

ASNC/AHA/ASE/EANM/HFSA/ISA/SCMR/SNMMI Expert Consensus Recommendations for Multimodality Imaging in Cardiac Amyloidosis: Part 1 of 2—Evidence Base and Standardized Methods of Imaging

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Preamble

Cardiac amyloidosis is a form of restrictive infiltrative cardiomyopathy that confers significant mortality. Because of the relative rarity of cardiac amyloidosis, clinical and diagnostic expertise in the recognition and evaluation of individuals with suspected amyloidosis is mostly limited to a few expert centers. Electrocardiography, echocardiography, and radionuclide imaging have been used for the evaluation of cardiac amyloidosis for over 40 years.^{1–3} Although cardiovascular magnetic resonance (CMR) has also been in clinical practice for several decades, it was not applied to cardiac amyloidosis until the late 1990s. Despite

an abundance of diagnostic imaging options, cardiac amyloidosis remains largely underrecognized or delayed in diagnosis.⁴ Although advanced imaging options for noninvasive evaluation have substantially expanded, the evidence is predominately confined to single-center small studies or limited multicenter larger experiences, and there continues to be no clear consensus on standardized imaging pathways in cardiac amyloidosis. This lack of guidance is particularly problematic given that there are numerous emerging therapeutic options for this morbid disease, increasing the importance of accurate recognition at earlier stages. Imaging provides noninvasive tools for follow-up of disease

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remission/progression complementing clinical evaluation. Additional areas not defined include appropriate clinical indications for imaging, optimal imaging utilization by clinical presentation, accepted imaging methods, accurate image interpretation, and comprehensive and clear reporting. Prospective randomized clinical trial data for the diagnosis of amyloidosis and for imaging-based strategies for treatment are not available. A consensus of expert opinion is greatly needed to guide the appropriate clinical utilization of imaging in cardiac amyloidosis.

Introduction

The American Society of Nuclear Cardiology (ASNC) has assembled a writing group with expertise in cardiovascular imaging and amyloidosis, with representatives from the American College of Cardiology (ACC), the American Heart Association (AHA), the American Society of Echocardiography (ASE), the European Association of Nuclear Medicine (EANM), the Heart Failure Society of America (HFSA), the International Society of Amyloidosis (ISA), the Society of Cardiovascular Magnetic Resonance imaging (SCMR), and the Society of Nuclear Medicine and Molecular Imaging (SNMMI). This writing group has developed a joint expert consensus document on imaging cardiac amyloidosis, divided into 2 parts. Part 1 has the following aims:

- 1) Perform and document a comprehensive review of existing evidence on the utility of echocardiography, CMR, and radionuclide imaging in screening, diagnosis, and management of cardiac amyloidosis.
- 2) Define standardized technical protocols for the acquisition, interpretation, and reporting of these noninvasive imaging techniques in the evaluation of cardiac amyloidosis.

Part 2 of this expert consensus statement addresses the development of consensus diagnostic criteria for cardiac amyloidosis, identifies consensus clinical indications, and provides ratings on appropriate utilization in these clinical scenarios.

Purpose of the Expert Consensus Document

The overall goal of this multi-societal expert consensus document on noninvasive cardiovascular imaging in cardiac amyloidosis is to standardize the selection and performance of echocardiography, CMR, and radionuclide imaging in the evaluation of this highly morbid condition, and thereby improve health-care quality and outcomes of individuals with known or suspected cardiac amyloidosis. We hope that research generated to validate the recommendations of this consensus document will form the basis for evidence-based guidelines on cardiac amyloidosis imaging within the next few years.

Overview of Cardiac Amyloidosis

Cardiac amyloidosis is a cardiomyopathy that results in restrictive physiology from the myocardial accumulation of misfolded protein deposits, termed amyloid fibrils, causing a clinically diverse spectrum of systemic diseases. Most cases of cardiac amyloidosis result from 2 protein precursors: amyloid immunoglobulin light chain (AL), in which the misfolded protein is a monoclonal immunoglobulin light chain typically produced by bone marrow plasma cells, and amyloid transthyretin (ATTR) amyloidosis, in which the misfolded protein is transthyretin (TTR), a serum transport protein for thyroid hormone and retinol that is synthesized primarily by the liver.³ ATTR amyloidosis is further subtyped by the sequence of the TTR protein into wild-type (ATTRwt) or hereditary (ATTRv), the latter resulting from genetic variants in the *TTR* gene.^{5,6} Cardiac involvement in systemic AL amyloidosis is common (up to 75%, depending on diagnostic criteria),⁷ and in the case of ATTRwt amyloidosis, is the dominant clinical feature seen in all cases.

The different types of cardiac amyloidosis display significant heterogeneity in clinical course, prognosis, and treatment approach.⁸ AL amyloidosis is characterized by a rapidly progressive clinical course, and if untreated, the median survival is less than 6 months. ATTRv amyloidosis follows a varied clinical course depending upon the specific mutation inherited with either cardiomyopathy and/or sensory/autonomic polyneuropathy.⁹ Furthermore, ATTR amyloidosis (both wild-type and hereditary) is characterized by an age-dependent penetrance, with the clinical phenotype developing as age advances.

The diagnosis of cardiac amyloidosis remains challenging owing to a number of factors, which include the relative rarity of the disease, clinical overlap with more common diseases that result in thickening of the myocardium (ie, hypertension, chronic renal failure, hypertrophic cardiomyopathy, aortic stenosis), unfamiliarity with the proper diagnostic algorithm, and a perceived lack of definitive treatment. Although systemic AL amyloidosis is indeed a rare disease affecting ~8 to 12^{10,11} per million person years, and as high as 40.5 per million person years in 2015,¹² ATTRwt cardiac amyloidosis appears quite common, with recent reports using contemporary diagnostic strategies that place the prevalence in as many as 10% to 16% of older patients with heart failure or with aortic stenosis.^{13–15} In addition, the most common mutation associated with ATTRv amyloidosis has been reproducibly demonstrated in 3.4% of African Americans.¹⁶ Although the penetrance remains disputed, this suggests there are ~2 million people in the United States who are carriers of an amyloidogenic mutation and are at risk for cardiac amyloidosis. It is clear that both ATTRv and ATTRwt cardiac amyloidosis are under-recognized, yet important causes of diastolic heart failure.¹⁷

Treatment options are rapidly expanding. Anti-plasma cell therapeutics have extended median survival in AL amyloidosis beyond 5 years,⁷ with increasing survival beyond

10 years. We are potentially nearing a similar sea change in the management of ATTR amyloidosis. ATTR amyloidosis was previously only treated by solid-organ transplantation, because conventional highly effective heart-failure therapy is poorly tolerated and contraindicated in advanced cardiac amyloidosis. Although early clinical trials of amyloid-specific antibodies have been unsuccessful to date,^{18–20} one remains under study in a Phase I clinical trial.²¹ Novel therapeutics that suppress TTR expression have been studied in Phase 3 clinical trials and received FDA approval^{18,19} for ATTRv with polyneuropathy. Additionally, a randomized clinical trial of TTR stabilizer therapy demonstrated a reduction in all-cause mortality in ATTR cardiomyopathy²²; this agent has recently received FDA approval for ATTR cardiomyopathy. As these exciting prospects move into the clinical realm, it is evident that early diagnosis will be essential to afford the most effective treatment options for both AL and ATTR cardiac amyloidosis.

Biomarkers and Biopsy in Cardiac Amyloidosis

Despite these advances in treatment, the challenge persists to increase recognition and achieve effective, timely diagnosis. In the past, a diagnosis of cardiac amyloidosis required an endomyocardial biopsy, which remains the gold standard, because it is virtually 100% accurate, assuming appropriate sampling, for the detection of amyloid deposits.²³ Specific identification of the precursor protein can be accomplished from the tissue specimen through immunohistochemistry, albeit with limitations,²⁴ or laser-capture tandem mass spectrometry (LC/MS/MS). This latter technique is considered the definitive test for precursor protein identification.²⁵ Although ATTR cardiac amyloidosis can now be diagnosed accurately without the need of cardiac biopsy,³ AL amyloidosis requires demonstration of light-chain amyloid fibrils in tissue (although not necessarily the heart) prior to administration of chemotherapy. Even for ATTR cardiac amyloidosis, a cardiac biopsy remains necessary in the context of equivocal imaging or the coexistence of a monoclonal gammopathy.

Clinical suspicion of cardiac amyloidosis can be raised by the constellation of clinical signs and symptoms, specific demographics (ie, age, race, country of family origin), electrocardiography, and suggestive noninvasive imaging findings. Endomyocardial biopsy, although highly sensitive (100%),²³ is impractical as a screening test for cardiac amyloidosis, given its inherent risk and requirement of pathologic expertise, which is limited to a few academic centers. Other limitations of endomyocardial biopsy include the following: inability to quantify whole-heart amyloid burden, inability to evaluate systemic disease burden, and, for these same reasons, limited assessment of response to therapy. Thus, contemporary imaging techniques, including CMR, radionuclide imaging with bone-avid radiotracers, and echocardiography with longitudinal strain quantification, have evolved as the principal means for diagnosis and management of cardiac amyloidosis.

The current diagnostic approach for cardiac amyloidosis involves the use of one or more of these imaging modalities

in conjunction with assessment of a plasma-cell disorder (Fig. 1).³ Serum plasma electrophoresis is an insensitive test for AL amyloidosis and thus is unreliable for diagnosing AL amyloidosis. Serum and urine immunofixation and the measurement of serum free light chains (FLCs) are necessary for the diagnosis of AL amyloidosis. In cases of confirmed ATTR amyloidosis, *TTR* gene sequencing is performed to establish ATTRwt versus ATTRv. In AL amyloidosis, the concentration of the affected FLC, in conjunction with serum N-terminal pro-brain natriuretic peptide (NT-proBNP) and cardiac troponin T or I, can be utilized to assign a disease stage that confers highly reproducible prognostic information.²⁶ Furthermore, a cardiac staging system based on NT-proBNP and cardiac troponins (along with differential FLC levels) allows the stratification of patients into stages widely used in clinical practice for modulating the

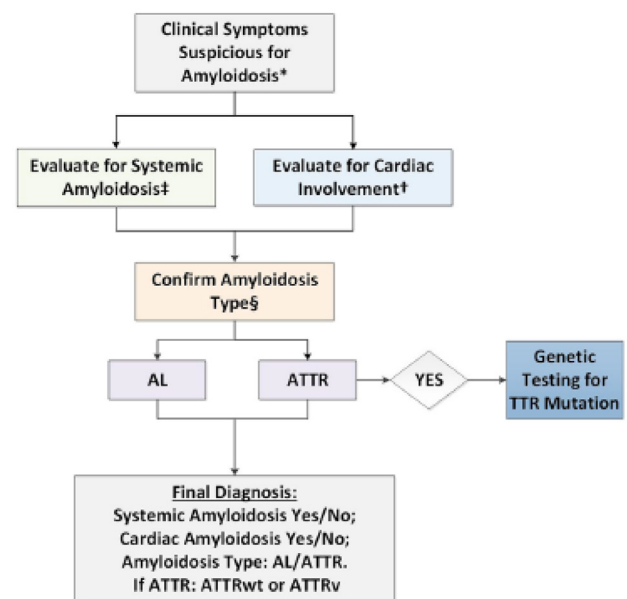


Fig. 1. Systematic evaluation of cardiac amyloidosis. A comprehensive evaluation of cardiac amyloidosis includes consideration of clinical symptoms, evaluation of cardiac involvement (biomarkers and cardiac imaging), evaluation of systemic amyloidosis (serum, urine testing, and biopsy), followed by typing of amyloid deposits into AL or ATTR, and documentation of mutations in patients with ATTR amyloidosis. *Clinical symptoms: heart failure, peripheral/autonomic neuropathy, macroglossia, carpal tunnel syndrome, periorbital bruising, stroke, atrial fibrillation, postural hypotension, fatigue, weight loss, pedal edema, renal dysfunction, diarrhea, constipation. †Evaluation for cardiac amyloidosis: ECG, ECHO, CMR, EMB, ^{99m}Tc-PYP/DPD/HMDP/¹²³I-mIBG/PET, NT-proBNP, troponin T. ‡Evaluation for systemic amyloidosis: AL: detect plasma cell clone: serum and urine immunofixation, serum FLC assay and immunoglobulin analysis; AL: detect systemic organ involvement: 24-hour urine protein, Alkaline phosphatase, eGFR, cardiac biomarkers (NT-proBNP, troponins); tissue biopsy: EMB/Fatpad/bone marrow/other with Congo red staining. §Confirm amyloidosis type: ATTR: IHC and MS of biopsy or ^{99m}Tc-PYP/DPD/HMDP Grade 2 or 3 if a clonal process is excluded; AL: MS or IHC of biopsy. ¶Confirm TTR mutation in patients with ATTR amyloidosis: genetic testing for TTR mutations. ECHO, echocardiogram; eGFR, estimated glomerular filtration rate; IHC, immunohistochemistry; v, hereditary; wt, wild-type.

therapy intensity in AL amyloidosis.²⁶ A European study identified a stage 3b subgroup with very advanced cardiac involvement; these patients had high concentrations of NT-proBNP (>8500 ng/L) and a very poor prognosis, which warrants further study.²⁷ Furthermore, a reduction in FLC following anti-plasma cell treatment, termed a hematologic response, is typically followed within 6 to 12 months by a reduction in NT-proBNP and troponin, termed an organ-specific response, which is associated with improved symptoms of heart failure and extended survival.²⁸ The FLC-based and NT-proBNP-based hematology and cardiac responses have been extensively validated in AL amyloidosis.²⁹ In ATTR cardiac amyloidosis, NT-proBNP, cardiac troponin, and estimated glomerular filtration rate have also been validated as diagnostic markers in different risk-prediction models,^{30–32} with changes in NT-proBNP useful to follow disease progression.^{18,22} Biomarker evaluation is an integral part of the management of patients with AL and ATTR cardiac amyloidosis.

