%, and tissue diagnosis by 5% of the included studies.

Study outcomes

Diagnosis of ICM depends on detection of the primary etiology. The most common cardiac condition leading to ICM is cardiac amyloidosis followed by cardiac sarcoidosis and Fabry disease. The mainstay of the diagnosis of cardiac amyloidosis was cardiac imaging by echocardiography, which could may be complemented by cardiac MRI and radionuclide imaging techniques. Echocardiographic imaging was used for characterizing LV dysfunction defined as increased LV wall thickness ≤ 12 mm and increased atrial septal thickness [61,73,76], and ventricular myocardium in infants defined as myocardial hyper-echogenicity [74]. Doppler Tissue echocardiography was also useful to detect changes in hemodynamics defined as significantly reduced basal strain rate, peak early velocity and increased late velocity and prolonged isovolumic relaxation time [61,75]. Cardiac MRI reveals abnormal LGE
Table 3: Summary of Data from Studies Investigating Diagnosis of ICM

<table>
<thead>
<tr>
<th>1st Author [Ref. #]</th>
<th>Period of Study</th>
<th>Form of ICM</th>
<th>Sample Size</th>
<th>Diagnostic Method(s)</th>
<th>Diagnostic Feature</th>
<th>Summary of Main Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>Murtagh [66]</td>
<td>2000-2005</td>
<td>Cardiac amyloidosis</td>
<td>127</td>
<td>Electrocardiography systemic amyloidosis and biopsy-proved cardiac involvement</td>
<td>-Low-voltage QRS witnessed in 50% of cases of AL amyloidosis, suggestive of cardiac amyloidosis.</td>
<td>-A pseudoinfarct pattern (poor R-wave progression Q5 waves) in the precordial leads observed in 50% of cases, suggestive of cardiac amyloidosis.</td>
</tr>
<tr>
<td>Lee [73]</td>
<td>1995-2012</td>
<td>Cardiac amyloidosis</td>
<td>117</td>
<td>Echocardiography Left ventricular wall thickness (LVWT)</td>
<td>-A third of patients diagnosed as having an LVWT of ≤ 12 mm.</td>
<td>-There is need for early detection using sensitive diagnostic methods even when LVWT is not increased.</td>
</tr>
<tr>
<td>Falk [74]</td>
<td>1982-1987</td>
<td>Cardiac amyloidosis</td>
<td>31</td>
<td>Echocardiography Ventricular myocardium</td>
<td>-Increased myocaridal echogenicity observed in 16 patients (sensitivity of 63%; specificity of 74% for diagnosis of amyloidosis).</td>
<td>-Increased atrial septal thickness observed in 60% of amyloid patients.</td>
</tr>
<tr>
<td>Klein [61]</td>
<td>1987-1989</td>
<td>Cardiac amyloidosis</td>
<td>64</td>
<td>Doppler echocardiography Left ventricular diastolic function.</td>
<td>-55 patients (with cardiac involvement) exhibited prominent abnormalities of left ventricular diastolic filling (LVWT: early &gt; 12 mm but &lt; 15 mm; advanced ≥ 15 mm).</td>
<td>-Relaxation appeared normal in early amyloidosis, with decreased peak early velocity (75 ± 20 versus 86 ± 16 cm/s; p &lt; 0.01), increased late velocity (71 ± 22 versus 56 ± 13 cm/s; p &lt; 0.01), decreased early to late velocity ratio (1.2 ± 0.6 versus 1.6 ± 0.5; p &lt; 0.01) and prolonged isovolumic relaxation time (87 ± 15 versus 73 ± 13 ms; p &lt; 0.01) compared with normal values.</td>
</tr>
<tr>
<td>Baccouche [75]</td>
<td>2007-2012</td>
<td>Cardiac amyloidosis</td>
<td>12</td>
