Wild-type transthyretin cardiac amyloidosis (ATTRwt-CA), previously known as senile cardiac amyloidosis: Clinical presentation, diagnosis, management and emerging therapies

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Clinical case

A 72-year-old male who presented with progressive dyspnea on exertion down to 100 feet. His prior cardiac history includes severe aortic stenosis for which he underwent bovine bioprosthetic aortic valve replacement. He also had a bypass graft with left internal mammary artery to left anterior descending artery at the time of aortic valve replacement. Patient further had atrial fibrillation and flutter for which he underwent direct current cardioversion. His other chronic condition has been hypertension (HTN) for which he was on β-blockers and angiotensin converting enzyme inhibitors (ACE-i) but had worsening fatigue, dyspnea and low blood pressures of 100/60 requiring cessation of their use. Of note, patient also has been treated for peripheral neuropathy, of unknown etiology, with diffuse peripheral neuropathic pain and left focal ulnar neuropathy (at the elbow) as well as left carpal tunnel syndrome.

Based on the patient’s clinical presentation and prior cardiac history, he was initially evaluated by electrocardiogram (EKG) which showed normal sinus rhythm with first degree atrioventricular (AV) block, right bundle branch block, and poor R-wave progression. A pharmacological stress test (exercise treadmill was attempted but patient became dyspneic after 20 seconds) with myocardial perfusion imaging showed normal myocardial perfusion with no evidence of stress induced ischemia. An echocardiography (Echo) showing normal left ventricular (LV) size with moderate hypertrophy, low normal systolic function and grade III diastolic dysfunction as well as apical sparing by longitudinal strain mapping

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Abstract: Cardiac amyloidosis is thought to be a rare group of diseases caused by extracellular deposition of misfolded proteins in the extracellular cardiac matrix resulting in heart failure with preserved ejection fraction (HFpEF). This review focuses on the similarities and differences between the pathophysiology, clinical presentation and diagnostic tests of wild-type transthyretin cardiac amyloidosis (ATTRwt-CA) compared to immunoglobulin light chain amyloidosis and hereditary cardiac amyloidosis. We address some obstacles to timely diagnosis and opportunities for management of the clinical symptoms as well as possibility of future novel disease modifying therapies.

Keywords: ATTRwt cardiac amyloidosis (ATTRwt-CA); heart failure with preserved ejection fraction (HFpEF); left ventricular hypertrophy (LVH); transthyretin; senile amyloidosis

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β-blockers/ACE-i intolerance and peripheral neuropathies as well as normal stress test and Echo findings of moderate hypertrophy, grade III diastolic dysfunction and apical sparing by longitudinal strain mapping pattern it was suspected that patient may have infiltrative cardiac process contributing to his condition. This was further evaluated by cardiovascular magnetic resonance (CMR) showing diffuse subendocardial infiltration by gadolinium (Figure 2) and marked increase in T1 relaxation, which has consistent with likely infiltrative process such as cardiac amyloidosis. After discussion with patient, he wanted to proceed with endomyocardial biopsy (EMB) which showed positive Congo red staining with apple-green birefringence (Figure 3) and mass spectroscopy analysis of deposited endocardial amyloid fibrils showing transthyretin with normal amino acid sequence. The final diagnosis was most consistent with wild-type transthyretin cardiac amyloidosis (ATTRwt-CA). After the diagnosis of ATTRwt-CA, the patient was offered to be evaluated for participation in several of the ongoing clinical trials, testing novel therapies for cardiac amyloidosis, but he declined. The patient has been treated with conservative management for the past few years, focused on symptomatic control.

**Introduction and epidemiology**

The incidence and prevalence of congestive heart failure (CHF) is increasing throughout the world, especially in
aging population (1). In US, it affected 5.7 million adults in 2016 (2), 16 million in EU in 2016 (http://www.ehnheart.org) and 4.2 million in China in 2011 (publication still pending from the China Heart Failure Registry Study in 2015) (3). There are three types of CHF: (I) LV ejection fraction (LVEF) of less than 40%, called heart failure with reduced EF (HFrEF); (II) LVEF of 40–50%, called heart failure with mid-range EF (HFmrEF); and (III) LVEF greater than 50% called heart failure with preserved EF (HFpEF). The epidemiology, pathophysiology and outcomes of HFmrEF are poorly understood due to limited studies. While, HFrEF has higher incidence and prevalence in males (4), with more than 60% caused by coronary artery disease (5), HFpEF is more common in women with incidence that increases with aging resulting in higher prevalence in older adults (4). The pathophysiology of HFpEF are poorly understood and are likely to be due to numerous factors resulting in a heterogeneous patient population. It is possible that due to the heterogeneity in patient population and conditions contributing to HFpEF there have been numerous negative clinical trials aimed at potential treatment strategies (6).

