Amyloidosis is a collection of diseases characterized by extracellular deposition of the fibrillar proteinaceous material, amyloid, in tissues and organs. Amyloid can be formed from several precursor proteins that misfold into toxic oligomers, which then aggregate with proteoglycan and serum amyloid proteins to form amyloid infiltrates. The vast majority of cardiac amyloidosis (CA) is caused by one of two proteins: light chain or transthyretin (TTR). Light-chain (AL) amyloidosis is a hematological disorder of proliferation of an abnormal plasma cell clone that overproduces lambda or, less commonly, kappa light chains. The abnormal circulating light chains form amyloid that can affect multiple organs. AL amyloidosis has equal prevalence in both sexes and typically manifests between the ages of 40 and 80 years. Therapy is primarily aimed at managing plasma cell dyscrasia using chemotherapy or stem cell transplantation.

TTR is a pre-albumin produced by the liver and functions as a transporter of thyroxine and retinol in its normal tetrameric form. The monomeric form of TTR tends to misfold to produce amyloid fibrils that are deposited in tissues. Amyloidosis resulting from TTR is termed ATTR. There are two types of TTR-related amyloidosis. One is ATTR wild-type (ATTRwt), in which amyloid from normal TTR protein is deposited over a period of decades. ATTRwt has a strong male predilection, and patients typically present between the ages of 60 years and 95 years with carpal tunnel syndrome preceding cardiac involvement. Because of its late presentation, ATTRwt is also referred to as senile amyloidosis. The second type of ATTR is due to mutant TTR (ATTRm) causing familial amyloidosis; patients are generally born with a pathological mutation in the TTR gene, resulting in accelerated breakdown of TTR to amyloid. Over 80 different amyloidogenic missense point mutations have been described. Val-30-Met is the most common pathogenic mutation globally and is the leading cause of familial amyloid polyneuropathy. Another important mutation, Val-122-Ile, is seen in 3% to 4% of African Americans and is the most common mutation leading to familial CA. Patients with ATTRm present at an earlier age (40 to 75 years) than those with ATTRwt and demonstrate a slight male predominance. The clinical manifestation of ATTRm is driven by the specific mutation, with phenotypes ranging from exclusive neurological involvement to CA with overlapping patterns; however, most affected patients have some degree of multisystem involvement. Chemotherapy has no role in ATTR. Clinical trials of various therapeutic agents that modify or inhibit amyloid fibril formation are in progress. Prognosis for ATTR is generally better than that for AL amyloidosis.
Arrhythmias in Cardiac Amyloidosis

Table 1: Types and Features of CA

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<td></td>
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Table 1: Types and Features of CA

CA: cardiac amyloidosis; AL: light chain amyloidosis; ATTRwt: wild-type transthyretin amyloidosis; TTR: transthyretin; VT: ventricular tachycardia; 99mTc-PyP: technetium pyrophosphate; 99mTc-DPD: technetium 3,3-diphosphono-1,2-propanodicarboxylic acid; AV: atrioventricular; IAA: isolated atrial amyloidosis. Adapted from Falk et al.1

The cardiac manifestations of amyloidosis include marked left ventricular (LV) thickening (due to amyloid fibril deposition) with electrocardiogram (ECG) voltage that is disproportionately low for the degree of LV thickening and associated diastolic (rather than systolic) heart failure. Clinical examination and echocardiography show restrictive cardiomyopathy features. Cardiac arrhythmias are common and are an important cause of morbidity and mortality. Diagnosis is based on imaging studies and biopsy. Cardiac magnetic resonance imaging (MRI) shows increased extracellular volume on T1 imaging, as well as patterns of late gadolinium enhancement in later stages.5 Bone-seeking radioisotopes have a high sensitivity for ATTR and are useful for the detection of subclinical CA when the MRI and ECG may still be unrevealing. 99mTc-pyrophosphate for myocardial imaging is readily available and is currently the tracer of choice for the diagnosis of TTR CA. It is also useful for differentiation from AL CA as there is minimal or no myocardial uptake of the tracer in AL CA.7 Endomyocardial biopsy typically shows deposition of amyloid in the extracellular and perivascular spaces that can be specifically stained with sulfated Alcian blue (Figure 1).