<td>Echocardiography Basal, midventricular, and apical.</td>
<td>-Patients with amyloidosis exhibited a significantly reduced basal radial strain (RS) (7.5 ± 19.7 vs. 22.3 ± 22.7; P &lt; 0.0001).</td>
<td>- 99mTc-DPDscintigraphy a useful step in the work-up of the differential diagnosis of TTR versus AL etiology in patients with documented CA.</td>
</tr>
<tr>
<td>Koyama [76]</td>
<td>2001-2002</td>
<td>Cardiac amyloidosis</td>
<td>119</td>
<td>Echocardiography Tissue velocity, strain, and strain rate at the basal, mid, and apical LV multiple walls.</td>
<td>-70 patients exhibited cardiac involvement defined as the mean value of LVWT &gt; 12 mm.</td>
<td>- 99mTc-PYP cardiac imaging distinguishes AL from ATTR CA and may be a simple, widely available method for identifying subjects with TTR amyloidosis.</td>
</tr>
<tr>
<td>Syed [81]</td>
<td>2006-2007</td>
<td>Cardiac amyloidosis</td>
<td>120</td>
<td>CMR Late gadolinium enhancement (LGE)</td>
<td>-Abnormal LGE found in 34 (97%) of the 35 patients with histologically verified cardiac amyloidosis (CA), and increased echocardiographic LVWT in 32 (91%) of the 35 patients.</td>
<td>-LGE is common in CA and detects interstitial expansion from amyloid deposition.</td>
</tr>
<tr>
<td>Perugini [70]</td>
<td>2004-2005</td>
<td>Cardiac amyloidosis</td>
<td>25</td>
<td>Radionuclide Imaging Left ventricular mass and renal function.</td>
<td>-Patients with TTR CA had a significantly higher semi-quantitative cardiac visual score than the AL cohort (2.9±0.06 vs. 0.8±0.27, p&lt;0.0001) as well as a higher quantitative score (1.80±0.04 vs.1.21±0.04, p&lt;0.0001).</td>
<td>-Combination of increased myocardial echogenicity and increased atrial thickness was 60% sensitive and 100% specific for the diagnosis of amyloidosis.</td>
</tr>
<tr>
<td>Bokhari [71]</td>
<td>2005-2008</td>
<td>Cardiac amyloidosis</td>
<td>45</td>
<td>Radionuclide Imaging Cardiac retention.</td>
<td>- Patients with TTR CA had a significantly higher semi-quantitative cardiac visual score than the AL cohort (2.9±0.06 vs. 0.8±0.27, p&lt;0.0001) as well as a higher quantitative score (1.80±0.04 vs.1.21±0.04, p&lt;0.0001).</td>
<td>- Heart and heart/whole-body tracer retention were significantly higher (p &lt; 0.05) in patients with cardiac involvement of TTR-related.</td>
</tr>
<tr>
<td>Ardehali [67]</td>
<td>1982-1997</td>
<td>Cardiac amyloidosis</td>
<td>845</td>
<td>Tissue Diagnosis Endomyocardial biopsy (EMBx)</td>
<td>-The final diagnosis differed from the initial clinical diagnosis in 264 (51%) of the 518 patients, with EMBs constitution 75% (evident in 196 patients) of the 264 patients.</td>
<td>-Abnormal LGE found in 34 (97%) of the 35 patients with histologically verified cardiac amyloidosis (CA), and increased echocardiographic LVWT in 32 (91%) of the 35 patients.</td>
</tr>
<tr>
<td>Burstow [77]</td>
<td>1983-1987</td>
<td>Cardiac sarcoidosis</td>
<td>88</td>
<td>Echocardiography LV systolic dysfunction</td>
<td>-Echocardiographic evidence of LV systolic dysfunction recorded in 14% (in 12 patients) of the 88 patients, suggestive of cardiac sarcoidosis.</td>
<td>-The 12 patients exhibited a high incidence of congestive HF (9 of 12) and conduction system disease (7 of 12).</td>
</tr>
</tbody>
</table>