Evolution of Imaging in Cardiac Amyloidosis

Despite the widespread utilization of serum biomarkers for risk assessment of cardiac amyloidosis, biomarkers themselves are nonspecific for the diagnosis of amyloidosis. This lack of specificity is primarily because of confounding by renal function and overlap with other cardiomyopathies that also result in abnormalities of NT-proBNP and troponin. For this reason, imaging remains a requisite component of the diagnostic algorithm for cardiac amyloidosis. In addition, imaging alone captures the cardiac functional impairment caused by amyloid infiltration and affords insight into hemodynamics. Finally, imaging has the potential to directly visualize cardiac remodeling that may result from both FLC reduction, TTR stabilization/suppression, and/or the anti-amyloid-specific therapies in development. This consensus document serves as means to summarize the interpretation and application of multimodal imaging in cardiac amyloidosis.

The first descriptions of echocardiographic findings in cardiac amyloidosis were reported more than 40 years ago.^{1,33} Since that time, echocardiography has become a standard part of the diagnostic assessment in patients with suspected or confirmed cardiac amyloidosis.^{34–37} The initial studies of echocardiography in cardiac amyloidosis occurred when only M-mode echocardiography was routinely available and predated the advent of clinical 2D and Doppler echocardiography. Nevertheless, these early studies recognized many of the findings of cardiac amyloidosis still used today in clinical practice,^{34–41} along with more recent advances as discussed in subsequent sections.^{1,33} Echocardiography has the advantage of portability, bedside availability, conspicuous presence, and superior diastolic function assessment. Thus, although echocardiography is not sufficient by itself to make the diagnosis of cardiac amyloidosis, it is an essential part of the diagnostic evaluation and ongoing management of patients with this disorder.

CMR in cardiac amyloidosis provides structural and functional information that complements echocardiography.⁴²

CMR may have advantages when acoustic windows are poor, for characterization of the right ventricle, tissue characterization based on the contrast-enhanced patterns of myocardial infiltration, and precise quantification of cardiac chamber volumes and ventricular mass. However, CMR with late gadolinium enhancement (LGE) may be relatively contraindicated in patients with suspected cardiac amyloidosis and concomitant renal failure—a frequent occurrence. Moreover, in centers where CMR scanning in patients with pacemakers is not yet routine, echocardiography may be the only option for imaging cardiac structure and function. Although both the echocardiographic and CMR assessment of structure and function alone may be nonspecific, some features provide more specificity, including biventricular long axis function impairment, apical sparing, reduced stroke volume index, pericardial effusion, marked biatrial enlargement, atrial appendage thrombus in sinus rhythm, sparkling texture of the myocardium, and/or disproportionate increase in left ventricular (LV) mass for electrocardiogram (ECG) voltages. Given the limitations of assessment of structure and function alone (by echo or CMR), tissue characterization by CMR adds high value, as discussed in subsequent sections.

Radionuclide imaging provides critical information on amyloid type that complements cardiac structural and functional characterization by echocardiography and CMR. It has long been appreciated that there is a unique myocardial uptake pattern in amyloid by scintigraphy with ^{99m}Tc-Technetium (Tc)-bisphosphonate derivatives (^{99m}Tc-pyrophosphate [PYP], ^{99m}Tc-3,3-diphosphono-1,2-propanodicarboxylic acid [^{99m}Tc-DPD), ^{99m}Tc hydroxymethylene-diphosphonate [^{99m}Tc-HMDP]). Many studies dating from the 1970s and 80s suggested that ^{99m}Tc-PYP could assist in diagnosing amyloidosis.^{2,43–48} However, there was variable diagnostic accuracy, which limited early use of the technique, owing to the study of mixed patients populations with undifferentiated ATTR and AL subtypes. Subsequent studies comparing ^{99m}Tc-bisphosphonate scintigraphy to gold standard endomyocardial biopsy discovered that ATTR cardiac amyloidosis has avidity for bone radiotracers, whereas AL cardiac amyloidosis has minimal or no avidity for these tracers. Therefore, bone-avid radiotracers can definitively diagnose amyloid type when a plasma cell dyscrasia is excluded. Recognition of preferential ATTR binding to bone-avid ^{99m}Tc-bisphosphonate-based radiotracers resulted in renewed interest and greater clinical application of cardiac scintigraphy with ^{99m}Tc-PYP, ^{99m}Tc-DPD, and ^{99m}Tc-HMDP. Although there is no direct comparison between these tracers, the information available suggests that they can be used interchangeably. This is fortunate, given that there is limited access to ^{99m}Tc-DPD and ^{99m}Tc-HMDP in the United States and ^{99m}Tc-PYP in Europe.

Evidence Base for Cardiac Amyloidosis Imaging

Diagnosis

Cardiac amyloidosis is substantially underdiagnosed because of varied clinical manifestations, especially in the early stages of disease. An ideal noninvasive diagnostic

method would identify cardiac involvement in amyloidosis and would also confirm the etiologic subtype. No existing diagnostic tools can provide this information individually, necessitating a multimodality cardiac imaging approach.

Echocardiography. Echocardiography plays a major role in the noninvasive diagnosis of cardiac amyloidosis because of its assessment of structure and function and its pervasive use in patients with concerning cardiac symptoms. The evaluation of cardiac amyloidosis using echocardiography focuses on morphologic findings related to amyloid infiltration, in particular, thickened LV walls >1.2 cm in the absence of any other plausible causes of LV hypertrophy (Fig. 2).²⁸ Although increased LV mass in the setting of low-voltage ECG is suggestive of cardiac amyloidosis, a definitive distinction by echocardiography of amyloidosis from hypertrophic cardiomyopathy or other causes of LV hypertrophy is challenging.⁴⁹ Other echocardiographic findings that suggest infiltrative disease include normal to small LV cavity size; biatrial enlargement and dysfunction⁴¹; left atrial and left atrial appendage stasis and

thrombi; thickened valves; right ventricular and interatrial septal thickening; pericardial effusion; and a restrictive transmitral Doppler filling pattern.^{50–54} Several of these features, including an overt restrictive mitral inflow pattern are uncommon until late in the disease process.^{34,51} However, reduced LV systolic thickening, filling pressures, cardiac output,³⁸ early diastolic dysfunction,^{39,40} and signs of raised filling pressures are commonly seen.^{34,52} A granular sparkling appearance of the myocardial walls may be appreciated, but it is not considered a highly specific finding and can be seen in other conditions, such as end-stage renal disease. The echocardiographic shift from fundamental to harmonic imaging has confounded this phenotype.

Tissue Doppler imaging (TDI) and speckle-tracking echocardiography (STE) refine the noninvasive recognition of cardiac amyloidosis by quantitating longitudinal systolic function.^{51,55,56} A pattern of reduced longitudinal shortening with preserved LV ejection fraction and radial shortening is characteristic of cardiac amyloidosis and can differentiate it from other causes of increased LV wall

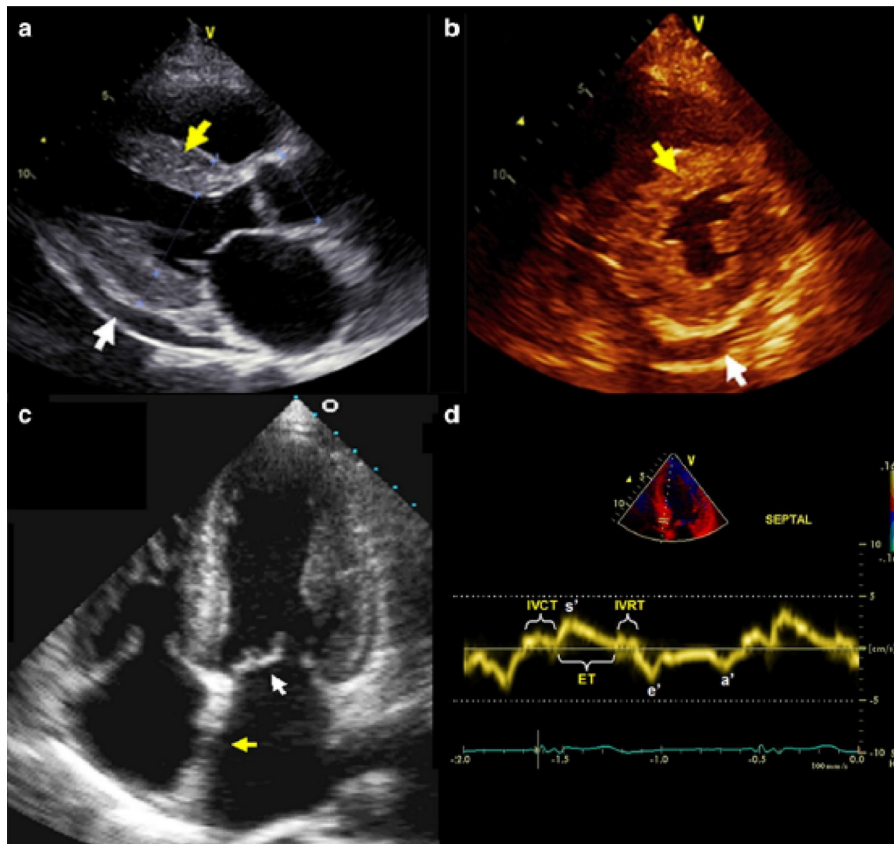


Fig. 2. Characteristic appearance of cardiac amyloidosis on echocardiography. (a–d) 2D echocardiography. (a) (parasternal long axis) and (b) (parasternal short axis) demonstrate increased LV wall thickness with a sparkling texture of the myocardium (yellow arrows) in a patient with primary (AL) cardiac amyloidosis. Also, note the small pericardial effusion (white arrows), which is often seen in patients with cardiac amyloidosis. (c) (apical 4-chamber view) demonstrates increased biventricular wall thickness, biatrial enlargement, and increased thickening of the interatrial septum (yellow arrow) and mitral valve leaflets (white arrow) in a patient with wild-type transthyretin cardiac amyloidosis. (d) TDI tracing taken at the septal mitral annulus in a patient with ATTR cardiac amyloidosis. The TDI tracings show the “5-5-5” sign (s' [systolic], e' [early diastolic], and a' [late (atrial) diastolic] tissue velocities are all <5 cm/s), which is seen in patients with more advanced cardiac amyloidosis. The dotted lines denote the 5 cm/s cutoff for systolic and diastolic tissue velocities. In addition to the decreased tissue velocities, isovolumic contraction and relaxation times (IVCT and IVRT, respectively) are increased and ejection time (ET) is decreased, findings also seen in patients with cardiac amyloidosis especially as the disease becomes more advanced.

thickness. Longitudinal systolic function is commonly impaired, even in the earlier phases of the disease, when radial thickening and circumferential shortening are still preserved.^{34,51,57–62} Both AL and ATTR cardiac amyloidosis patients demonstrate a typical pattern of distribution of STE-derived longitudinal strain in which basal LV segments are severely impaired while apical segments are relatively spared (Fig. 3).^{51,63} Conversely, patients with other causes of LV hypertrophy (ie, aortic stenosis, hypertrophic cardiomyopathy) typically show reduced LV longitudinal strain in the regions of maximal hypertrophy.^{63,64}

Another abnormal quantitative measure of LV contractility in cardiac amyloidosis is the myocardial contraction fraction (MCF), the ratio of stroke volume to myocardial volume. The MCF is an index of the volumetric shortening of the myocardium that is independent of chamber size and geometry and highly correlated with LV longitudinal strain.^{65–67} Abnormalities beyond the left ventricle can also suggest cardiac amyloidosis. Recently, it has been reported that the stroke volume index has a prognostic performance similar to LV strain in predicting survival in AL cardiac amyloidosis, independently of biomarker staging. Because

the stroke volume index is routinely calculated and widely available, it could serve as the preferred echocardiographic measure to predict outcomes in AL cardiac amyloidosis patients. Left atrial reservoir and pump functions measured by strain are frequently impaired, irrespective of left atrial size, suggesting that both raised LV filling pressures and direct atrial amyloid infiltration (as documented by CMR studies) contribute to left atrial dysfunction.^{41,68} This dysfunction may result in the formation of atrial and atrial appendage thrombi, even in the setting of normal sinus rhythm, exposing patients to higher relative risk for embolic strokes. Although data are not available, clinical experience from major amyloidosis centers suggests that the highly thrombogenic milieu of the left atrium increases cardioembolic risk in these patients.⁶⁹ The right ventricle is often affected due to a combination of increased afterload from pulmonary hypertension and intrinsic right ventricular amyloid infiltration, resulting in reduced tricuspid annular plane systolic excursion, tissue Doppler systolic velocity, and longitudinal strain.⁷⁰

Because echocardiographic findings lack the tissue characterization provided by CMR, echocardiographic diagnosis of

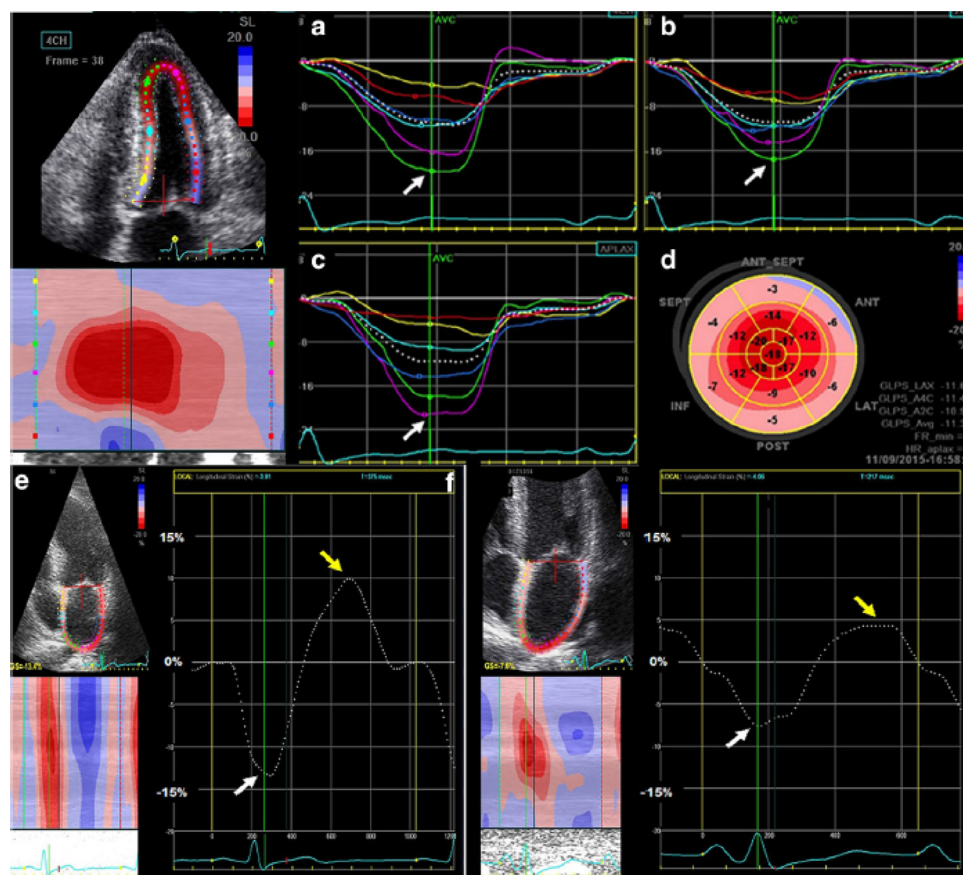


Fig. 3. Left ventricular longitudinal strain abnormalities. (a) (apical 4-chamber view), (b) (apical 2-chamber view), (c) (apical 3-chamber view) all show abnormal longitudinal strain in the basal and mid segments with relative preservation in the apical segments (purple and green curves, white arrows) in a patient with ATTRv cardiac amyloidosis. (d) The corresponding bullseye map of the longitudinal strain pattern throughout the left ventricle with the “cherry-on-the-top” sign (red denotes normal longitudinal strain at the apex and pink/blue denotes abnormal longitudinal strain at the mid/basal left ventricle).