Pathophysiology of arrhythmias in cardiac amyloidosis

Cardiac involvement is invariable in ATTRwt. In the familial TTR amyloidoses (ie, ATTRm), the prevalence of CA varies with the specific mutation. AL amyloidosis may have minimal or severe cardiac involvement, with 50% of patients showing CA. Regardless of the cause of amyloid production, CA is characterized by extracellular amyloid deposition throughout the heart, including the cardiac conduction tissue and valves. The ventricles are non-dilated and thickened, with restrictive filling. Elevated filling pressures result in atrial dilation despite atrial wall thickening from amyloid deposition. Atrial arrhythmias and loss of atrial mechanical function with atrial thrombus formation are important risk factors for cardioembolic stroke.8,9
The histology of CA is amyloid infiltration of the extracellular spaces separating and distorting the myocardial cells. Myocardial scarring and patchy fibrosis that are typical of chronic ischemic cardiomyopathy or other non-ischemic cardiomyopathies are not described in CA. Hence, the exact mechanism of arrhythmias in CA is less well defined and is likely to be multifactorial. Small vessel disease due to perivascular amyloid infiltration associated with impaired vasodilation is a likely substrate for myocardial ischemia, especially in AL.10 Patients with CA occasionally report angina-type chest pains. Persistent cardiac troponin elevation is a feature of AL CA and, to a lesser extent, of ATTR. This increase in troponin may represent myocardial ischemia from small vessel occlusion or direct toxic effects of AL light chains. AL amyloid has been demonstrated to impair myocyte function and calcium release by increasing oxidative stress.11 In a zebrafish model, injecting AL light chains isolated from AL cardiomyopathy patients impaired cardiac function and led to cell death.12 Thus AL amyloidosis is a toxic, infiltrative cardiomyopathy. Inflammatory cell damage and separation of myocytes by amyloid fibrils would explain conduction abnormalities, atrial arrhythmias, and loss of atrial contractility. In the ventricles, non-sustained ventricular tachycardia (VT) is the most common form of arrhythmia. Sustained monomorphic VT is uncommon, as most unexpected lethal ventricular arrhythmias are due to polymorphic VT or ventricular fibrillation (see later).

TTR amyloidosis, especially ATTRwt, has a protracted course with a median survival of 60 months following presentation with heart failure symptoms.13 Conversely, AL amyloidosis has a high mortality rate once cardiac involvement becomes apparent. Several adverse prognostic indicators have been described, including elevated N-terminal pro-brain natriuretic peptide (NT-pro-BNP) and troponin levels, diastolic dysfunction on echocardiography, and the extent of extracellular volume and late gadolinium enhancement on MRI.14 In general, arrhythmia interventions are not likely to be beneficial in cases of severe cardiac involvement, as myocardial dysfunction leads to electromechanical dissociation, the predominant terminal event in patients with CA.

There are few systematic studies of electrophysiological abnormalities and arrhythmias associated with CA and their appropriate management. Most available data are based on case reports or series from single centers. The nature, incidence, and management of arrhythmias differ based on the type of amyloidosis and extent of cardiac involvement.