Infiltrative cardiomyopathies refer to a heterogeneous group of cardiac conditions characterized by restrictive cardiac physiology. As a result, the basis of diagnosis is the detection of the causative cardiac condition, mainly cardiac amyloidosis, cardiac sarcoidosis, Fabry disease and Friedreich’s ataxia. Diagnosis of cardiac amyloidosis requires a combination of clinical examination and cardiac imaging. Echocardiography-defined increased LV wall thickness ≥ 2 mm and atrial dilatation may suggest cardiac amyloidosis. Cardiac LGE-MRI may be useful in the detection of the extent of cardiac involvement while radionuclide imaging, although insensitive, is useful in the differentiation of AL amyloidosis (immunoglobulin light chain amyloidosis) and transthyretin (TTR) amyloidosis. The present findings are consistent with AHA recommendation on the diagnosis of cardiac amyloidosis in patients suspected with infiltrative cardiomyopathies [64]. The AHA recommends echocardiography imaging-modality in the diagnosis of cardiac amyloidosis because of the uncertainty of the accuracy of cardiac LGE-MRI and low sensitivity of nuclear imaging.

### Discussion

Infiltrative cardiomyopathies refer to a heterogeneous group of cardiac conditions characterized by restrictive cardiac physiology. As a result, the basis of diagnosis is the detection of the causative cardiac condition, mainly cardiac amyloidosis, cardiac sarcoidosis, Fabry disease and Friedreich’s ataxia. Diagnosis of cardiac amyloidosis requires a combination of clinical examination and cardiac imaging. Echocardiography-defined increased LV wall thickness ≥ 2 mm and atrial dilatation may suggest cardiac amyloidosis. Cardiac LGE-MRI may be useful in the detection of the extent of cardiac involvement while radionuclide imaging, although insensitive, is useful in the
Clinical management of cardiac amyloidosis targets treatment of HF caused by restrictive cardiomyopathy and therapy targeting the underlying protein disorder [1]. The use of loop diuretics helps in attaining euvoemla [1]. However, beta-blockers and angiotensin converting enzyme (ACE) inhibitors are typically avoided in the clinical management of cardiac amyloidosis [1]. Beta-blockers are avoided because cardiac output is largely dependent on heart rate in the setting of a fixed stroke volume while ACE-inhibitors increases the likelihood of neurohormonal blockade resulting in significant hypotension in the setting of the underlying automatic neuropathy [86, 87]. Device therapy – implantable cardioverter defibrillators (ICD) may be considered for preventing sudden cardiac death but its protective value has not been established in patients with cardiac amyloidosis [1].

For the treatment of AL amyloidosis, a melphalan-based cytotoxic chemotherapy regimen with or without autologous stem cell transplantation is often recommended [88]. However, a recent study by Mikhail [89] reported that cyclophosphamide-bortezomib-dexamethasone (CyBorD) generates rapid and complete hematologic response in patients diagnosed with AL amyloidosis. Even though rare, cardiac transplantation has been found to be a viable clinical management option for patients with AL amyloidosis, with isolated cardiac disease that makes them inappropriate candidates for neoadjuvant chemotherapy [90]. Liver transplantation has been found to be a potential curative clinical management method for ATTR amyloidosis, particularly among patients without existing cardiac involvement [100]. Finally, a number of pharmacotherapies such as the nonsteroidal anti-inflammatory drugs (NSAIDs) diflunisal and its non-NSAID analog tafamidis [101, 102], as well as RNAs [103], antisense oligonucleotides [104], and green tea extracts [105] have been used for purposes of reducing, stabilizing, or silencing TTR activity or production.

Cardiac sarcoidosis

Glucocorticoids are the mainstay in clinical management of cardiac sarcoidosis initiated promptly following diagnosis [1]. Due to paucity of randomized controlled trials confirming the efficacy of glucocorticoids, clinical management strategies involving glucocorticoids rely on observational studies [106, 107]. The ultimate duration of treatment is determined by clinical response and can be guided by monitored improvement in LGE on CMR [108]. For patients indicated for pacemaker and ICD, careful consideration is necessary due to the high incidence of SCD [1]. The existence of CHB or high-grade AV block is a demonstration of permanent pacemaker implantation, even in if the AV block reverses transiently [109]. Cardiac implantation is a potential option for managing cardiac sarcoidosis in young patients with endstage. New York Heart Association (NYHA) Class IV HF, despite optimal medical therapy, and for patients diagnosed with intractable ventricular arrhythmias [1].