cardiac amyloidosis relies on the presence of highly suggestive findings that can confirm diagnostic suspicion.^{54,71} Table 1 lists the echocardiographic parameters for acquisition, interpretation, and reporting in cardiac amyloidosis. Moreover, abnormal parameters are provided that suggest cardiac amyloidosis and warrant further evaluation. The

combination of these echocardiographic “red flags” with other parameters, such as abnormal cardiac biomarkers and electrocardiographic findings, maximizes diagnostic accuracy.⁷² For instance, the combination of pericardial effusion and symmetric LV wall thickening in the presence of low or normal QRS voltages should prompt a strong suspicion of

Table 1. Standardized Acquisition, Interpretation, and Reporting of Echocardiography for Cardiac Amyloidosis

Parameter for acquisition and reporting	Abnormal parameter	Notes	Recommendations for reporting
2D, Color, and Spectral Doppler Imaging			Required
LV wall thickness	Increased LV wall thickness (>1.2 cm) and increased relative wall thickness (>0.42)	Increased LV wall thickness relative to ECG QRS voltage is particularly suggestive	Required
Myocardial echogenicity	Increased echogenicity of the myocardium (sparkling, hyper-refractile “texture” of the myocardium)	Not highly specific (differential diagnosis includes ESRD or other infiltrative cardiomyopathies). However, this finding in conjunction with severely reduced longitudinal function of the LV is highly suggestive.	Required
Atrial size and function	Atrial enlargement and dysfunction	Non-specific but important finding to support the diagnosis and potentially provide insight into risk for stroke or arterial embolism	Required
Interatrial septum and valves	Thickening of the interatrial septum and valves (>.5 cm)	Non-specific but suggestive of the diagnosis	Required
Pericardial effusion	Pericardial effusion	Non-specific, but when coupled with other echo signs is suggestive of the diagnosis	Required
Diastolic function	Grade 2 or worse diastolic dysfunction with high E/A ratio (>1.5) and reduced E deceleration time (<150 ms)	Doppler diastolic function is helpful in determining prognosis. Severely reduced A wave velocity can be due to LA failure, which can be helpful in determining risk of stroke.	Required
Estimated PA systolic and right atrial pressure	Increased pressures (>35mmHg for PA, ≥10mmHg for RA)	These are important parameters to estimate volume status and optimize diuretic dosing.	Required
Tissue Doppler Imaging			Required
Tissue Doppler velocities	Reduced tissue Doppler’s*, e*, and a* velocities (all <5 cm/s)	If present, the “5-5-5” sign (all TDI velocities <5 cm/s) can be useful and is typically highly suggestive of the diagnosis but may not be sensitive for the diagnosis in early forms of the disease	Required
Strain Imaging			Recommended
Longitudinal LV strain	Decreased global longitudinal LV strain (absolute value less than -15%)	2D and STE shows characteristic appearance of myocardial deformation in patients with cardiac amyloidosis	Recommended
Longitudinal LV strain bullseye map	“Cherry-on-the-top” sign on STE longitudinal strain bullseye map (preservation of apical longitudinal strain with severely abnormal basal and mid-LV longitudinal strain)	Characteristic bullseye pattern is likely the most specific sign to rule in the diagnosis of cardiac amyloidosis (but still does not differentiate ATTR vs. AL amyloidosis)	Recommended

REPORTING	
An overall interpretation of the echo findings into categories of:	Required
Not suggestive: Normal LV wall thickness, normal LV mass normal atrial size, septal or lateral tissue Doppler e* velocity >10 cm/s Strongly suggestive: Increased LV wall thickness, increased LV mass, typical LV longitudinal strain pattern, mitral annular TDI < 5 cm/sec, biatrial enlargement, small A wave in sinus rhythm, small pericardial and/or pleural effusions Equivocal: Findings not described above	
Interpret the echo results in the context of prior evaluation.	Recommended
Provide follow-up recommendations: Strongly suggestive echocardiographic findings cannot distinguish AL from TTR cardiac amyloidosis. Endomyocardial biopsy is not always indicated in patients with strongly suggestive echo findings. Please see Part 2, Table 1 “Expert Consensus Recommendations for Diagnosis of Cardiac Amyloidosis” for indications for endomyocardial biopsy. Consider evaluation (1) to exclude AL amyloidosis, evaluate for plasma cell dyscrasia (serum and urine immunofixation, serum FLC assay) and (2) to exclude ATTR cardiac amyloidosis, consider imaging with ^{99m} Tc-PYP/DPD/HMDP.	Recommended

2D, 2 dimensional; A, late (atrial) mitral inflow velocity; AL, amyloid light chain; ATTR, amyloid transthyretin; E, early mitral inflow velocity; E/A, ratio of early to late (atrial) mitral inflow velocities; ECG, electrocardiogram; ESRD, end-stage renal disease; IVCT, isovolumic contraction time; IVRT, isovolumic relaxation time; LA, left atrial; LV, left ventricular; PA, pulmonary artery; RA, right atrium; STE, speckle-tracking echocardiography; TDI, tissue Doppler imaging

cardiac amyloidosis.^{50,72,73} In particular, the ratio of QRS voltage to echocardiographic LV wall thickness is useful in diagnosing cardiac amyloidosis.⁴⁹

Key Recommendations for Diagnosis: Echocardiography.

- Comprehensive 2D echocardiography, including quantitative tissue Doppler and speckle-tracking strain analysis (when available) should be performed in all patients with unexplained LV wall thickening and a clinical suspicion of cardiac amyloidosis.
- To increase identification of this underdiagnosed disease, any echocardiographic abnormalities suggestive of cardiac amyloidosis should prompt further evaluation.
- Echocardiographic parameters should be combined with electrocardiographic, clinical, biomarker, and other imaging findings to maximize diagnostic accuracy.

Cardiac Magnetic Resonance. CMR has a central role in the noninvasive diagnosis of cardiac amyloidosis because of its ability to provide tissue characterization in addition to high-resolution morphologic and functional assessment. CMR offers value in 2 clinical scenarios: the differentiation of cardiac amyloidosis from other cardiomyopathic processes with increased wall thickening and potentially in detection of early cardiac involvement in patients with evidence of systemic amyloidosis. A comprehensive CMR evaluation for cardiac amyloidosis includes morphologic and functional assessment of the left and right ventricles and atria using cine imaging, evaluation of native T1 signal (assessed on non-contrast T1 mapping), assessment of LGE, and extracellular volume (ECV) measurement. Overall, current published reports from single-center studies demonstrated heterogeneity in study design, sample size, and types of amyloidosis included. See the Appendix for a summary of the published literature on diagnosis of cardiac amyloidosis using CMR.

Maceira et al⁷⁴ described a typical LGE pattern in cardiac amyloidosis of global subendocardial enhancement. Initial observations were that nulling—rendering remote myocardium dark—was difficult in cardiac amyloidosis. The blood pool and myocardium null together due to expansion of the extra cellular myocardial volume (from amyloid infiltration), which approaches plasma volume. An inversion time scout (TI-scout) technique (obtaining a series of images with various inversion time values) could be useful to select the optimal inversion time for the LGE sequence.⁷⁵ Traditional LGE imaging techniques, however, can be difficult to acquire and interpret in cardiac amyloidosis. LGE using the widely available relatively new phase-sensitive inversion recovery sequence (PSIR) eliminates the need to optimize null-point settings, making LGE in cardiac amyloidosis more robust and operator independent. Using the PSIR technique, LGE is significantly more specific and sensitive than echo or CMR functional assessment. Although multiple LGE distributions have been described in cardiac amyloidosis, subendocardial and transmural LGE patterns

predominate. Both patterns are present in AL and ATTR cardiac amyloidosis, but to different extents, with subendocardial LGE being more prevalent in AL and transmural LGE more prevalent in ATTR cardiac amyloidosis.⁷⁶ LGE shows an initial basal predilection but with biventricular transmurality in advanced disease.^{77–80}

At 4 minutes post-gadolinium administration, a subendocardial-blood T1 difference of 191 ms detected cardiac amyloidosis at 90% and 87% sensitivity and specificity, respectively.⁷⁴ In several studies where the results of an endomyocardial biopsy have been used as a reference standard, a typical LGE pattern has consistently been shown to have a diagnostic sensitivity of 85% to 90%.^{74,77,78,80–82} However, the true specificity of LGE in diagnosing cardiac amyloidosis with reference to histologic evidence cannot be accurately determined, given verification bias (typically only positive CMR cases are referred for endomyocardial biopsy). A recent meta-analysis based on 7 published studies, estimated a sensitivity and specificity of 85% and 92%, respectively, for CMR-based LGE in diagnosing cardiac amyloidosis.⁸³

Other CMR methods include native (non-contrast)^{82,84} and post-contrast T1 mapping,⁸⁵ left atrial LGE,⁶⁸ and qualitative visual T1 comparison between the myocardium and cardiac blood pool.⁷⁹ The method of a nulling comparison between the myocardium and the blood pool allows a rapid confirmation of cardiac amyloidosis diagnosis as an adjunct to LGE findings, at an excellent sensitivity but a moderate specificity.⁷⁹ LGE in non-ischemic cardiomyopathies, especially cardiac amyloidosis, is not easy to quantify; therefore, using LGE to track changes over time can be difficult. T1 mapping is a new technique where a direct quantitative signal from the myocardium is measured, either pre-contrast (native T1) or post-contrast (ECV).⁸⁶ T1 mapping before and after contrast administration allows a quantitative measure of the contrast exchange between the blood pool and the expanded extracellular compartment, thus permitting an incremental characterization and detection of the degree of infiltration.

Native T1 may find particular utility when administration of contrast is contraindicated. Of note, a recent report demonstrated that native myocardial T1 measured by the shortened modified look-locker inversion recovery (ShMOLLI) method achieved a diagnostic sensitivity and specificity of 92% and 91%, respectively.⁸² Native T1, however, is a composite signal from the extracellular and intracellular space, and administration of contrast with ECV measurement enables us to isolate the signal from the extracellular space.⁸⁶ Amyloidosis is an exemplar of interstitial disease, and this is reflected by substantial elevation of ECV in patients with AL and ATTR cardiac amyloidosis.^{85,87} ECV is also elevated even when conventional testing and LGE suggest no cardiac involvement, highlighting a potential role of ECV as an early disease marker.⁸⁸ Both native T1 and ECV track a variety of markers of disease activity, and there is early evidence they could be used to track changes in amyloid burden over time.

Advanced techniques, such as T2 mapping and perfusion are being used to assess additional aspects of the cardiac amyloidosis phenotype, including myocardial edema⁸⁹ and coronary microvascular dysfunction. Using a combination of CMR features, a measure of the likelihood of cardiac amyloid type (ATTR vs AL), and likelihood of ATTR versus AL can be gleaned^{90,91}; but, this is typically not sufficient for excluding AL cardiac amyloidosis. FLCs combined with cardiac scintigraphy with bone tracers have advantages over echo and CMR for differentiation of the type of cardiac amyloidosis.³

Key Recommendations for Diagnosis: Cardiac Magnetic Resonance.

- 1) Comprehensive CMR-based evaluation of cardiac structure, function, and myocardial tissue characterization is helpful for diagnosis of cardiac amyloidosis, particularly when echocardiographic findings are suggestive or indeterminate.
- 2) In patients with biopsy-proven systemic amyloidosis, typical CMR findings, including diffuse LGE, nulling of myocardium before or at the same inversion time as the blood pool, and extensive ECV expansion are combined with structural findings of increased wall thickness and myocardial mass to diagnose cardiac involvement. In the absence of documented systemic amyloidosis, typical CMR features should prompt further evaluation for cardiac amyloidosis.
- 3) CMR, however, is typically unable to definitively distinguish AL from ATTR cardiac amyloidosis.
- 4) CMR parameters should be combined with electrocardiographic, clinical, biomarker, and other imaging findings to maximize diagnostic accuracy.

Radionuclide Imaging. Radionuclide imaging plays a unique role in the noninvasive diagnosis of cardiac amyloidosis. A variety of ^{99m}Tc-labeled diphosphonate and PYP (bone-avid) compounds diagnose ATTR cardiac amyloidosis with high sensitivity and specificity.³ Targeted amyloid binding ¹⁸F-positron emission tomography (PET) tracers are highly specific to image amyloid deposits and appear to bind to both AL and ATTR.^{92–96} ¹²³I-meta-iodobenzylguanidine (*mIBG*), an established tracer for imaging myocardial denervation, has been utilized to image myocardial denervation in familial ATTR cardiac amyloidosis.^{97,98} A substantial additional benefit of radionuclide evaluation of cardiac amyloidosis is that whole-body imaging can be performed concurrently, allowing evaluation of multi-organ systemic involvement.