Conduction system disease and the role of cardiac pacing

The conduction system is affected in all forms of CA. Atrioventricular (AV) conduction delay or block is more common than sinus node disease: despite the high frequency of atrial involvement with amyloidosis, sinus node disease appears to be less common and relevant reports are mostly limited to isolated events in patients with ATTRm. In some reports, transient sinus node dysfunction occurred in association with autonomic dysfunction occurring spontaneously or during general anesthesia induction.15 An intracardiac electrophysiology (EP) evaluation of sinus node function in 25 patients with AL amyloidosis revealed normal sinus node function in 88% of those examined.16 Sayed et al. reported data from implanted loop recorders in 20 patients with Mayo stage III AL CA who were symptomatic with syncope or presyncope. Persistent sinus bradycardia with pauses requiring cardiac pacing was only detected in one patient.17

Conduction defects in the His-Purkinje system are more common and are associated with symptomatic AV block. First-degree AV block is often due to a delay in the His-Purkinje system level with preserved conduction at the nodal level. EP studies show an abnormal His bundle-ventricular (HV) interval in most patients with AL and ATTR CA.16,18 However, the incidence of symptomatic AV block is higher in ATTR, possibly because these patients are older and have better survival. In our series of 18 patients with advanced CA who underwent EP studies primarily for supraventricular arrhythmias, the mean (± standard deviation) HV interval was 87 ± 27 ms, despite a relatively narrow QRS duration (119 ± 32 ms).18 Prolongation of the HV interval with preserved QRS duration is well recognized in CA (Figures 2 and 3) and may represent diffuse amyloid infiltration of the bundles, creating equal delays in both the right and left branches, yielding a disproportionately narrow QRS.16

Approximately 25% to 35% of ATTRm and 45% of ATTRwt patients receive pacemaker implants. While cardiac pacing provides symptomatic relief, it does not change the

Figure 1: A high-powered view of a section of an endomyocardial biopsy in a patient with transthyretin amyloidosis stained with sulfated Alcian blue. The green staining is amyloid and the yellow staining is the myocardium. Note the extensive replacement of the myocardium by extracellular amyloid deposits.

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Figure 2: Intracardiac recording of a His bundle in a patient with ATTRwt cardiac amyloidosis and syncope. First-degree heart block is present with a PR interval of 230 ms. The PR prolongation is due to a conduction delay in the His-Purkinje system with an HV interval of 119 ms. Note the relatively narrow QRS duration of 98 ms despite marked prolongation of the HV interval. ATTRwt: wild type transthyretin amyloidosis; HV: His bundle–ventricular; QRS: QRS duration; RA: right atrium.

Figure 3: Hematoxylin and eosin staining of the region of the central fibrous body and His bundle in a patient who died of heart failure without a premortem diagnosis of cardiac amyloidosis. The patient had first-degree AV block but no higher grades of AV block. Autopsy showed extensive amyloid infiltration of the myocardium. The yellow circle represents the area of the His bundle. The pink amorphous eosin staining material marked by asterisks (*) represent amyloid deposits. The black arrow points to a small blood vessel with perivascular amyloid infiltration. CFB: central fibrous body; IVS: interventricular septum; LV: left ventricle; RA: right atrium; TV: tricuspid valve.

When the predominant rhythm is sinus, consideration should be given to atrial synchronized ventricular pacing, as maintenance of AV synchrony with atrial synchronized ventricular pacing may be important for maintaining cardiac output in the setting of diastolic dysfunction and preload dependence. A combination of peripheral vasoconstrictors and atrial pacing can be helpful in maintaining an adequate blood pressure in patients with severe autonomic neuropathy and chronotropic incompetence. When ventricular pacing is necessary, there is concern that dysynchrony from right ventricular (RV)-only pacing may cause further deterioration of ventricular function. Observations during echocardiography suggest improved cardiac output from biventricular pacing (personal communication). However, there are currently no formal studies to support this premise.