Fabry disease

Enzyme replacement is the mainstay method of clinical management of the Fabry disease, which should be initiated prior to the development of tissue injury and fibrosis [110].

Danon disease

The performance of cardiac transplantation is a potentially lifesaving clinical management procedure for patients diagnosed with the Danon disease [111]. However, patients with the disease often die early in life [1].

Friedreich ataxia

The clinical management approach to the Friedreich's disease is largely supportive, and limited to conventional HF medication, device implantation, and antiarrhythmic drugs [112].

Meta-analysis of clinical management methods

The present meta-analysis combines findings from previous research articles on clinical management of ICM. The intention is to advance the knowledge and understanding on the most common clinical management approaches in use today and their clinical effectiveness.

Study selection

Studies investigating clinical management methods of ICM were searched on online electronic search engines - PubMed, EMBASE, Scopus and Google Scholar. Relevant medical subject headings (MESH) terms and a combination of key words to ensure no relevant study was left out. The key search words used were "infiltrative cardiomyopathy" OR "Cardiac Amyloidosis" OR "Cardiac Sarcoidosis" AND "Clinical Management" OR “Treatment” OR “Implantation”, “Chemotherapy” OR “Medical Therapy”. The search was not limited to publication year, language or the age of the recruited patients. Study selection began with screening of titles and abstracts useful for excluding case studies (involving one or a few ICM patients) animal studies and conference papers (still subject to revision) and compared abstracts to the inclusion criteria. The inclusion criteria were the study (a) recruited patients diagnosed with ICM or one of the underlying cardiac conditions; (b) included at least one clinical management methods; and (c) reported outcomes on clinical management. After selection, relevant data was extracted from all the included studies. The extracted data was recorded in Microsoft Excel Worksheet. The extracted data included first author, study period, sample, mean age, type of ICM, treatment, follow-up period, and the main outcomes. Table 4 provides a summary of data extracted from the included studies.