The explanation for this differential uptake in ATTR versus AL cardiac amyloidosis is unknown, but it has been suggested that the preferential uptake by ATTR may be a result of higher calcium content.^{99,100} Furthermore, the type of mutation and the result of the proteolysis of myocardial fibers (full-length only vs full-length plus C-terminal ATTR fragments) also modulate uptake of bone radiotracers by amyloid fibrils.¹⁰⁰

Bone-Avid Radiotracers for Cardiac Scintigraphy: ^{99m}Tc-PYP/DPD/HMDP. Systematic evaluation of diphosphonate radiotracers suggests

that cardiac uptake of ^{99m}Tc-PYP, ^{99m}Tc-DPD, and ^{99m}Tc-HMDP are remarkably sensitive (but not completely specific) for ATTR cardiac amyloidosis.^{3,31,100–106} Notably in the absence of cardiac amyloidosis (or previous myocardial infarction), there is no myocardial uptake of bone tracers; therefore, cardiac scintigraphy with bone-avid radiotracers may reliably distinguish cardiac amyloidosis from other entities that mimic cardiac amyloidosis, such as hypertrophic cardiomyopathy.^{31,102} Cardiac scintigraphy with bone-avid radiotracers is particularly sensitive in the early identification of ATTR cardiac amyloidosis, including carriers without apparent cardiac involvement by other diagnostic techniques.^{105,107,108} Furthermore, ^{99m}Tc-DPD/HMDP allows the possibility of detecting extra-cardiac (skeletal muscle and lung) amyloid infiltration.^{109,110} See the Appendix for a full summary of the published literature on diagnosis of ATTR cardiac amyloidosis using ^{99m}Tc-PYP/DPD/HMDP.

A multicenter experience in 1498 patients showed a positive predictive value for ATTR cardiac amyloidosis of 100% (95% confidence interval, 98.0–100) in patients with an echocardiogram or CMR consistent with or suggestive of cardiac amyloidosis, and absence of monoclonal protein using urine and serum, with serum FLC assay and immunofixation electrophoresis.³ A recent bivariate meta-analysis confirmed the accuracy of bone scintigraphy in the assessment of ATTR cardiac amyloidosis.¹¹¹ Again, these high sensitivities and specificities were reported from major centers of expertise and in patients with advanced stages of the disease, and often with New York Heart Association (NYHA) heart failure greater than Class II. The yield of ^{99m}Tc-PYP/DPD/HMDP cardiac scintigraphy in patients with earlier stages of disease or with preclinical disease is yet to be confirmed.

Several diagnostic parameters have been evaluated on cardiac scintigraphy with bone-avid tracers. The ratio of heart-to-contralateral (H/CL) lung uptake (semiquantitative scoring), heart-to-whole-body (H/WB) retention, and a heart-to-bone ratio (visual grade) have been assessed at both 1 and 3 hours (see Standardized Imaging Techniques). Early work by Perugini et al¹¹² found that a visual grade ≥ 2 on ^{99m}Tc-DPD (ie, moderate or strong myocardial uptake) was 100% sensitive to identify ATTR cardiac amyloidosis and 100% specific to distinguish from AL and control subjects. Subsequent studies have confirmed the high sensitivity to detect ATTR cardiac amyloidosis and showed that mild uptake of ^{99m}Tc-DPD (Grade 1) may be noted in patients with other subtypes of cardiac amyloidosis (ie, AL, amyloid A amyloidosis, and apolipoprotein A1).^{43,106} Rapezzi et al¹⁰⁵ evaluated the ratio of H/WB retention of ^{99m}Tc-DPD, on the late (3-hour) images, in patients with TTR mutation, and demonstrated that individuals with increased LV myocardial wall thickness >1.2 cm had much higher H/WB retention ratio compared to individuals with normal LV wall thickness. In a single-center experience, Bokhari et al¹¹³ identified a very high diagnostic accuracy (area under the curve of 0.992, $P < .0001$) for visual Grade ≥ 2 and a H/CL ratio ≥ 1.5 on 1-hour images to distinguish ATTR from AL cardiac amyloidosis.^{3,111} A H/CL ratio ≥ 1.3 has

been proposed to distinguish ATTR accurately from AL cardiac amyloidosis on the late (3-hour) ^{99m}Tc -PYP images.¹¹⁴

The recently developed consensus algorithm for noninvasive diagnosis of cardiac amyloidosis attributes a central role to ^{99m}Tc -PYP/DPD/HMDP cardiac scintigraphy (Fig. 4).³ If cardiac amyloidosis is suspected clinically or based on echocardiography/CMR, blood and urine should be analyzed for evidence of a monoclonal protein and ^{99m}Tc -PYP/DPD/HMDP cardiac scintigraphy should be considered if ATTR cardiac amyloidosis is suspected. If both tests are negative, then current evidence suggests that cardiac amyloidosis is very unlikely. It is still possible, however, for patients with ATTRv to have negative findings on DPD scintigraphy¹⁰⁵ in case of some rare non-V30M mutations and in some V30M mutations with early onset and only full-length TTR fibrils.¹¹⁵ In the presence of a Grade 2 or 3 positive ^{99m}Tc -PYP/DPD/HMDP cardiac scan (see Standardized Imaging Techniques) without evidence for monoclonal proteins in blood and urine, a diagnosis of ATTR cardiac amyloidosis can be made without a biopsy (specificity and positive predictive value >98%).³ For those patients with evidence of a plasma cell dyscrasia, a histologic diagnosis is still required because the presence of low-grade uptake on a ^{99m}Tc -PYP/DPD/HMDP scan is not 100% specific for ATTR cardiac amyloidosis, and substantial uptake (Grade 2 or 3) has been reported in more than 20% of patients with AL cardiac amyloidosis.³ The writing

group would like to emphasize the importance of excluding monoclonal process with serum/urine immunofixation and a serum FLC assay in all patients with suspected amyloidosis.

^{99m}Tc -PYP/DPD/HMDP scintigraphy has been recently used to detect ATTR cardiac amyloidosis in previously unexplored clinical settings, including heart failure with preserved ejection fraction (prevalence 15%)^{13,15} and severe degenerative aortic stenosis,^{14,116} including the “paradoxical low-flow low-gradient” subtype (18%).¹¹⁷

Based on the utility of cardiac scans with SPECT bone-avid radiotracers, there has been interest in ^{18}F -NaF, a PET bone radiotracer, for imaging cardiac amyloidosis.^{118,119} Early reports, however, suggest limited utility for imaging ATTR cardiac amyloidosis, and further studies are warranted to examine its utility.

Amyloid Binding Radiotracers. Several amyloid binding SPECT and PET radiotracers are available for amyloidosis imaging. ^{99m}Tc -aprotinin^{120–122} and ^{123}I -serum amyloid P-component (^{123}I -SAP)¹²³ were originally developed to image systemic amyloidosis but have limited availability. They have not been useful to image cardiac amyloidosis due to poor signal-to-noise ratio¹²³ and concerns for risk of bovine encephalopathy.¹²¹ In contrast, several PET amyloid-binding radiotracers, structurally similar to thioflavin-T and likely binding to the amyloid fibril structure, approved for

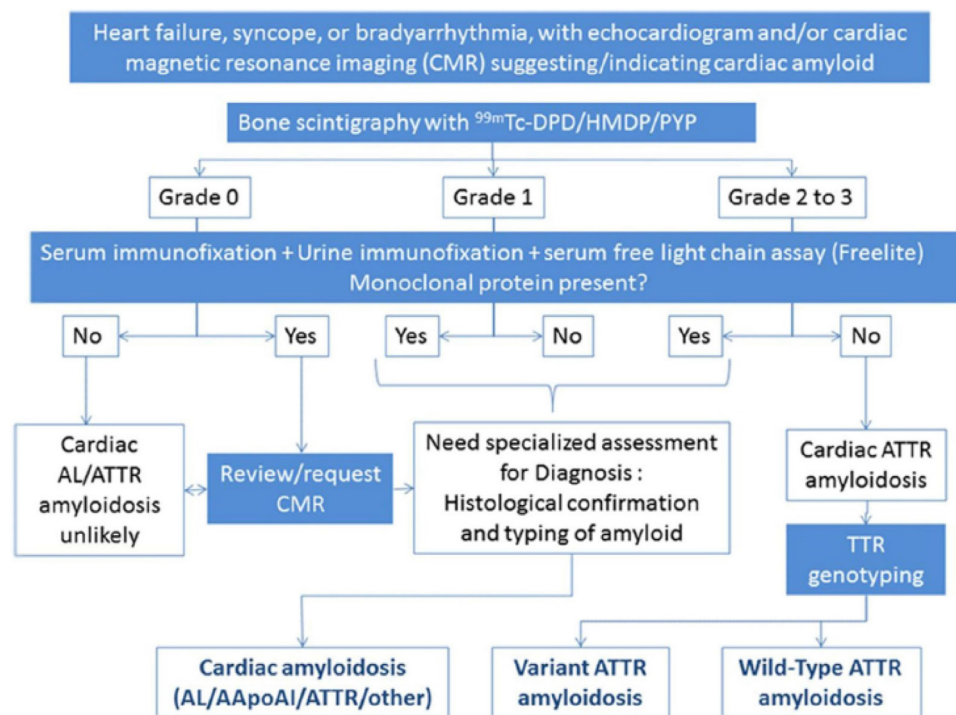


Fig. 4. Consensus algorithm for noninvasive diagnosis of cardiac amyloidosis. This algorithm provides an approach to the evaluation of patients with cardiac amyloidosis. Among patients with suspected cardiac amyloidosis, Grade 2 or 3 uptake of ^{99m}Tc -PYP/DPD/HMDP uptake in the absence of a clonal abnormality is highly specific to diagnose ATTR cardiac amyloidosis avoiding the need for endomyocardial biopsy. Patients with any abnormal serum/urine immunofixation or a positive serum FLC assay should be referred for further evaluation to a hematologist ideally with amyloidosis experience. (Figure reproduced with permission from Gillmore, et al. *Circulation* 2016;133:2404–2412.)

imaging beta amyloid in Alzheimer's disease,¹²⁴ have been successfully used to image cardiac amyloidosis. ¹¹C-Pittsburgh compound B (PIB) was one of the first PET radiotracers developed for beta-amyloid imaging but is limited in availability to sites with a cyclotron. ¹⁸F-florbetapir, ¹⁸F-florbetaben, and ¹⁸F-flutemetamol developed subsequently and are currently FDA approved for beta-amyloid imaging and are widely commercially available. Several additional tracers are still under development.¹²⁵

¹¹C-PIB,^{92,95} ¹⁸F-florbetapir,^{93,96} and ¹⁸F-florbetaben⁹⁴ have been evaluated in patients with AL and ATTR cardiac amyloidosis. In these pilot studies, high cardiac radiotracer uptake was consistently reported in patients with cardiac amyloidosis compared to controls, including hypertensive controls. A target to background (LV myocardium to blood pool) ratio >1.5 and a retention index of >0.025 min⁻¹ with ¹⁸F-florbetapir⁹³ and ¹⁸F-florbetaben⁹⁴ have been shown to separate patients with cardiac amyloidosis from controls without amyloidosis. Myocardial retention of ¹¹C-PIB,⁹² ¹⁸F-florbetapir,⁹³ and ¹⁸F-florbetaben⁹⁴ was significantly greater in cardiac amyloidosis patients compared with controls. In one study, although ¹⁸F-florbetapir myocardial retention index was lower in ATTR compared to AL cases, definitive subtype differentiation was not feasible⁹³; similar findings were confirmed with ¹⁸F-florbetaben.⁹⁵ Although not studied serially, retention of ¹¹C-PIB was lower in AL cardiac amyloidosis patients treated with chemotherapy compared to those who did not undergo treatment,⁹⁵ suggesting that it is possible this radiotracer will be useful for disease monitoring. Finally, unlike echocardiography or CMR, amyloid-binding PET tracers can image systemic amyloid deposits in various other organs^{126,127} and offer the potential to quantify the load of amyloidosis in the whole body.

As literature on PET amyloid-binding radiotracers is limited, sections on risk assessment and standardized protocols are not provided for these radiotracers.

Autonomic Myocardial Innervation Imaging. Patients with amyloidosis are prone to autonomic dysfunction from amyloid infiltration of myocardial and nerve conduction tissue, resulting in rhythm disorders.¹²⁸ Autonomic dysfunction is most common in ATTR cardiac amyloidosis, particularly ATTRv, where it has been studied extensively.^{129,130} Notably, cardiac dysautonomia may occur independent of the presence of a typical restrictive cardiomyopathy.¹³¹ In patients with ATTRwt cardiac amyloidosis, polyneuropathy and dysautonomia are less common, seen in ~9%.¹³² Although AL cardiac amyloidosis patients less commonly manifest autonomic dysfunction,¹³³ it may develop as a complication of AL amyloidosis treatment.¹³⁴ Therefore, autonomic denervation is a nonspecific finding.¹²³ ¹²³I-mIBG scintigraphy is not able to discriminate between cardiac amyloidosis subtypes nor differentiate cardiac amyloidosis from other forms of cardiomyopathy.¹³⁵ However, cardiac denervation evidenced by mIBG occurs earlier than amyloid deposit detection by diphosphonate scintigraphy in TTR mutation carriers.¹³⁶ Although secondary (amyloid A, AA)

amyloidosis rarely shows cardiac manifestations, myocardial denervation has been reported in one study.¹³⁵

Although amyloid infiltration of the cardiac autonomic system cannot be directly imaged, multiple tracers assess autonomic myocardial denervation, including ¹²³I-mIBG, ¹²⁴I-mIBG, N-[3-Bromo-4-3-[¹⁸F]-fluoro-propoxy)-benzyl]-guanine LM1195, and ¹¹C-hydroxy-ephedrine. ¹²³I-mIBG, a chemically modified analogue of norepinephrine, is stored in vesicles in presynaptic sympathetic nerve terminals and is not further catabolized.¹²³ ¹²³I-mIBG has been specifically studied in cardiac amyloidosis, and semiquantitative analysis of ¹²³I-mIBG cardiac uptake compared to background (heart-to-mediastinal ratio [HMR]), provides indirect information of amyloid infiltration in the sympathetic nerve system.^{97,98,130,135,137–141} Decreased HMR at 4 hours after tracer administration (late HMR) reflects the degree of sympathetic dystonia, and is an independent prognostic factor in the development of ventricular dysrhythmia. PET imaging of sympathetic innervation in cardiac amyloidosis has not yet been studied. See the Appendix for a summary of the published literature on assessment of autonomic myocardial innervation imaging in amyloidosis using ¹²³I-mIBG.