Cardiac involvement of systemic amyloidosis is one of the causes of HFpEF. Systemic amyloidosis is thought to be a rare group of diseases, as it is likely underdiagnosed, and is characterized by progressive tissue infiltration of insoluble fibrillar proteins resulting in extracellular architecture disruption that causes organ dysfunction. There are three main types of systemic amyloidosis that are known to cause cardiac dysfunction and are identified by the amyloid fibril protein deposited in the heart, namely: (I) immunoglobulin light chain (AL) amyloidosis, (II) ATTRwt-CA, previously known as senile cardiac amyloidosis; and (III) a mutated ATTR causing hereditary cardiac amyloidosis (ATTRm-CA). AL amyloidosis is most commonly diagnosed amyloidosis due to the dysfunction it causes to organs, other than the heart, by the amyloid light chain deposition. The diagnosis of ATTRm and ATTRwt cardiac amyloidosis is more elusive as the clinical presentation and severity of organ dysfunction is gradual and variable. Studies have shown that 10–25% of patients with HFpEF have LV ATTRwt deposition at the time of autopsy (7-9).

In this review, our aim was to summarize the main pathophysiology changes caused by ATTRwt amyloidosis, their clinical presentation and diagnostic evaluation. The early identification and diagnosis of this condition can result in better current symptomatic management and identification of novel emerging therapies aimed at altering the progression and hopefully the prognosis of this condition.

Pathophysiology and clinical features

ATTRwt-CA is likely significantly underdiagnosed. Although it thought to affect 10–25% of patients with HFpEF, suggesting a high prevalence in the elderly population, there are less than 100 cases annually at the National Amyloidosis Center in UK (10) resulting in limited understanding of the natural history of the disease due to small patient cohorts. The reason for the low diagnostic rate is likely due to the nature of the disease pathophysiology that results in a subtle clinical presentation in an elderly population who also has frequent comorbid conditions and the belief that diagnosis will not affect the patients’ outcomes.

In contrast to other forms of amyloidosis, ATTRwt-CA almost exclusively causes clinically evident pathophysiology to the heart and the peripheral nerves as well as non-clinically relevant deposits in the liver and lungs. The cardiac manifestations of amyloid wild-type amyloid proteins deposition are LV hypertrophy (LVH) with increased filling pressure and the “typical” HFpEF with symptoms of breathlessness, reduced exercise tolerance, fatigue, lower extremities/abdominal swelling, early satiety and erectile dysfunction. However, the constellation of these cardiac signs and symptoms are very similar to other conditions affecting the ATTRwt-CA patient population of mostly males who are often greater than 65 years of age (11), such as HTN, coronary artery disease, aortic stenosis, diabetes, arrhythmias, obstructive sleep apnea and/or obesity. Hypertrophic cardiomyopathy or hypertensive heart disease are also close mimickers of ATTRwt-CA clinically and by traditional Echo imaging findings, leading to frequent misdiagnosis (12). Additionally, patients at that age group often have concomitant monoclonal gammopathy of undetermined significance (MGUS) (13) and the extracardiac ATTRwt deposition results in bilateral carpal tunnel syndrome (14) that precedes cardiac manifestations leading to evaluations by non-cardiovascular specialists and delay from definitive diagnosis.

High index of suspicion is necessary for the timely diagnosis and treatment of ATTRwt-CA. The key factors that may help increase the sensitivity for diagnosis of ATTRwt-CA are: LVH without presence of prior history of severe or uncontrolled HTN, new onset of hypotension, inability to tolerate ACE-i or β-blockers, atrial
arrhythmias (15,16), AV nodal dysfunction (13) requiring permanent pacemakers and the presence of bilateral carpal tunnel syndromes (14) and/or spinal stenosis. Right heart failure symptoms are also associated with ATTRwt-CA with persistently elevated internal jugular veins, hepatic congestion, ascites, abdominal bloating with early satiety and chronic lower extremity edema.