Atrial arrhythmias

Atrial fibrillation (AF) or atrial tachycardia (AT) is common in CA, especially as the disease progresses. In a group of patients from Sweden with familial amyloid polyneuropathy, 20% had previously undetected atrial arrhythmias suggesting possible amyloid atrioatrial, even in the absence of overt CA. Arrhythmia is more common in ATTRwt due to the older age at presentation and the higher prevalence of age-related AF. In one series, 62% of patients with ATTRwt CA had AF. AF or AT is often highly symptomatic and poorly tolerated, mostly due to rapid ventricular rates and an irregular ventricular response that impairs ventricular filling and contractility. Given that atrial contractility is often diminished in CA, the loss of atrial contribution is less likely to be the mechanism for deterioration. Ventricular rate control can be difficult since β-blockers and calcium blockers are not well tolerated due to hypotension and their negative inotropic effects. Although digoxin is known to bind amyloid tissue and increase the risk of toxicity, its use in low doses is usually tolerated, although its role in rate control is limited. Amiodarone provides a good option for AF control and is fairly well tolerated when administered orally. Alternative drugs such as dofetilide can be useful for maintaining sinus rhythm when used cautiously with close monitoring of the QT interval. Anticoagulation is usually well tolerated and should be used for all atrial arrhythmias. If there is evidence for non-contractility of the atria, there is a case to be made for anticoagulation even in sinus rhythm, as thromboembolism may occur with stasis of blood in the left atrium and appendage.

AF in amyloidosis tends to have longer cycle lengths and may appear organized. This is likely due to amyloid deposition separating out the atrial myocyte bundles and creating a marked delay in intra-atrial conduction. In our series, left atrial voltage mapping revealed significantly lower voltages in amyloid patients compared with an age-matched patient population with persistent AF. However, ablation attempts to correct what is often interpreted as left atrial tachycardia on surface
ECG often yield only limited success. Despite acute ter-
mination and restoration of sinus rhythm with ablation,
the recurrence rate is high. In our series, the recurrence
rate at one year was 83% for CA patients compared
with 25% in the non-CA, persistent AF patient group.18
In addition, procedural complications are higher in CA
patients with poor tolerance of fluid shifts and general
anesthesia. Acute chronic renal failure and pulmonary
congestion are common in the postprocedural period and
may require lengthy hospitalization for correction.

When the heart rate is difficult to control, AV nodal abla-
tion with ventricular pacing has the benefits of rate con-
rol and rate regularization that improve cardiac output
and offer symptomatic improvement.22 For ventricular
pacing following AV nodal ablation, our practice is to
aim for biventricular pacing or placement of the RV lead
in the high septal or para-His regions to minimize fur-
ther depression of ventricular function due to ventricular
dysynchrony from RV pacing.

Isolated atrial amyloidosis due to atrial natriuretic pep-
tide depositing as amyloid fibrils is seen in up to one-third
of patients with persistent AF undergoing valve surgery
and is unrelated to the atrial arrhythmias of generalized
CA. Isolated atrial amyloidosis is a disease of the elderly,
with a female preponderance.23

**Syncope**

Postural hypotension resulting from autonomic dysfunc-
tion is a common finding in AL amyloidosis and a cause
of syncope in this patient population. Diuretics tend to
aggravate symptoms. Midodrine, an alpha 1 adrenergic
stimulant, is often necessary in high doses to maintain
blood pressure. The supine hypertension that is com-
mon with midodrine use is not seen in CA, so large doses
can be employed without concern. Cardiac arrhythmias
such as heart block and ventricular arrhythmias are more
important causes of syncope and may be a harbinger of
sudden death, but careful history taking and examination can often differentiate one from the other. Ambulatory monitoring and EP studies can be helpful, especially for the detection of infra-Hisian AV block. Atrial arrhythmias with rapid ventricular rates can cause a precipitous drop in cardiac output, resulting in syncope. In advanced CA, another mechanism for syncope is vasodilation with an inability to increase cardiac output due to poor contractility reserve. Patients may present with exertional syncope. Prophylactic midodrine prior to planned exercise can be helpful.