Myocardial Perfusion Imaging. Angina, in the absence of coronary artery disease, is common in patients with cardiac amyloidosis. Endothelial¹⁴² and microvascular dysfunction¹⁴³ have been described and may precede the clinical diagnosis of cardiac amyloidosis.^{143,144} In one study, focal and global sub-endocardial hypoperfusion at rest and post-vasodilator stress were ubiquitous in patients with AL and ATTR cardiac amyloidosis.¹⁴⁵ Absolute myocardial blood flow¹⁴⁵ and coronary flow reserve^{144,145} are substantially reduced in patients with cardiac amyloidosis, despite absence of epicardial coronary artery disease. Whether coronary microvascular dysfunction improves after successful anti-amyloid therapy is not known.

Key Recommendations for Diagnosis: Radionuclide Imaging.

- Myocardial imaging with ^{99m}Tc-PYP/DPD/HMDP, in the appropriate clinical context, is highly sensitive and specific to diagnose ATTR cardiac amyloidosis and may aid in its early detection.
- In the absence of a light-chain clone, myocardial uptake of ^{99m}Tc-PYP/DPD/HMDP of Grade ≥2 is diagnostic of ATTR cardiac amyloidosis, obviating the need for endomyocardial biopsy.
- To facilitate early diagnosis of ATTR cardiac amyloidosis, cardiac ^{99m}Tc-PYP/DPD/HMDP scintigraphy should be more broadly considered in patients with unexplained increased LV wall thickness, heart failure with preserved ejection fraction, familial amyloid polyneuropathy, family history of amyloidosis, degenerative aortic stenosis with low-flow low gradient in the elderly, and a history of bilateral carpal tunnel syndrome.
- ¹²³I-mIBG can detect cardiac denervation in patients with hereditary ATTR amyloidosis.

Assessment of Prognosis

Cardiac involvement is common in systemic AL and ATTR amyloidosis and markedly impacts quality of life and outcome.¹⁴⁶ Thus, cardiac assessment in patients with systemic amyloidosis is crucial for risk stratification and treatment decisions. Imaging plays a key role in risk stratification of patients with AL and ATTR cardiac amyloidosis and may add to the existing clinical and biomarker-based risk stratification as discussed previously (see Biomarkers and Biopsy in Cardiac Amyloidosis).

Echocardiography. Abnormalities in several echocardiographic imaging parameters (eg, LV longitudinal strain, early mitral inflow [E-wave], deceleration time, myocardial performance index, pericardial effusion) are associated with worse outcomes and should alert the clinician to the potential of advanced disease.^{9,50,59,65,70,147–165} At the present time, however, there is no formal staging system for ATTRv, ATTRwt, or AL cardiac amyloidosis that uses echocardiographic parameters. Therefore, echocardiography findings in isolation should not be used to determine risk in the individual patient with cardiac amyloidosis. Additional studies to assess the optimal risk-stratification algorithm that incorporates multiple echocardiographic parameters are needed. Moreover, further study is needed to demonstrate the incremental value of echocardiographic parameters over simple clinical markers (eg, NYHA functional class, B-type natriuretic peptide, troponin, glomerular filtration rate) and radionuclide and CMR imaging findings. See the Appendix for a summary of the published literature on the prognostic value of echocardiography in cardiac amyloidosis.

Cardiac Magnetic Resonance. Multiple CMR measures have prognostic significance in cardiac amyloidosis, including LGE presence and pattern, native T1, post-contrast T1, and multiple morphologic parameters.¹⁶⁶ Despite the excellent discriminative capacity of LGE, conflicting results were initially reported describing its prognostic impact in cardiac amyloidosis.^{78,80,81,167} At that time, LGE patterns of cardiac amyloidosis were heterogeneous due to non-standardized acquisition and analysis. The transition to more robust LGE approaches, such as PSIR,¹⁶⁸ has markedly improved image quality. This tool has provided insight into progression of both AL and ATTR cardiac amyloidosis through visualization of a continuum of amyloid accumulation as determined by progression of the LGE pattern from normal to subendocardial to transmural.^{76,169} As a result, several studies now show that the LGE pattern can serve as an independent predictor of prognosis after adjustments for echocardiographic characteristics and blood biomarkers (NT-proBNP and troponin) have been performed.¹⁷⁰ Importantly, the LGE pattern confers prognosis in both AL and ATTR cardiac amyloidosis. Despite its prognostic usefulness, LGE does not lend itself readily toward quantification of myocardial infiltration, owing to different patterns and signal intensities. Thus, the capacity of LGE to track changes accurately over time and monitor response to

treatment is unknown. Parametric T1 mapping has the potential to overcome these limitations.⁸⁶ Recent studies have shown that higher native myocardial T1 can accurately stratify worse prognosis in AL cardiac amyloidosis¹⁷¹ but not in ATTR cardiac amyloidosis.⁸⁷ Alternatively, T1-derived ECV has been associated with prognosis in AL and ATTR cardiac amyloidosis after adjustment for known independent predictors.^{171,172} T2 mapping, a measure of myocardial edema, adds a third dimension to the tissue characterization; in patients with AL cardiac amyloidosis, it is an independent predictor of prognosis.⁸⁹ See the Appendix for a summary of the published literature on the prognostic value of CMR in cardiac amyloidosis.

Radionuclide Imaging. The prognostic role of ^{99m}Tc-PYP/DPD/HMDP scintigraphy and ¹²³I-mIBG have been explored in several studies. ^{99m}Tc-PYP/DPD/HMDP cardiac uptake moderately correlates positively with LV wall thickness and mass, troponin T, NT-proBNP, and ECV; it correlates negatively with LV ejection fraction.^{31,102,105,173–175} The degree of cardiac uptake correlates with overall mortality and survival free from major adverse cardiac events. Multiple semiquantitative markers of cardiac uptake have been studied, including heart and H/WB retention,^{105,174} heart/skull ratio, 102 H/CL ratio,¹⁷³ and visual scoring.¹⁷⁵ In a multicenter study using ^{99m}Tc-PYP, an H/CL ratio of >1.5 was associated with worse survival among patients with ATTR cardiac amyloidosis.¹⁷³ Similar data was found in a single-center study in patients with suspected ATTR cardiac amyloidosis,¹⁷⁵ and these same authors found that regional variability of ^{99m}Tc-PYP uptake may also predict mortality.¹⁷⁶ In all these studies, combining the degree of cardiac uptake with an anatomical (interventricular septal thickness) or functional (Class NYHA, NT-proBNP) variable improved prognostic risk stratification. Of note, visual grading of ^{99m}Tc-PYP/DPD/HMDP has not been shown to be an independent predictor of outcomes.^{31,114}

Cardiac sympathetic denervation is associated with decreased survival in ATTRv cardiac amyloidosis.^{129,131} A late decreased HMR < 1.6 portends a poor prognosis and can be used to identify ATTRv cardiac amyloidosis patients who would benefit from liver transplantation.¹³¹ After liver transplantation, cardiac sympathetic denervation does not appear to progress¹³⁸ and has questionable independent prognostic significance.¹³¹ The prognostic relevance of late-HMR reduction is less clear in AL and ATTRwt cardiac amyloidosis.^{135,138–140} See the Appendix for a summary of the published literature on the prognostic value of radionuclide imaging in ATTR cardiac amyloidosis.

Key Recommendations for Assessment of Prognosis.

- Multiple imaging parameters predict a worse prognosis, including increased LV mass, lower global longitudinal strain, increased right ventricular wall thickness, higher native T1 and ECV, higher H/CL ratio, and ¹²³I-mIBG increased HMR and delayed washout rate.
- Although not formally incorporated into current risk-assessment algorithms, radionuclide results should be

combined with electrocardiographic, clinical, biomarker, and other imaging findings for optimal prognostication.

Management. The ideal method for evaluating the time course of the disease and the response to treatment, particularly disease-modifying treatments, should provide a precise quantitative measure of systemic and cardiac amyloid burden. In AL cardiac amyloidosis, cardiac response is assessed using the serum NT-proBNP concentration, a substantial reduction of which consistently predicts clinical improvement and extended survival.¹⁷⁷ However, the advent of anti-amyloid therapies demands the development of imaging techniques that can estimate the amyloid burden in the heart.

Echocardiography remains the cornerstone of serial assessment of LV dysfunction in patients with heart failure. However, there is relatively little information in the echocardiographic cardiac amyloidosis literature regarding assessment of disease progression and response to therapy. A few studies have shown potential benefit for the use of echocardiography in the following areas: 1) to demonstrate changes in cardiac disease in response to treatment in patients with AL cardiac amyloidosis;^{29,178,179} 2) to determine whether patients with cardiac amyloidosis need to be anticoagulated for stroke prophylaxis; 3) to diagnose progressive cardiac involvement after liver transplantation in patients with ATTRv amyloidosis^{180–182}; and 4) to assess LV ejection fraction in patients with AL amyloidosis being considered for stem-cell transplantation.¹⁸³ Because of a higher incidence of cardiac thrombi in patients with cardiac amyloidosis, some centers consider a transesophageal echocardiogram prior to cardioversion of atrial arrhythmias, even in patients on therapeutic anticoagulation. Emerging data suggests that echocardiographic LV global longitudinal strain may be a marker of disease progression and response to therapy.¹⁸⁴ In contrast, T1 mapping with ECV measurement by CMR can track multiple parameters of structural change (amyloid burden and cardiomyocyte response). In a small retrospective study, the prevalence of a decrease in LV mass and ECV on CMR was higher in patients with AL cardiac amyloidosis and a complete response or very good partial response to chemotherapy.⁸⁸ The quantitative nature of CMR makes it a promising tool to monitor disease progression and response to therapy. Although ^{99m}Tc-PYP/DPD/HMDP scintigraphy correlates well with anatomic and functional variables, this technique has not been definitively proven to quantify changes in response to current therapies, and thus repeat studies are not typically clinically useful.¹⁸⁵ PET is inherently more sensitive and quantitative, and holds the possibility of monitoring response to therapy with PET amyloid-binding tracers once adequately studied. Serial myocardial denervation studies have been studied in ATTRv amyloidosis to guide timing of liver transplantation.¹⁸⁶ Experience with implantable cardioverter defibrillators (ICDs) in cardiac amyloidosis is limited,^{187,188} and the indication for ICD implantation in these patients is unclear even in the setting of myocardial denervation. Prospective

studies are needed in this area. The role of imaging to guide referral to cardiac transplantation and monitor for recurrence post-transplant is not well elucidated and needs further study.

Notably, none of the imaging techniques has been validated for assessing response to therapy, and no study has correlated changes in imaging findings after therapy with survival.

Key Recommendations for Management.

- Transthoracic echocardiography is reasonable to monitor disease progression and/or response to therapy in cardiac amyloidosis because echocardiography is often done clinically for other reasons (ie, heart failure management).
- Transthoracic echocardiography (for the evaluation of left atrial size and function) and transesophageal echocardiography (for the evaluation of the left atrial appendage) are useful to guide initiation and management of anticoagulation in patients with cardiac amyloidosis.
- Cardiovascular magnetic resonance assessment of LV wall thickness, LV mass, and particularly ECV is emerging as a tool to assess disease progression and response to therapy.
- Serial SPECT ^{99m}Tc-PYP/DPD/HMDP scintigraphy is currently not recommended to assess disease progression or response to therapy.

Standardized Imaging Techniques

Extensive research has been performed in cardiac amyloidosis using varied protocols without a clear consensus. This section will provide recommendations for standardized image acquisition, interpretation, and reporting in the assessment of cardiac amyloidosis using echocardiography, CMR, and radionuclide imaging. Standardization would facilitate comparability and reproducibility within and across institutions and enable pooling of data for research purposes.

Echocardiography

2D Echocardiography. 2D and Doppler echocardiographic acquisition in patients with suspected or known cardiac amyloidosis should follow the ASE/European Association of Cardiovascular Imaging (EACVI) guidelines,^{40,189} and all standard transthoracic echocardiography views should be obtained. Required and optional reporting recommendations are provided in Table 1. When reporting results of the echocardiogram in this population, it is important to distinguish other forms of LV hypertrophy from increased LV wall thickness due to amyloid infiltration. Therefore, the report should include not only wall-thickness measurements but also qualitative assessment of the “texture” of the myocardium. Other morphologic features that can be helpful for the diagnosis of cardiac amyloidosis

(eg, atrial enlargement; increased relative wall thickness defined as 2 times posterior wall thickness/LV end-diastolic dimension; thickening of the interatrial septum and/or valves; and the presence of a pericardial effusion) should also be reported. The visual assessment of the loss of longitudinal motion of the heart on 2D imaging (ie, minimal descent of the base in the apical views) can be helpful to include in the report as it increases the likelihood of cardiac amyloidosis.

In patients with cardiac amyloidosis, right ventricular involvement confers a worse prognosis; thus, right ventricular wall thickness (measured in the subcostal view at end-diastole) and assessment of right ventricular systolic function should be included in the report.¹⁸⁹ On Doppler assessment, evaluation of diastolic function (mitral inflow velocities, early mitral inflow [E] deceleration time, and early diastolic relaxation velocity on TDI; see next section below) should be reported.^{39,40} In addition, estimation of hemodynamics (including right atrial pressure, pulmonary artery systolic pressure, LV filling pressure [based on E/e' ratio], and cardiac output [based on LV outflow tract diameter, velocity-time integral (on pulse wave Doppler)]) is helpful for the management of heart failure.³⁸

Tissue Doppler Echocardiography. Accurate tissue Doppler images should be obtained per ASE and EACVI recommendations.⁴⁰ As shown in Fig. 2, in the setting of cardiac amyloidosis, s', e', and a' velocities are all often reduced, and should be reported. The right ventricular free wall TDI should be measured, and the s' velocity reported as a measure of right ventricular longitudinal systolic function (<10 cm/s is abnormal).¹⁹⁰ In addition, isovolumic relaxation and contraction times are increased, and ejection time is decreased. Although not widely used in clinical practice, these 3 indices can be combined to calculate the myocardial performance index (ejection time/[isovolumic relaxation time + isovolumic contraction time]), which is also reduced in the majority of patients with overt cardiac amyloidosis.¹⁶⁵

Speckle-Tracking (Strain) Echocardiography. High-quality longitudinal strain STE LV curves should be obtained in the apical 2-, 3-, and 4-chamber views at frame rates of 50–80 fps with good endocardial border definition (Fig. 3). Right ventricular free-wall strain is calculated as the average of the basal, mid, and apical longitudinal segmental strains. The curves for the left atrium should be generated using P–P gating, if the patient is in normal sinus rhythm. In patients with atrial fibrillation or other rhythm with a lack of P waves, there will be no booster component to the left atrial strain curve, and the left atrial conduit and reservoir strains will be equal to each other.^{191,192} Emerging literature (scientific abstract not yet published) suggests that transesophageal echocardiography should be considered in patients with suspected cardiac amyloidosis and distal embolization to rule out left atrial and left atrial appendage thrombi even in the setting of normal sinus rhythm.