Diagnosis

The definitive diagnosis of ATTRwt-CA is made by histology of cardiac biopsy and subsequent genotyping. However, EMB is an invasive procedure with associated potential risks and is not readily available in many centers. There are several non-invasive testing modalities that have variable sensitivities and specificities for the diagnosis of ATTRwt-CA, including EKG, biomarkers, Echo, scintigraphy, CMR, as well as liver and fat pat biopsies that should be discussed.

EKG

EKG is one of the oldest, best described and most widely used cardiac diagnostic modalities worldwide. However, its use as diagnostic tool for ATTRwt-CA has likely lead to falsely reduced clinical suspicion and its underdiagnoses of the disease. This is largely from the “classical” teaching which dictates that cardiac amyloidosis is associated with low voltage (defined as ≤5 mV amplitude in the limb leads or ≤10 mV amplitude in the precordial leads) despite the presence of L VH and comes from studies of AL amyloid (16,17). In recent ATTRwt-CA studies, low voltage has been found to have poor independent sensitivity (~30%) for its diagnosis (18,19).

There are numerous other EKG findings that are seen in patients with ATTRwt-CA but have low sensitivity and specificity for the disease because they are also common in age and comorbidities matched patient cohorts: (I) 38–63% have pseudoinfarct pattern (17,19); (II) 34% have poor R-wave progression (19); (III) 16–80% have atrial fibrillation (15,18); (IV) 21% have first degree AV block (20); and (V) 16% have nonspecific ST-T-wave abnormalities (20). Left bundle branch block can be potentially useful to differentiate ATTRwt-CA and AL-CA as it occurs in 40% vs. 4%, respectively (14).

Voltage-to-mass ratio, calculated by sum of S wave in lead V1 plus R wave in lead V5 or V6 (SV1 + RV5 or V6) divided by the echocardiographic muscle cross-sectional area (21), has been shown to have high sensitivity and specificity for ATTRwt-CA (14) as compared to ATTRm-CA, AL-CA and other age related cardiac conditions. Additionally, patients with HTN or LVH usually have normal or increased EKG voltage while patients with tamponade, pericardial effusion or emphysema usually have low EKG voltages with normal LV mass (21).

Echo

Echo is the most frequently used noninvasive imaging test for evaluation of suspected cardiomyopathies, like ATTRwt-CA, as it provides both structural and functional information with minimal risk to the patient. Two-dimensional (2D) imaging, pulse wave and tissue Doppler velocities findings in ATTRwt-CA by Echo are similar to other forms of amyloidosis and other causes of HFpEF, like HTN, obesity, hypertrophic cardiomyopathy, Fabry’s disease and aortic stenosis.

The most common 2D finding in cardiac amyloidosis is the presence of increased left ventricular thickness or LVH, however, unlike the other forms of HFpEF mentioned above, the LVH associated with cardiac amyloidosis is not due to cardiac myocyte hypertrophy but rather due to amyloid fibrils deposited in the extracellular matrix. The increase in LV wall thickness associated with ATTRwt-CA is significantly greater than either AL (13) and ATTRm-CA (14). Another 2D finding that is associated with extracellular deposition of amyloid fibrils is a visually discernable “speckled” or granular “sparkling” texture appearance that seen by initial Echo machines, however, it is now less evident with modern echocardiographic machines due to almost exclusive use of harmonic frequency imaging.

Diastolic dysfunction with preserved EF is a hallmark of cardiac amyloidosis by Echo. It is characterized by the evidence of grade I–II mitral valve inflow Doppler patterns with concomitant reduction of S’ and e’ tissue Doppler velocities at the basal septal and lateral walls and impaired systolic dysfunction assessed by myocardial strain (22,23) at the early stages of the disease. Forty percent of ATTRwt-CA of patients have moderate LV systolic dysfunction compared to 8–22% patients with ATTRm-CA and AL-CA (14). At the later stages, ATTRwt-CA patients develop severe diastolic dysfunction characterized by grade III mitral valve inflow pattern associated with very low S’ve’ velocities, and eventually reduction of EF (24). In fact, the development of systolic dysfunction may be present with preserved EF and predictive of poor outcomes in patients with cardiac
amyloidosis, measured by a reduction of myocardial contraction fraction (MCF), which is the calculated stroke volume divided by myocardial volume (25). Other non-specific findings with ATTRwt-CA are: (I) biatrial enlargement; (II) thickening of the mitral/aortic leaflets and intra-atrial septum; (III) small pericardial effusion; (IV) moderate pulmonary HTN; and (V) intracardiac thrombus (22,26).