### Table 4. Summary of Results of Studies on Clinical Management of ICM

<table>
<thead>
<tr>
<th>1st Author [Ref #]</th>
<th>Study Period</th>
<th>Sample Size</th>
<th>Mean Age (Yrs.)</th>
<th>ICM Form</th>
<th>Treatment</th>
<th>Follow-Up Period (Months)</th>
<th>Outcome(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Varr [116]</td>
<td>2009-2012</td>
<td>31</td>
<td>68</td>
<td>Cardiac amyloidosis</td>
<td>Implantable cardioverter-defibrillator (ICD)</td>
<td>23</td>
<td>-ICD successful in most patients, with therapy resulting in the termination of the arrhythmia in 80% of the patients, and hence demonstrating its potential for primary prevention of sudden cardiac death (SCD).</td>
</tr>
<tr>
<td>Mikhail [89]</td>
<td>2008-2010</td>
<td>17</td>
<td>59</td>
<td>Confirmed and symptomatic AL</td>
<td>Cyclophosphamide, bortezomib, and dexamethasone (CyBorD)</td>
<td>21</td>
<td>-16 of 17 (94%) of patients responded to CyBorD, with 71% achieving complete hematologic response and 24% a partial response. - CyBorD produced rapid and complete hematologic responses in most of the patients.</td>
</tr>
<tr>
<td>Maurer, [90]</td>
<td>1997-2004</td>
<td>25</td>
<td>63</td>
<td>Systemic amyloidosis and HF</td>
<td>Cardiac transplantation followed by stem cell or liver transplantation.</td>
<td>6</td>
<td>-Patients who received cardiac transplantation followed by stem-cell or liver transplantation showed improved survival, and hence a novel therapeutic option for amyloid patients.</td>
</tr>
<tr>
<td>Dubrey [100]</td>
<td>1990-1995</td>
<td>11</td>
<td>63</td>
<td>Familial amyloidosis (FAP)</td>
<td>Liver transplantation</td>
<td>102</td>
<td>-5 patients showed progression of LVWT, with increased LV mass after liver transplantation. Three of the five patients exhibited reduction in electrocardiographic voltage despite abolition of the mutant protein from the serum. Despite 9 of the 11 patients showing symptomatic improvement in neurologic symptoms, 1 patient developed HF and a second patient suffered SCD.</td>
</tr>
<tr>
<td>Berk [101]</td>
<td>2006-2012</td>
<td>130</td>
<td>59.7</td>
<td>Familial amyloid polyneuropathy</td>
<td>Pharmacotherapy (diflunisal)</td>
<td>24</td>
<td>-Reduced rate of progression of neurologic endpoints observed.</td>
</tr>
<tr>
<td>Coelho [102]</td>
<td>2010-2012</td>
<td>125</td>
<td>61</td>
<td>Transthyretin familial amyloid polyneuropathy (TTR-FAP)</td>
<td>Pharmacotherapy (Tafamidis treatment)</td>
<td>18</td>
<td>-A higher-than-anticipated liver transplantation dropout rate was evident. -Reduced rate of progression of neurologic endpoints observed.</td>
</tr>
<tr>
<td>Kristen [105]</td>
<td>2011-2012</td>
<td>9</td>
<td>65</td>
<td>amyloidosis transthyretin (ATTR) 7 of 9 cardiomypathy</td>
<td>Green tea (GT) and/or green tea extracts (GTE)</td>
<td>12</td>
<td>-No increase of LVWT and LV myocardial mass was observed by echocardiography after 12 months of follow-up. Observations suggestive of inhibitory effect of GT and/or GTE on the progression of cardiac amyloidosis.</td>
</tr>
</tbody>
</table>

| Total/Average     | 368          | 57           | 31              |          |           |                           |            |

### Study characteristics

After all the potentially relevant studies were screened against the inclusion criteria, nine (9) studies were included in this meta-analysis. The studies included six (6) retrospective and three (3) prospective studies. The nine studies recruited a combined population of 368 patients diagnosed with various ICM-related disorders with a mean age of 57 years followed up for a mean period of 31 months, range 6 to 102 months. A greater majority of the studies (78%) investigated cardiac amyloidosis, while the remaining studies (11% each) investigated treatment option for cardiac sarcoidosis and Danon Disease. The most common clinical management method was medical therapy assessed by a third of the studies (45%), followed by cardiac/liver transplantation (33%), ICD (11%) and clinical monitoring (11%).

### Study outcomes

Medical therapy was the most common treatment for cardiac amyloidosis. A combination of cyclophosphamide, bortezomib, and dexamethasone (CyBorD) had 94% positive response with a rapid and complete hematologic response in 71% of the patients and a partial response in 24% of the patients [89]. Nonsteroidal anti-inflammatory drug (NSAID) – diflunisal – reduced the rate of progression [101]. Tafamidis, a specific transthyretin (TTR) stabilizer or dissociation inhibitor, was affective in the treatment of TTR amyloidosis. The drug preserved LV wall thickness and LV myocardial mass after 12 months [102]. Green tea extracts (GTE) was also effective in inhibiting the progression of cardiac TTR amyloidosis [105]. Besides medication, cardiac or liver transplantation was a common clinical management approach in ICM patients. In patients with systemic amyloidosis and heart failure, heart transplantation followed by liver transplantation significantly improve survival [90]. Liver transplantation alone is less effective in cardiac amyloidosis since about half of patients with cardiac amyloidosis exhibit progression of LV wall thickness. Cardiac transplantation also showed great improvement in clinical outcomes of patients with Danon Disease [111]. For patients with cardiac sarcoidosis, routine monitoring reveal improvement in LV systolic function imaged using cardiac LGE-MRI [108].