Ideally, in all patients with suspected or known cardiac amyloidosis, the global LV longitudinal strain value (which

is calculated using the peak negative instantaneous average of the 18 longitudinal segmental strains [6 in each of the apical views]) should be reported.^{193,194} In addition, a description and assessment of the pattern displayed on the global longitudinal strain bullseye map (as shown in Fig. 3) should be included in the report.⁶³ Right ventricular free wall strain can also be reported. If left atrial strain is performed, the values of the reservoir, conduit, and booster strains can be reported.

Key Recommendations for Standardized Imaging Techniques: Echocardiography.

- Echocardiograms in patients with suspected or known cardiac amyloidosis should be obtained using ASE/EACVI guidelines on comprehensive echocardiography.
- Reporting should include assessment of wall thickness and myocardial “texture”; thickening of other cardiac structures; pericardial effusion; tissue Doppler velocities (s', e', and a'); diastolic function; and hemodynamics.
- STE should be performed routinely in patients with suspected or known cardiac amyloidosis when available, and efforts should be made to optimize the apical 2D imaging views for speckle-tracking analysis. The global longitudinal strain and pattern of segmental strains (ie, “bullseye” map) should be reported. RV and LA strain can be reported when performed.
- An overall reporting on likelihood of amyloidosis based on imaging findings is recommended (not suggestive, strongly suggestive, or equivocal for cardiac amyloidosis).

Cardiac Magnetic Resonance

Structure and Function. CMR assessment of structure and function in patients with suspected or known cardiac amyloidosis follows well-standardized protocols (Table 2).¹⁹⁵ Image interpretation and reporting should highlight effusions, atrial thrombi, long axis function, and stroke volumes in addition to LV and right ventricular ejection fraction. These and other parameters are specified in Table 3.

Late Gadolinium Enhancement. Protocols for LGE assessment in cardiac amyloidosis are likewise well defined.¹⁹⁵ LGE visualizes the extracellular space expansion that occurs in cardiac amyloidosis. LGE imaging depends on “nulling” of normal myocardium in order to detect LGE from slowed gadolinium washout (thus signal enhancement) in abnormal tissue. Initial LGE evaluation of cardiac amyloidosis was challenging due to similar nulling of both the myocardium and blood pool. The more recent PSIR technique, which ensures appropriate nulling, overcomes this limitation.^{76,168} There are 2 phenomena that are unique to the LGE assessment of cardiac amyloidosis. First, there is rapid movement of gadolinium into the ECV due to the high burden of amyloid protein. This results in myocardial nulling prior to or concurrent with the blood pool,

Table 2. Recommendations for Standardized Acquisition of CMR in Cardiac Amyloidosis

#	Protocol Step	Sequence Technique	Note
1	Cine function	Retrospectively gated cine	2-, 4- and 3-chamber and short-axis stack cines per SCMR guidelines
2	Native T1 mapping (pre-contrast)	Quality controlled T1 mapping sequence	Mid and basal short-axis and apical 4-chamber views as per SCMR clinical recommendations
3	T2		Minimum mid-short axis, consider multiple views
4	Contrast type	Gadolinium-based non-protein bound cyclic contrast agent (0.1–0.2 mmol/kg)	
5	T1 mapping post-contrast (ECV estimation)	Quality controlled T1 mapping sequence	Mid and basal short-axis and apical 4-chamber Should be acquired at least 10-minutes post-contrast Sampling scheme can be varied post-contrast to optimize for short T1 times post-contrast
6	TI scout	TI scout	
7	LGE	Phase-sensitive inversion recovery (PSIR) LGE imaging is recommended	2-, 4-, and 3-chamber and short-axis stack per SCMR

The overall imaging protocol as described above will take approximately 45–60 minutes. This table provides a general guide to the steps of a CMR imaging protocol. Some variation between sites may exist. Each of these sequences assesses a unique myocardial characteristic as discussed in the text and Table 3
ECV, extracellular volume; SCMR, Society for Cardiovascular Magnetic Resonance

which can be identified visually on the TI scout.⁷⁹ Second, there is a global delayed washout of gadolinium from the ECV, resulting in diffuse LGE at time points at which LGE are typically assessed in scar imaging.⁷⁴

A limitation of LGE assessment in cardiac amyloidosis is the requirement for gadolinium administration in the setting of a high coincidence of renal failure in ATTR and AL amyloidosis because of age and multiple myeloma and renal involvement, respectively. Cyclic gadolinium agents need to be administered to decrease risk of nephrogenic systemic fibrosis and other complications. Partially protein-bound contrast agents (gadolinium-BOPTA MultiHance, Bracco Diagnostics) should not be used, as neither the ECV technique nor the characteristic amyloid LGE pattern are reliable.¹⁶⁹

T1 and T2 Mapping. In contrast to LGE, T1 and T2 mapping techniques are quantitative tools. Their acquisition has been standardized in a recent consensus statement.^{86,196} Per this guideline, T1 map acquisition is recommended in 2 short-axis slices and a 4-chamber view before and after contrast; T2 map acquisition is recommended in 1 mid-short-axis slice. Use of local reference ranges and quality control phantoms has been emphasized. A potential concern is the time required for these multiple acquisitions.

T1 mapping can measure the longitudinal magnetization of the myocardium before contrast (native T1). In addition, by measuring T1 before and after contrast and correcting for the blood volume of distribution (1-hematocrit), ECV can be derived (Fig. 5). In combination with pre-contrast T1, an approach using one post-contrast T1 has been validated in cardiac amyloidosis¹⁹⁷ and is used by many centers. Other centers perform serial post-contrast measurements, as the fidelity of mapping the myocardial versus blood exchange of contrast may be improved.¹⁹⁸ T1 mapping has advanced from a cumbersome multi-breath-hold technique with contrast infusion; current techniques require a single breath-hold and generate an ECV map automatically, in some cases without the need for hematocrit sampling or off-line processing.^{199,200}

More recently, CMR with multiparametric mapping has been driving a change in disease understanding: cardiac amyloidosis is not a disease of solely infiltration. T2 mapping, a marker of myocardial edema, has been highlighting other processes in the myocardium—a possible new aspect of the evolution of the myocardial phenotype in cardiac amyloidosis.⁸⁹

Other techniques may also add value: perfusion is profoundly abnormal in cardiac amyloidosis with vasodilator stress revealing marked endo to epicardial gradients (Fig. 5).²⁰¹

Table 3. Recommendations for Standardized Interpretation and Reporting of CMR for Cardiac Amyloidosis

Parameter for acquisition and reporting	Criteria	Notes	Recommendations for reporting
LV function and morphology			
LV function	Biventricular long-axis impairment with relative apical functional sparing	Although LV ejection fraction is typically preserved in cardiac amyloidosis, a reduced LV ejection fraction may be seen in advanced cases	Required
LV wall thickness	Increased LV wall thickness: > laboratory ULN for sex on SSFP cine CMR ²⁰⁵ and increased relative wall thickness >0.42 cm	Increased LV wall thickness is suggestive in the presence of normal or low QRS voltage on ECG and/or concomitant increased right ventricular wall thickness. While increased LV wall thickness is typically concentric, it can be asymmetric in ATTR cardiac amyloidosis ¹⁷²	Required
Stroke volume index	LV stroke volume index (<35 ml/m ²)	A low stroke volume index is non-specific but suggestive of cardiac amyloidosis	Required
LV mass	LV mass ≥91 g/m ² for men and ≥78 g/m ² for women (with papillary muscle included as part of LV mass measurement) ²⁰⁶	To quantify myocardial and amyloid mass	Required
Atrial size and function (based on Simpson's method)	Increased left atrial volume >163 ml for men and >131 ml for women ²⁰⁶ Increased right atrial volume >85 ml/m ² ²⁰⁶ Reduced atrial function: <29% for men and <35% for women. ²⁰⁶	Non-specific but important finding to support the diagnosis and potentially provide insight into risk for stroke or arterial embolism	Required
Pericardial effusion	Pericardial effusion	Non-specific, but when coupled with other CMR signs is suggestive of the diagnosis, especially in the setting of normal LV ejection fraction	Required
Amyloid Imaging			
LGE imaging	Abnormal LGE Pattern • Diffuse LGE • Subendocardial LGE • Patchy LGE • Difficulty in achieving myocardial nulling over a range of inversion times • Dark blood pool signal	Standard mag-IR LGE imaging is not recommended given difficulty in selecting the optimal inversion time (TI). Phase-sensitive reconstruction is preferred. Data acquisition should be obtained in every other RR interval. Quantification of LGE is challenging in amyloidosis and is not recommended for routine clinical practice.	Required
Myocardial signal suppression pattern	Abnormal myocardial signal suppression pattern Myocardium nulls before blood pool on Look Locker, Cine IR, or TI scout sequences		Recommended
Amyloid quantitation			
Native T1 mapping (pre-contrast)	Abnormal T1 mapping (criteria may vary based on the sequence used [MOLLI, ShMOLLI] and the field strength of the magnet)	Assess interstitial amyloid accumulation without gadolinium. Reference range should be based on a site's local calibrated values on specific field strengths.	Recommended
T1 mapping post-contrast (ECV estimation)	ECV >0.40 is highly suggestive of cardiac amyloidosis	Assess expansion of ECV from interstitial amyloid accumulation A. 1 pre- and 1 post-contrast measurement (15-minute post-contrast injection) B. 1 pre- and 3 post-contrast measurements (5-, 15-, and 25-minutes post contrast injection)	A. Recommended B. Optional
Reporting of CMR Findings in Cardiac Amyloidosis			
An overall interpretation of the CMR findings into categories of: • Not suggestive: Normal LV wall thickness, normal LV mass, no ventricular LGE, normal atrial size • Strongly suggestive: Increase LV wall thickness, increased LV mass, atrial enlargement, typical diffuse or global LGE pattern, difficulty in achieving myocardial nulling, significantly increased ECV (>0.40), small pericardial and/or pleural effusions • Equivocal: Findings not described above.			Required
Interpret the CMR results in the context of prior evaluation.			Recommended
Provide follow-up recommendations: Strongly suggestive CMR findings cannot distinguish AL from ATTR cardiac amyloidosis. Endomyocardial biopsy is frequently unnecessary in patients with strongly suggestive CMR findings and histologically defined systemic amyloidosis or diagnostic ^{99m} Tc-PYP/DPD/HMDP imaging. Consider evaluation (1) to exclude AL amyloidosis, evaluate for plasma cell dyscrasia (serum and urine immunofixation, serum FLC assay) and (2) to exclude ATTR cardiac amyloidosis, consider imaging with ^{99m} Tc-PYP/DPD/HMDP.			Recommended

2 mapping is currently not part of the standard clinical amyloidosis imaging protocol
 AL, amyloid light chain; ATTR, amyloid transthyretin; CMR, cardiac magnetic resonance imaging; ECV, extracellular volume; EF, ejection fraction; FLC, free light chain; LGE, late gadolinium enhancement; LV, left ventricular; MOLLI, modified Look-Locker inversion recovery; SSFP, steady state free precession; ShMOLLI, Shortened Modified Look-Locker Inversion Recovery; ULN, upper limit of normal and per Ref. 205 at mid-cavity level ULN for women/men were 7 mm/9 mm (long axis) and 7 mm/8 mm (short axis), respectively

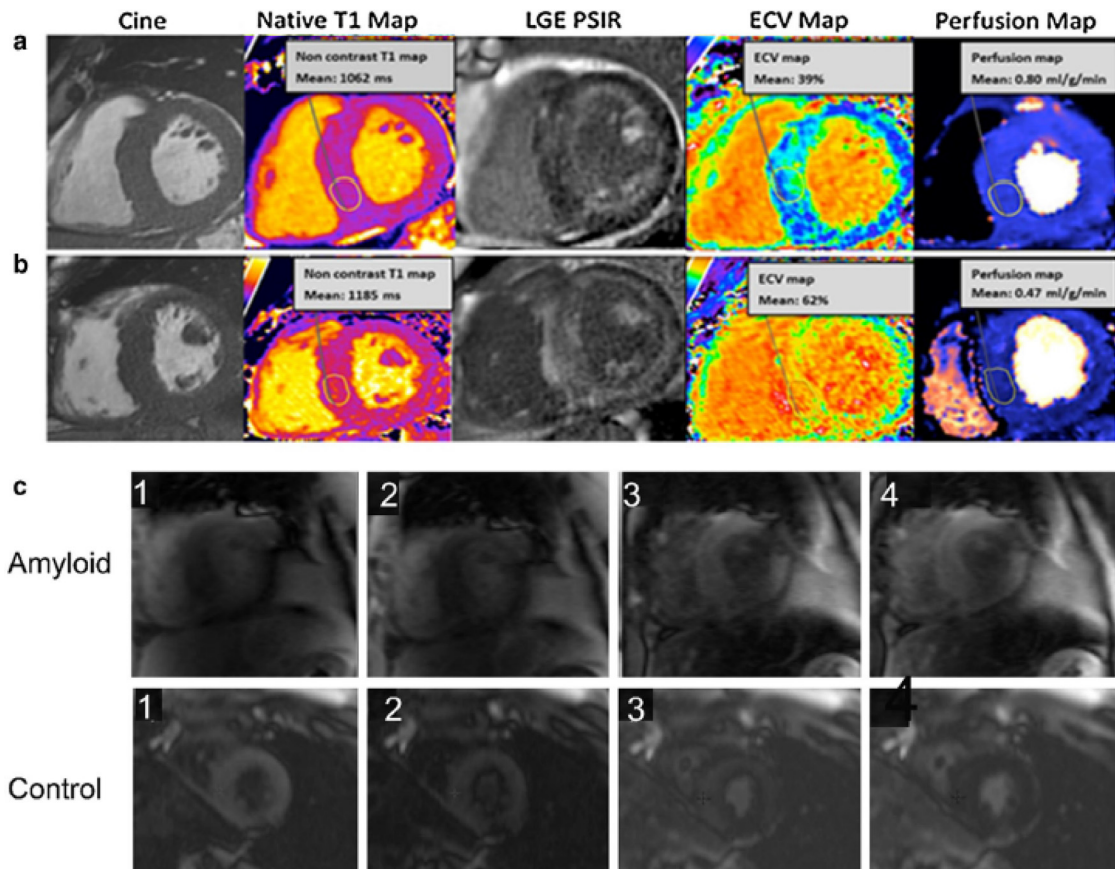


Fig. 5. Characteristic appearance of cardiac amyloidosis on CMR. Two patients [top and bottom rows, (a) and (b)] with cardiac amyloidosis: similar mass (cine), but significantly different amyloid burden, with the patient at the bottom (b) showing a significant higher amyloid burden (higher native T1, higher ECV, transmural LGE) and lower myocardial resting perfusion (also, after adjusting for ECV expansion). (c) Inversion scout images in 2 patients: top row, amyloid; bottom row, non-amyloid control. These images show a distinct pattern of myocardial and blood pool nulling. In the non-amyloid subject, the blood pool nulls prior to myocardium; in contrast, in the subject with cardiac amyloidosis, the myocardium nulls prior to the blood pool.