Advanced echocardiographic techniques, such as strain and strain rate imaging, have shown promise to help differentiate cardiac amyloidosis from hypertrophic cardiomyopathy or other causes of LVH. Strain and strain rate are echocardiographic techniques that use vendor specific proprietary software to track the longitudinal shortening of speckles within the 2D image of the left ventricular wall during systole and are quickly becoming clinically readily available. With all forms of cardiac amyloidosis, there is much greater restriction of basal speckle longitudinal movement compared to apical movement resulting in a relative ‘apical sparing’ pattern of longitudinal strain with bulls-eye plot (Figure 1) (27). Furthermore, Pagourelias et al. reported that EF global longitudinal strain ratio (EFSR), calculated by ratio of left ventricular EF and global longitudinal strain, had nearly 90% sensitivity and 92% specificity for all form of cardiac amyloidosis in the challenging subgroups with left ventricular wall thickness <16 mm with preserved EF (28).

**Cardiac biomarkers**

Brain natriuretic peptide (BNP), N-terminal fragment of BNP (NT-proBNP) and cardiac troponins have been widely used to help with diagnosis, monitor treatment and prognostic evaluation both HFrEF and HFpEF. Elevation of NT-proBNP above >82 pg/mL has been shown to have sensitivity of 92% and specificity of 90% for presence of left ventricular abnormalities in patients with familial mutation in ATTR thus could be used as a potential screening test to initiate workup for ATTRm-CA (29). This observation may translate to patients with ATTRwt-CA in whom the log of BNP has been shown to have direct correlation with left ventricular thickness (30). Increasing serum levels of BNP, NT-proBNP and troponins were shown to have progressive correlation with disease severity and worse outcomes at increasing levels in patients with ATTRwt-CA (31). Furthermore, monitoring of these cardiac biomarkers is being employed to monitor treatment in ongoing clinical trials using disease modifying approaches for the treatment of ATTR cardiac amyloidosis (see “investigational medications” section later in the article).

**Nuclear imaging**

Nuclear imaging has emerged as important non-invasive tool in the diagnosis of suspected ATTR cardiac amyloidosis due to wide availability at a low cost, having few contraindications and the capacity to differentiate from other cardiomyopathies. Bone-avid tracers, like 99mTc-DPD (technetium-3,3-diphosphono-1,2-propanodicar-boxylic acid), 99mTc-PYP (technetium-pyrophosphate) and 99mTc-HMDP [technetium-hydroxymethylene diphosphonate (Tc-HMDP)] have been shown to have high sensitivity and specificity for differentiating patients with ATTR CA, irrespective of genotype, from patients with AL-CA or others with HFpEF (32-37). The exact mechanism by which these radiotracers differentially accumulate in myocardium is not completely clear but may be due to differences in deposited amyloid proteins (38,39), higher calcium levels seen during the repair process (40) and/or higher degree of tissue microcalcifications in ATTR compared to AL cardiac amyloidosis (41). Irrespective of the mechanism, an international consensus document has confirmed that the combination of grade 2 or 3 cardiac uptake on a bone-avid tracer scan in the setting of absent monoclonal protein by serum immunofixation electrophoresis (IFE), urine IFE, and serum free light chain assay is diagnostic of ATTR cardiac amyloidosis as compared to AL-CA or other wall thickening diseases (42). Bokhari et al. (37) described a standardized imaging protocol using 99mTc-PYP to diagnose ATTR CA using Heart/Contralateral ratio ≥1.5. Nuclear imaging with 99mTc-DPD (43) and 99mTc-PYP (44) can diagnose the presence of cardiac involvement prior to any overt echocardiographic abnormalities and can predict major adverse cardiac events (34,35,39,43). Future studies are needed to establish the role of bone-avid nuclear tracers for the early identification of ATTRwt-CA, differentiating it from ATTRm-CA (confirmed by genetic testing) and evaluation of subsequent outcomes.