### Discussion

The principal target of clinical management approaches for patients diagnosed with ICM is treating the primary underlying etiology. In the present findings, clinical management of ICM focused on treating cardiac amyloidosis, sarcoidosis and Danon disease, which represent the main primary etiologies of ICM. The target of therapy was relieving restrictive filling (reducing LV wall thickness), positive hematologic response and the treatment of heart failure. The primary therapy was medication to achieve hematologic response and reverse LV wall thickening. Cardiac transplantation followed by or liver transplantation conveyed a protective effect improving survival in patients with systemic amyloidosis and symptomatic heart failure. Consistent with these findings, the current recommendations by AHA propose medication,
cardiac transplantation and liver transplantation as effective therapy for TTR amyloidosis without significant cardiac involvement. However, the recommendations caution that conventional HF medication such as ACE-inhibitors and beta-blockers should be avoided because of the risk or neurohormonal blockade and preservation of heart rate [64]. Although the present analysis finds ICD is important in protecting against sudden cardiac death, the AHA recommendations suggests its clinical value on preventing sudden cardiac death has not been conclusively demonstrated [64].

The treatment of cardiac sarcoidosis-induced ICM has limited clinical trials. Whereas the present findings suggest monitoring of cardiac function by cardiac LGE-MRI to predict subsequent risk of developing systolic dysfunction, the AHA guidelines recommend corticosteroids are the mainstay of cardiac sarcoidosis therapy in symptomatic patients. However, randomized clinical trials on the efficacy of corticosteroids are lacking but retrospective studies suggest a clinically beneficial effect in improving LV function. Furthermore, treatment of cardiac sarcoidosis in asymptomatic or minimally symptomatic patients remains controversial [64]. Standard HF therapy should be considered for patients with cardiac sarcoidosis and significantly reduced LV ejection fraction [106, 107]. Finally, clinical guidance for under underlying etiologies of ICM such as Fabry Disease and Danon Disease are lacking, with management varying in between studies. In summary, the mainstay clinical management of ICM is the identification and treatment of the underlying etiologies, principally, cardiac amyloidosis and sarcoidosis. The primary treatment is medical therapy. Additional therapies are cardiac and/or liver transplantation for cardiac amyloidosis, and corticosteroids and conventional HF treatment for cardiac sarcoidosis. 

Conclusion

Infiltrative cardiomyopathy define a heterogeneous group of myocardial disorders characterized by the deposition of abnormal substances within the intercellular spaces leading to diastolic and/ or systolic dysfunction. The cardiac condition manifests as increased rigidity of ventricular walls disrupting ventricular filling. It can be idiopathic or secondary to cardiac or systemic disorders. It is a rare form of cardiomyopathy with a prevalence of 2 to 7% of reported cardiomyopathies. Risk factors include advanced age, female gender and the presence of diabetes, hypertension, chronic kidney disease, and obesity. Clinically, ICM mainly manifests as cardiac amyloidosis and cardiac sarcoidosis and in fewer cases, Fabry disease, Danon Disease and Friedreich’s ataxia. Prognosis of ICM is challenging since it varies considerably based on the primary underlying etiology. The etiopathogenesis of ICM remains incompletely understood but cardiac amyloidosis and sarcoidosis have been implicated as the key etiopathogenic factors. The heterogeneity of underlying etiologies complicates diagnosis of ICM. Although endomyocardial biopsy is the gold standard for diagnosis of cardiac amyloidosis, echocardiography is the most common imaging method for diagnosis. Cardiac LGE-MRI is valuable in the diagnosis of cardiac sarcoidosis. The two cardiac imaging methods diagnose ICM by characterizing increasing LV wall thickness and restrictive ventricular filling. Finally, clinical management approaches target to relieve hemodynamics (reduce LV wall thickness), achieve complete hematologic response and relieve symptoms of heart failure. The common treatment options include medical therapy, implantable cardioverter defibrillator and cardiac/ liver transplantation.

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