Key Recommendations for Standardized Imaging Techniques: CMR.

- CMR should be performed using standard parameters, as listed in this section.
- Cardiac structure, function, and PSIR LGE should be assessed and reported per SCMR guidelines.
- Cardiac amyloidosis-specific CMR markers, such as native T1 mapping and ECV, should be assessed and reported when available, as discussed in this document.
- An overall reporting on likelihood of cardiac amyloidosis based on imaging findings is recommended (eg, not suggestive, strongly suggestive, or equivocal for cardiac amyloidosis).

Radionuclide Imaging

^{99m}Tc-PYP/DPD/HMDP Imaging. Recommendations for standardized radionuclide image acquisition for cardiac amyloidosis using ^{99m}Tc-PYP/DPD/HMDP are provided in Table 4. Images should be acquired early (1 h) or late (2–3 h). There is a stepwise approach to interpretation as shown in Table 5. The first step of interpretation is to visually confirm diffuse myocardial radiotracer uptake and differentiate

this uptake from residual blood pool activity or overlapping bone using SPECT and planar images.

If myocardial uptake is confirmed visually, there are 2 approaches to differentiate AL from ATTR cardiac amyloidosis, depending on the tracer used and time between injection and scan acquisition. The 1-hour approach has been validated for ^{99m}Tc-PYP and involves generation of an elliptical/circular region-of-interest (ROI) over the heart on the anterior planar images with care to avoid sternal overlap and with size adjusted to maximize coverage of the heart without inclusion of adjacent lung. This ROI should be mirrored over the contralateral chest to adjust for background and rib uptake (Fig. 6). A semiquantitative H/CL ratio is calculated as a ratio-of-heart ROI mean counts to contralateral chest ROI mean counts; a ratio of ≥ 1.5 at 1 hour can accurately differentiate ATTR cardiac amyloidosis from AL cardiac amyloidosis.¹¹³

Alternatively, a 2- or 3-hour approach can be used (as typically performed for ^{99m}Tc-DPD/HMDP) in which a visual grading scale is used (Table 5). Grade 2 or Grade 3 myocardial uptake of ^{99m}Tc-PYP/DPD/HMDP, in the absence of a clonal disorder, is diagnostic of ATTR cardiac amyloidosis (Fig. 7). Both planar and SPECT imaging

Table 4. Recommendations for Standardized Acquisition of ^{99m}Tc -PYP/DPD/HMDP for Cardiac Amyloidosis

Imaging procedures	Parameters	Recommendation
Preparation	No specific preparation. No fasting required.	Required
Scan	Rest scan	Required
Dose	^{99m}Tc -PYP: 10-20 mCi (370-740 MBq) intravenously ^{99m}Tc -DPD: 10-20 mCi (370-740 MBq) intravenously ^{99m}Tc -HMDP: 10-20 mCi (370-740 MBq) intravenously	Recommended
Time between injection and acquisition: ^{99m}Tc -PYP	1 hour 3 hours	Recommended Optional, recommended if excess blood pool activity noted on 1-hour images
Time between injection and acquisition: ^{99m}Tc -DPD/HMDP	2 or 3 hours	Recommended
General imaging parameters		
Field of view	Heart Chest	Required Optional
Image type	Planar SPECT if planar is positive	Recommended Required
Position	Supine Upright	Required Optional
Energy window	140 keV, 15–20%	Required
Collimators	Low energy, high resolution	Recommended
Matrix-Planar	256 x 256	Recommended
Matrix-SPECT	128 x 128 (at least 64 by 64 is required)	Recommended
Pixel size	3.5–6.5 mm	Recommended
Planar imaging specific parameters		
Number of views*	Anterior and lateral	Required
Detector configuration	90°	Recommended
Image duration (count based)	750,000 counts	Recommended
Magnification	1.46	Recommended
SPECT imaging specific parameters		
Angular range	180°	Required
Detector configuration	90°	Recommended
Angular range	360°	Optional
Detector configuration	180°	Optional
ECG gating	Off; Non-gated imaging	Recommended
Number of views/detector	40/32	Recommended
Time per stop	20 seconds / 25 seconds	Recommended
Magnification	1.46 (180° angular range) 1.0 (360° angular range)	Recommended

Adapted from Ref. 207

ECG, electrocardiogram; PYP, pyrophosphate

*Anterior and lateral views are obtained at the same time; lateral planar views or SPECT imaging may help separate sternal from myocardial uptake

Table 5. Recommendations for Interpretation ^{99m}Tc -PYP/DPD/HMDP for Cardiac Amyloidosis

Step 1: Visual interpretation to diagnose ATTR cardiac amyloidosis	
- Evaluate planar and SPECT images to confirm diffuse radiotracer uptake in the myocardium. - Differentiate myocardial radiotracer uptake from residual blood pool activity, focal myocardial infarct, and overlapping bone (e.g., from rib hot spots from fractures). If excess blood-pool activity is noted on the 1-hour SPECT images, recommend repeat SPECT imaging at 3 hours. - If myocardial tracer uptake is visually present on SPECT, proceed to step 2, semi-quantitative grading to distinguish ATTR from AL cardiac amyloidosis using either the 1- or 3-hour approach.	
Step 2: Semi-quantitative grading to distinguish AL from ATTR cardiac amyloidosis (1- or 3-hour approach)	
1-Hour Approach (validated for ^{99m}Tc -PYP):	
- An elliptical/circular ROI should be drawn over the heart on the anterior planar images with care to avoid sternal overlap and with size adjusted to maximize coverage of the heart without inclusion of adjacent lung. This ROI (same size) should be mirrored over the contralateral chest to adjust for background and rib uptake (see Figure 6). - A H/CL ratio is calculated as the fraction of heart ROI mean counts to contralateral chest ROI mean counts. - H/CL ratios of ≥ 1.5 at one hour can accurately identify ATTR cardiac amyloidosis if systemic AL amyloidosis is excluded.¹¹³	
3-Hour Approach:	
Examine 3-hour images for relative tracer uptake in the myocardium relative to ribs and grade using the following scale:	
Grade 0	No myocardial uptake and normal bone uptake
Grade 1	Myocardial uptake less than rib uptake
Grade 2	Myocardial uptake equal to rib uptake
Grade 3	Myocardial uptake greater than rib uptake with mild/absent rib uptake
See Figure 7. Grade 2 or Grade 3 uptake is consistent with ATTR cardiac amyloidosis if a monoclonal plasma cell dyscrasia is excluded, as this degree of uptake can be seen in >20% of patients with AL cardiac amyloidosis.³ Grade 0 and Grade 1 uptake may be observed in AL cardiac amyloidosis and warrants further evaluation to exclude AL amyloidosis.³ The writing group would like to emphasize the importance of excluding a monoclonal process with serum/urine immunofixation and a serum free light-chains assay in all patients with suspected amyloidosis.	

Adapted from Ref. 207

AL, amyloid light chain; ATTR, amyloid transthyretin; H/CL, heart/contralateral lung; ROI, region of interest

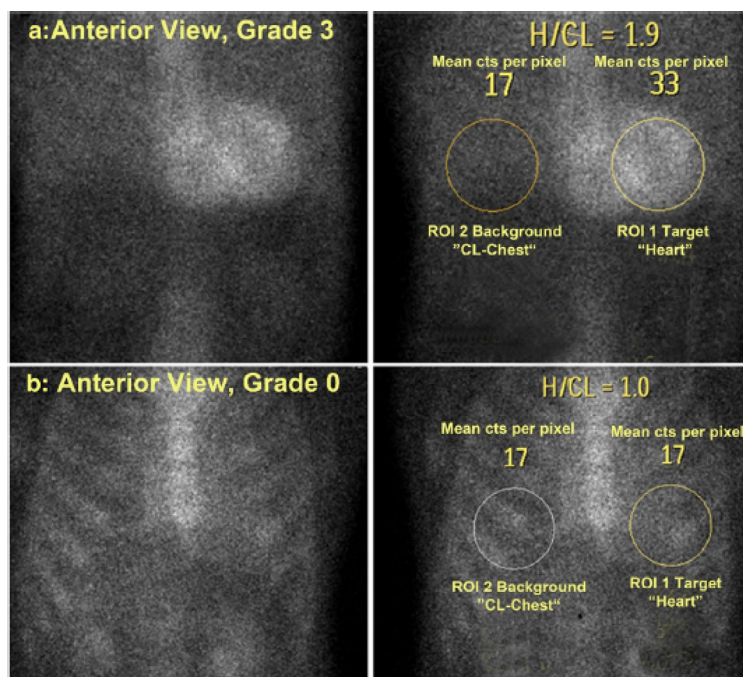


Fig. 6. Characteristic appearance of cardiac amyloidosis on ^{99m}Tc -PYP/DPD/HMDP imaging. Semiquantitative H/CL Ratio on ^{99m}Tc -PYP planar imaging. Anterior planar chest views 1 hour after injection of ^{99m}Tc -PYP a patient with Grade 3 (a), and Grade 0 (b) ^{99m}Tc -PYP uptake. On the right are the corresponding H/CL (heart/contralateral lung) lung-ratio methodology with measurement of mean counts per pixel for target (heart) and background (contralateral chest). As shown in this figure, the ROI should be positioned to minimize overlap with sternal or focal rib uptake and maximize coverage of the heart without including adjacent lung.

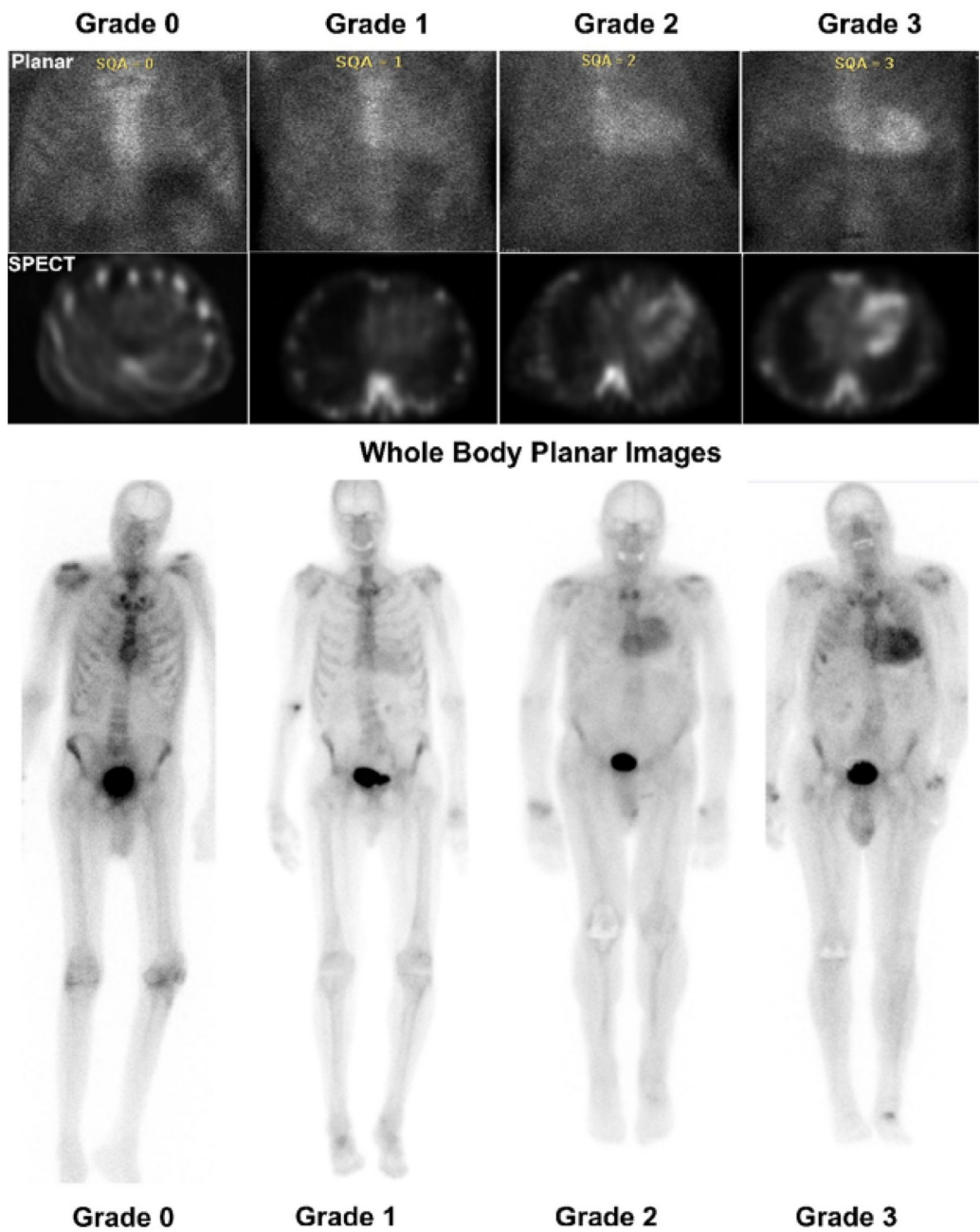


Fig. 7. ^{99m}Tc -PYP/DPD/HMDP. Anterior planar chest images (top row), SPECT cardiac imaging (middle row), and planar whole-body imaging (bottom row). Cardiac uptake is visually compared with surrounding ribs for a visual grading score as described in Table 5. Images with Grade 0, Grade 1, Grade 2, and Grade 3 myocardial uptake of ^{99m}Tc -PYP are shown. (Top panel provided by ASNC Cardiac Amyloidosis Practice Points²⁰⁹).

should be reviewed and interpreted using visual and quantitative approaches irrespective of the timing of acquisition. SPECT imaging is necessary for studies that show planar myocardial uptake because they can help differentiate myocardial uptake from blood pool or overlying bone uptake. Interpretation should also include comment on focal versus diffuse radiotracer uptake; diffuse uptake is typically consistent with cardiac amyloidosis, whereas focal uptake may represent early cardiac amyloidosis but has also been described in acute or subacute myocardial infarction.