Scintigraphy imaging, which is inherently qualitative, falls short of being able to quantify the radioactivity at the affected sites thus cannot be used in assessing disease burden and response to therapy. Positron emission tomography (PET) is a nuclear modality that can circumvent this problem, emerging as a promising tool in the monitoring and management of cardiac amyloidosis. Recently discovered
amyloid binding PET tracers 18-F florbetapir (45,46) and 11C-Pittsburgh compound B (PiB) (47,48) can identify both AL and ATTR cardiac amyloidosis. Case reports have shown cardiac uptake of another PET tracer [18F]-sodium fluoride only in patients with ATTRwt-CA and ATTRm-CA, but not in ones with AL-CA (49,50).

Cardiovascular magnetic resonance

CMR is an important diagnostic and prognostic tool for cardiac amyloidosis. Various techniques utilizing different tissue imaging timing, with and/or without gadolinium contrast, and strain can provide detailed information about the presence, location, and distribution of hypertrophy, as well as cardiac function. One major drawback of using CMR is that gadolinium contrast is contraindicated in patients with moderate to severe kidney disease.

T1 sequence of CMR, a composite measure of extracellular matrix and myocardial cells, with the use of gadolinium can help differentiate extracellular tissue thickening due to myocardial hypertrophy vs. extracellular deposition. Utilizing pre- and post-contrast T1 mapping, extracellular volume (ECV) can be calculated and is a direct measurement of the cardiac interstitium (51). ECV expansion can detect amyloid fibrils infiltration in AL and ATTR cardiac amyloidosis earlier than conventional testing and is quantitative marker of the amyloid burden lending itself with the potential use in early diagnosis and disease monitoring (52,53). Marked increased non-contrast T1 relaxation times also seen in patients who have interstitial infiltration by amyloid fibrils and has good correlation with disease severity, future prognosis, and the potential to track changes over time (from natural progression or disease modifying therapy) (54-56). One of the major advantages of T1 mapping is that it does not require contrast which is contraindicated in advanced kidney disease, however, currently this technique has limited clinical availability due to technical challenges related to sequence- and vendor-specific differences (57,58).

The myocardial deposition of amyloid fibrils in ATTR-CA increases ECV which serves as a reservoir for gadolinium accumulation leading to characteristic continuum of late gadolinium enhancement (LGE) (59). In the early stages of the disease there usually is no LGE enhancement that progresses to subendocardial and finally transmural (in non-ischemic pattern) LGE enhancement at the late stages (Figure 2), which also tracks with increasing ECV and a worse prognosis (58,60). In contrast, a diffuse-subendocardial patterns is most commonly described in AL-CA disease (58,60), which has led to CMR based LGE scoring system that seems to differentiate between ATTR and AL cardiomyopathies (61). Future studies are necessary to establish similarities and differences between the LGE in patients with ATTRm-CA and ATTRwt-CA.

Traditional LGE imaging technique depends on normal myocardium to enhance the diseased area and has been limited in ATTR-CA due to difficult nulling, which have led to early and advanced disease misclassification. Phase-sensitive inversion recovery (PSIR) sequence that reduces the need for an optimal null point setting [initially described CMR evaluation of myocardial infarction (62)] makes LGE assessment in cardiac amyloidosis faster and operator-independent (58).

Feature tracking software applied to cine CMR datasets for assessment of left ventricular strain has shown good agreement between CMR and 2D Echo-derived myocardial longitudinal strain measurements (63) with apical sparing in longitudinal strain (64).

Histopathological diagnosis

EMB with histopathology remains the gold standard for diagnosis of cardiac amyloidosis by showing deposition of amorphous deposits of amyloid fibrils in the heart. The binding of Congo red stain to the deposited amyloid fibrils leads to characteristic apple-green birefringence under polarized light microscopy (Figure 3) and an intense yellow-green fluorescence is seen when binding to thioflavin (image not shown). Precursor protein identification can be accomplished by immunohistochemistry, electron microscopy, or mass spectrometry (preferred), depending upon institutional expertise. Fat pat biopsy with Congo red staining has sensitivity of 70–90% for diagnosis of systemic AL amyloidosis (65,66), but only 45% and 15% sensitivity for diagnosis of ATTRm-CA and ATTRwt-CA respectively (67). Adjunctive laboratory tests to rule out AL include assaying for other organ dysfunction (e.g., proteinuria, alkaline phosphatase) and directly measuring the circulating light chains in plasma.