**Ventricular arrhythmias**

As end-stage heart failure is the major driver of mortality in CA, managing ventricular arrhythmias has assumed a lesser role in overall outcome. AL amyloid, with its more precipitous downward course after heart failure onset, has a higher incidence of ventricular arrhythmia compared with that of ATTR disease. However, there are few systematic studies on the prevalence of ventricular arrhythmias in CA available to date. In a study of 195 patients with AL amyloidosis, 24-hour Holter recordings revealed non-sustained VT in 27% of patients with advanced AL CA. In this study, mortality was high (88%), suggesting a patient population in a late stage of the disease. The incidence of sudden death was no different between those with and without non-sustained VT, implying little prognostic importance regarding the finding of non-sustained VT alone. In a more recent study with implanted loop monitors in 20 patients with AL CA, non-sustained VT was observed in only one transmission. In both studies, recorded sudden deaths were mostly associated with terminal bradycardia followed shortly by pulseless electrical activity.

Monomorphic VT is occasionally inducible during EP study or is noted in patients with an implantable cardioverter-defibrillator (ICD), but it is an infrequent event in CA compared with other cardiomyopathies. Ventricular fibrillation storm provoked by monomorphic premature ventricular complexes of Purkinje origin was reported in two patients with possible AL amyloidosis, supporting the hypothesis of an inflammatory response to AL amyloid. Catheter ablation targeting the Purkinje fibers to trigger fibrillation is helpful in acute arrhythmia control.

**Sudden death and the role of implantable cardioverter-defibrillators**

The most frequent documented terminal event in CA is pulseless electrical activity or agonal bradycardia. Even in sudden death, the mechanism is commonly electromechanical dissociation from end-stage heart failure. As a result, there has been very little enthusiasm for the ICD in patients with CA. Arrhythmic sudden death is far more common in AL cardiomyopathy. However, the prognosis of AL amyloidosis with heart failure has historically been dismal, with survival lasting less than 12 months after diagnosis, representing a relative contraindication to ICD implantation. When life expectancy is greater than one year, secondary prevention ICD is a reasonable consideration, following a discussion of the risks and benefits. There is a tendency for a higher defibrillation threshold in patients with CA, but first shock efficacy is generally comparable to that reported in the SCD-Heart Failure Trial study. Primary prevention ICD implantation in CA patients is associated with a high rate of appropriate discharges for ventricular arrhythmias in AL patients (32% in the first year, in one series) but does not translate to improved survival. With improved treatment options for AL amyloidosis, it is time to revisit the issue of primary prevention ICDs. However, the prediction of risk for arrhythmic sudden death does not follow the available guidelines for other forms of heart disease. An ICD would be a reasonable consideration for unexplained syncope or recurrent episodes of non-sustained VT in the early stages of AL amyloidosis, when the levels of biomarkers such as troponin and NT-pro-BNP are low. Primary prevention ICD may also have a role in patients being considered for cardiac transplantation.

**Future directions and conclusions**

A combined multicenter effort to define the cardiac arrhythmias associated with CA is essential to gauge the survival benefits from early arrhythmic prophylaxis in CA. The formation of the Amyloid Research Consortium is an encouraging move in this direction (http://www.arc.org). Early pacemaker implantation in patients at high risk for heart block may also serve as arrhythmia monitoring with respect to data gathering. With the high rate of pacemaker implantation in ATTR amyloidosis, a trial of biventricular pacing in patients with heart block is feasible and important to clarify the role of ventricular synchrony in this unique form of cardiomyopathy. In the more malignant AL amyloidosis, the role of ICDs has thus far been assessed largely in the later stages, when severe heart failure negates any benefit from defibrillation. ICD should be evaluated in the earlier stages of AL CA, when arrhythmic sudden death is likely to be a greater risk.

There has been considerable progress in our understanding of amyloidosis pathophysiology and treatment in recent years. Drugs aimed at preventing new amyloid formation achieve very favorable responses in some patients, and formal trial results are awaited. Early diagnosis and management by experts are key to improving outcomes in CA. The disease can no longer be dismissed as a terminal illness with limited therapeutic options.

**References**