EMB is an invasive procedure that is associated with risks of complications including ventricular free-wall perforation of up to 0.4%, arrhythmia 0.5–1.0%, conduction abnormalities 0.2–0.4% (68) and requires expertise that is not readily available in many centers. With recent advance in Echo, CMR and nuclear imaging, multiple diagnostic
Halatchev et al. Wild-type transthyretin cardiac amyloidosis

algorithms have been proposed for patients with suspected ATTRwt-CA (12,42,59,69,70).

Genetic testing

Once diagnosis of ATTR is confirmed via EMB with histology, it is important to offer DNA sequencing for the transthyretin gene (especially in clinical research studies) since the presence of a pathologic mutation can affect clinical trial options, predict sites of organ involvement, and have relevance for family members. The absence of amino acid sequence abnormalities by mass spectrometry is consistent with ATTRwt-CA, although mass spectrometry cannot detect all transthyretin mutations.

Treatment

Limitations in treatment due to insufficient evidence-based approaches

While there are great advances in the treatment of AL amyloidosis by chemotherapy (71,72) and emerging new therapies for ATTRm-CA aimed at the reduction of mutant gene expression (73), TTR tetramer stabilization (74,75) and dissolution (76), currently there are no proven curative or disease modifying therapies for ATTRwt-CA. There are several possible reasons why ATTRwt-CA does not have any proven therapies: (I) it is a underrecognized and underdiagnosed leading to small patient cohorts to study and incomplete understanding of its natural history of the disease (13); (II) the disease presents late in life with multitude of age related comorbidities and mortality (13,14); and (III) the disease process is likely gradual thus curative or disease modifying therapeutic effects are difficult to measure (13). Consequently, most of the therapeutic considerations for ATTRwt-CA are based on expert opinion and observations from therapeutic effects on the other type of amyloidosis affecting the heart.

Symptomatic relief and supportive care

The mainstay of therapy is supportive care aimed at symptomatic relief. The main symptoms of patients with ATTRwt-CA are congestion, fatigue, peripheral nerve pain, and hypotension. Relief of congestive symptoms and associated fatigue can be accomplished with the use of diuretics, including loop diuretics and thiazides in combination with mineralocorticoid receptor antagonist to help with potassium reabsorption. Diuretics should be used sparingly as these patients are preload dependent due to high filling pressures and can be used with weight-based parameters. Other medications typically used in cardiomyopathies, such as β-blockers, ACE-i and angiotensin receptor blockers (ARB), do not seem to modify the disease progression and often result in worsening fatigue and hypotension. B-blockers are usually not tolerated as they can reduce the inotropy of the heart resulting in decreased stroke volume and have a maladaptive blunting of chronotropy on which ATTRwt-CA patients relay for augmentation of their cardiac output. ACE-i and ARB medications often worsen hypotension in ATTRwt-CA patients due pre-existing peripheral neuropathy affecting the autonomic nervous system of these patients. Hypotension can be managed with α-1 blocker midodrine and compression stockings.

Atrial arrhythmias

The symptoms associated with ATTRwt-CA are further exacerbated by atrial arrhythmias which are frequently associated with the condition (15). Rate control is difficult in these patients as β-blockers can worsen hypotension at higher doses, calcium channel blockers are contraindicated as they bind to the amyloid fibrils causing sustained worsening of CHF (77-79) and digoxin can cause cardiac toxicity due to progressive accumulation in the amyloid rich heart despite normal serum levels (80). Antiarrhythmic medications, such as amiodarone, can be used for rhythm control as patients are older and is overall well tolerated because it is usually not associated with deleterious hemodynamic changes. Symptomatic relief from atrial arrhythmias has been reported in a small cohort of ATTRwt-CA patients with catheter ablation (81). However, catheter ablation for atrial arrhythmias have high recurrence rate, necessitating AV ablation with permanent pacemaker placement in refractory cases.

Investigational medications

Currently there are no known curative or disease modifying agents for the treatment of ATTRwt-CA. However, there are several investigational medications that may have the potential to modify or even reverse the disease process. The strategies employed are ATTR disruption to reduced amyloid fibril aggregation, ATTR destabilization, and ATTR suppression. The combination of bile acid
Cardiac infiltration with amyloid fibrils can precipitate conduction system abnormalities, like bundle branch blocks and AV node dysfunction (84). A significant portion of patients with ATTRwt-CA have pacemakers at the time of diagnosis or have one implanted for high degree AV block (13). ACC/AHA guidelines suggest a case by case consideration for implantable cardioverter-defibrillator (ICD) implantation in patients for primary and secondary prevention due to limited clinical trials to show associated outcomes (85).

**Permanent pacemaker and defibrillator implantation**

Cardiac infiltration with amyloid fibrils can precipitate conduction system abnormalities, like bundle branch blocks and AV node dysfunction (84). A significant portion of patients with ATTRwt-CA have pacemakers at the time of diagnosis or have one implanted for high degree AV block (13). ACC/AHA guidelines suggest a case by case consideration for implantable cardioverter-defibrillator (ICD) implantation in patients for primary and secondary prevention due to limited clinical trials to show associated outcomes (85).

**Advanced therapies: left ventricular assist devices (LVAD) and heart transplantation**

ATTRwt-CA patients have advanced heart failure symptoms and reduced life expectancy, findings which may qualify them to be considered for advanced heart failure options such as LVAD and/or heart transplantation. In a small single center report of 9 patients, LVAD implantation is technically feasible for ATTRwt-CA but it was associated with higher 24-month mortality and morbidity compared to other indications for LVAD implantation (86). A review of the Interagency Registry for Mechanically Assisted Circulatory Support (INTERMACS) database is necessary to be able to better evaluate outcomes of LVAD implantation for ATTRwt-CA. Patients with ATTRwt-CA are usually not considered for heart transplantation due to their advanced disease presentation, age, and associated comorbidities. Indeed, in a literature review there was a single case report found of a 68-year-old patient who underwent heart transplantation for biopsy proven ATTRwt-CA who was reported to have a good 3-year outcome (87,88).

**Conclusions**

Cardiac amyloidosis is a group of diseases caused by extracellular deposition of misfolded proteins. ATTRwt-CA is currently thought to have lowest prevalence among the different types of systemic amyloidosis, which is likely due to a low incidence from underdiagnosis, as it is found in 10–25% of all patients with HFpEF on autopsy studies. High clinical suspicion is necessary to initiate discussion with the affected patient about the risk, cost and benefit of early diagnosis and potential management, both of which often requiring a referral to a tertiary center. The total cost of testing, including EKG, Echo, CMR, and EMB is $5,000 to $15,000 (the variable cost to the patient) with the potential risks inherent to of each procedure. However, as in many rare or underdiagnosed conditions, an accurate diagnosis and appropriate level of follow up is needed to increase the patients able to participate in goal directed therapies and improve our understanding of the disease process as well as future possible treatments. Patients with signs and symptoms of heart failure (especially right sided symptoms), LVH out of proportion of duration and severity of HTN, arrhythmias, bundle branch blocks, have extracardiac manifestations of carpel tunnel syndrome and show intolerance to common cardiac medications should be evaluated for presence of ATTRwt-CA. Initial, diagnostic evaluation the highest sensitivity followed by ones with highest specificity should be performed to help diagnoses ATTRwt-CA: (I) Echo strain patterns with characteristic apical sparing and reduction of ratio of stroke volume to myocardial volume; (II) quantitative and qualitative characterization by nuclear scintigraphy; and (III) CMR tissue characterization with LGE, ECV and T1 mapping. Only if those non-invasive modalities are highly suggestive of ATTRwt-CA, EMB, followed by genetic testing should be performed as the diagnostic gold standard as it is not readily available and is invasive. Once patients are diagnosed to have ATTRwt-CA clinical consideration should be taken to adjust their medical therapy to best support their unique pathophysiology and patients should be considered to participate in the ongoing or upcoming clinical trials testing possible novel disease modifying therapies.
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Footnote

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