Guidelines for standardized reporting are provided in Table 6. An H/CL ratio may be falsely low in patients who had suffered a prior large remote myocardial infarction; myocardial uptake of the tracer will be limited to non-infarcted zone. Careful evaluation of these imaging using SPECT and nonplanar image display are recommended to visualize regional uptake. ^{123}I -mIBG Sympathetic Innervation Tracer. An overview of the imaging acquisition parameters for ^{123}I -mIBG

Table 6. Recommendations for Standardized Reporting of ^{99m}Tc -PYP/DPD/HMDP Imaging for Cardiac Amyloidosis

Parameters	Elements
Demographics	Patient name, age, sex, reason for the test, date of study, prior imaging procedures, biopsy results if available (Required)
Methods	Imaging technique, radiotracer dose and mode of administration, interval between injection and scan, scan technique (planar and SPECT) (Required)
Findings	Image quality Visual scan interpretation (Required) Semi-quantitative interpretation in relation to rib uptake (Required) Quantitative findings H/CL lung ratio (Optional; recommended for positive scans)
Ancillary findings	Whole-body imaging if planar whole-body images are acquired (Optional) Interpret CT for attenuation correction if SPECT/CT scanners are used (Recommended)
Conclusions	- An overall interpretation of the findings into categories of 1) not suggestive of ATTR cardiac amyloidosis; 2) strongly suggestive of ATTR cardiac amyloidosis or 3) equivocal for ATTR cardiac amyloidosis Not suggestive: A semi-quantitative visual grade of 0. Equivocal: If myocardial uptake of ^{99m}Tc-PYP/DPD/HMDP is visually confirmed, a semi-quantitative visual grade of 1 or H/CL ratio 1-1.5. Strongly suggestive: If myocardial uptake of ^{99m}Tc-PYP/DPD/HMDP is visually confirmed, a semi-quantitative visual grade of 2 or 3 - Interpret the results in the context of prior cardiac evaluation and evaluation for systemic AL amyloidosis using serum-free light chain assay, serum immunofixation, and urine immunofixation studies. A ^{99m}Tc-PYP/DPD/HMDP scan does not exclude AL cardiac amyloidosis. Therefore: - If echo/CMR are strongly suggestive of cardiac amyloidosis, and ^{99m}Tc-PYP/DPD/HMDP is negative or equivocal, consider further evaluation for AL amyloidosis by serum FLCs, serum, and urine immunofixation and referral to a hematologist or an amyloidosis expert. Endomyocardial biopsy may be considered. - A positive ^{99m}Tc-PYP/DPD/HMDP scan with abnormal FLC evaluation, consider referral to a hematologist or an amyloidosis expert.

Adapted from Ref. 207

AL, amyloid light chain; ATTR, amyloid transthyretin; CMR, cardiovascular magnetic resonance; echo, echocardiography; FLC, free light chain; H/CL, heart-to-contralateral lung ratio

is available in the Appendix. Sources of variability in late HMR include non-homogeneity in ^{123}I -mIBG imaging acquisition; differing gamma camera systems; and low-versus medium-energy collimators.^{131,186,202,203} Recommendations for the reporting of ^{123}I -mIBG are provided in the Appendix and are predominately based on the HMR and washout-rate quantification. As with ^{99m}Tc -PYP/DPD/HMDP, SPECT imaging is of value in addition to planar imaging to evaluate regional cardiac sympathetic innervation abnormalities. The majority of patients (in both AL and ATTR cardiac amyloidosis) with low HMR show reduced tracer accumulation in the inferolateral segments.^{97,98,130,135,137,139} This, however, is not a finding specific to cardiac amyloidosis; reduced radiotracer uptake in the inferolateral myocardial wall is also reported in healthy control subjects because of physiological over projection of ^{123}I -mIBG accumulation of the liver into this region.²⁰⁴ Also, this technique should be avoided in

patients with suspected cardiac amyloidosis and prior myocardial infarction.

Key Recommendations for Standardized Image Techniques: Radionuclide Imaging.

- ^{99m}Tc -PYP/DPD/HMDP and ^{123}I -mIBG imaging should be performed using standard protocols as discussed in this section.
- SPECT imaging is useful particularly in positive or equivocal cases to differentiate myocardial from blood pool signal and to describe regional heterogeneity.
- Visual and semiquantitative interpretation of ^{99m}Tc -PYP/DPD/HMDP planar and SPECT images should be employed to evaluate heart-to-bone ratio and/or H/CL lung ratio. The HMR is used to interpret ^{123}I -mIBG images.
- An overall reporting on likelihood of amyloidosis based on imaging findings is recommended (eg, not suggestive,

strongly suggestive, or equivocal for cardiac amyloidosis and for extra-cardiac findings).

Future Directions

The field of imaging in cardiac amyloidosis is expanding rapidly and more research is needed in several key areas.

- Early detection with imaging remains an unmet need in cardiac amyloidosis, and techniques that identify disease at an earlier stage are needed. ^{99m}Tc PYP/DPD/HMDP has the potential for early detection of ATTR cardiac amyloidosis prior to echocardiography and CMR. This needs to be further validated.
- Molecular imaging techniques, including amyloid binding PET radionuclide tracers and ECV by CMR are particularly well suited to detect early disease. Further studies are needed.
- Early detection of cardiac amyloidosis could allow targeted therapy prior to symptom onset and improve clinical outcomes. This needs to be studied further.
- Methods for quantitative assessment of systemic and cardiac burden of amyloidosis are needed. ECV assessment by CMR, and ^{18}F -labeled PET tracers have the potential to provide accurate quantification but require additional evaluation and more widespread dissemination of technology and broader clinical use to reach their full potential.
- Precise detection of changes in the burden of cardiac amyloidosis using imaging can allow evaluation of the efficacy of emerging novel therapies aimed at stabilization and even resorption of amyloid fibrils.
- Advanced echocardiography, including 3D echocardiographic strain, dynamic echocardiography, left atrial

mechanics, and automated, machine learning-based methods over standard approaches are being investigated.

- Prospective studies evaluating the incremental diagnostic and prognostic value of noninvasive imaging techniques, including advanced echocardiographic methods, ^{99m}Tc PYP/DPD/HMDP, ^{123}I -*m*IBG, and CMR should be undertaken. The incremental value of imaging markers over clinical and laboratory markers needs to be studied further.
- The majority of existing literature arises from small, single-center studies of highly selected patients. Multicenter studies, including larger patient cohorts and standardized imaging methods, are needed to advance the evaluation and management of cardiac amyloidosis. In particular, large prospective studies are needed to validate the clinical utility of cardiac imaging in assessing the response to therapy and predicting clinical outcome.

Summary

The purpose of Part 1 of this consensus statement has been to establish the available diagnostic and prognostic literature for imaging in cardiac amyloidosis and provide comprehensive expert recommendations based on this evidence and expert opinion regarding the role of imaging in cardiac amyloidosis, including standardized image acquisition, interpretation, and reporting. We hope that use of these consensus recommendations on standardized imaging techniques will improve patient care and outcomes. We also hope we have identified gaps in the literature that can spur relevant research to broaden our understanding of this complex disease and support guideline development.

Disclosures

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Jamieson M. Bourque, MD Angela Dispenzieri, MD		Astellas Celgene, Takeda, Janssen, Pfizer, Alnylam Pharmaceuticals, Prothena Bioscience	Pfizer		Locus Health
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Appendix

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A summary of literature that supports the recommendations provided in this consensus statement on the prognostic value of echocardiography (Table 7); diagnostic and prognostic value of CMR (Tables 8 and 9), and diagnostic and prognostic value of radionuclide imaging with ^{99m}Tc -PYP/DPD/HMDP (Tables 10, 11, 12, 13, and 14) in the evaluation of cardiac amyloidosis are provided in the Appendix. The diagnostic value, prognostic value, standardized image acquisition and reporting of ^{123}I -mIBG in cardiac amyloidosis are provided in Tables 15, 16, 17, and 18.

Table 7. Key Literature Summarizing the Utility of Echocardiography for Risk Assessment in Cardiac Amyloidosis

First author	Year	N patients	N controls	Design	Follow-up period (months)	Outcome	Event rate (annualized)	Hazard ratio	Comments
Left ventricle									
Mohy ¹⁶⁰	2017	63	87	Retrospective	24 (range 0–216)	All-cause mortality	Year 1: 31%	2.29	Left-sided valve thickening
Siepen ¹⁶⁴	2018	191	0	Retrospective	26.2 ± 1.7	All-cause mortality	Year 1: 3.7%	0.173	MAPSE ≤ 8.8 mm
Liu ¹⁵⁶	2014	41	21	Retrospective	16 (quartiles 5–35)	All-cause mortality	Year 1: 33%	7.5	Longitudinal early diastolic strain rate Cut-off 0.85
Barros-Gomes ¹⁴⁷	2017	63	87	Retrospective	40.8 (31.2–51.6)	All-cause mortality	Year 1: 11%	4.71	GLS _{GE} ≥ 14.81%
Ochs ¹⁶¹	2016	36	53	Retrospective	12	Transplant-free survival	Year 1: 36%	0.66	AAPSE < 5 mm
Senapati ¹⁶³	2016	49	48	Retrospective	21.2 (quartiles 5.7–34.3)	All-cause mortality or heart transplantation	Year 1: 31%	2.45	RRSR ≥ 1.19; 59 AL, 38 ATTR
Tendler ⁶⁵	2015	36 AL 34 ATTR 32	22	Retrospective	22.3 ± 21.4	Event-free survival	Year 1: 45%	2.841 AL 3.39 ATTR 1.26	MCF < 30, AL and ATTR
Riffel ¹⁶²	2015	50	70	Retrospective	12	All-cause mortality or heart transplantation	Year 1: 42%	0.67	Long axis shortening ≥ 5.8
Hu ¹⁵¹	2015	8	16	Retrospective	16 (7.3–15.7)	All-cause mortality	67% during follow-up	5.47	LSys > 3%
Liu ¹⁵⁵	2017	19	39	Retrospective	12	All-cause mortality	Year 1: 21%	8.48	Tei index ≥ 0.9
Perlini ⁹	2014	221	121	Retrospective	18.4	All-cause mortality	n. a.	X ² 58.2	Midwall fractional shortening ≤ 12.04%
Migrino ¹⁵⁷	2014	27	15	Prospective	60 months	All-cause mortality	Year 1: 40.9%	5.07	Left ventricular ejection time ≤ 240 ms
Bellavia ⁵⁹	2011	23	28	Prospective	34 (0.9–64)	All-cause mortality	32% during follow-up	6.5	V ₁ II–III or the V ₁ VI gene family vs V ₁ or V ₂ –I families
Koyama ¹⁵²	2010	70	49	Prospective	6.2 ± 4.5	All-cause mortality	Year 1: not given (26.9% during follow-up)	n. a.	Basal systolic strain ≤ 13%
Kristen ²⁰⁸	2007	17	22	Retrospective	24.7 ± 3.1	All-cause mortality and heart transplant	Year 1: 27.9%	n. a.	Progression of LV wall thickness < 0.2 mm/month
Koyama ¹⁵³	2002	133	75	Retrospective	10.2 ± 5.2	All-cause mortality	n. a.	25.6	CV-IB ≤ 5.35 dB
Tei ¹⁶⁵	1996	45	45	Retrospective	36 months	All-cause mortality	n. a.	Chi square 4.6	Tei index ≤ 0.77
Cueto-Garcia ⁵⁰	1985	71	44	Retrospective	n. a.	All-cause mortality	CHF 83% no CHF 41%	n. a.	CHF
Left atrium									
Mohy ¹⁶⁰	2017	77	39	Prospective	19 ± 10 (IQR 9–26)	All-cause mortality	80 ± 5	0.94	3D positive atrial longitudinal strain > 14%
Mohy ¹⁵⁹	2011	53	58	Retrospective	33.6 ± 34.8	All-cause mortality	Year 1: 28.9%	2.47	AL
Right ventricle									
Bodez ¹⁴⁸	2016	82	47	Prospective	8 (2–16)	Death, heart transplant, acute heart failure	Year 1: 16.3%	0.85	TAPSE ≥ 14 mm
Bellavia ⁷⁰	2012	47	59	Prospective	53 (0.6–75)	All-cause mortality	Year 1: 49%	1.3	Strain rate of the RV free wall middle segment ≥ – 1.37/s
Cappelli ¹⁴⁹	2012	52	31	Prospective	19 ± 12 (median 20)	Cardiac death	Year 1: 11.5% (missing values not mentioned)	1.128	RV longitudinal strain > 9%
Other									
Damy ¹⁵⁰	2016	84	149	Retrospective	17 (6–35)	All-cause mortality	Year 2: 53% (AL) 58% (ATTRwt) 28% (ATTRv)	2.25	Pericardial effusion

AL, light-chain amyloidosis; ATTR, transthyretin amyloidosis; ATTRv, amyloidosis due to transthyretin gene mutations; ATTRwt, amyloidosis due to deposition of wild-type transthyretin (senile amyloidosis); AAPSE, anterior aortic plane systolic excursion; CV-IB, cyclic variation of integrated back scatter; CHF, congestive heart failure; GLS, global longitudinal strain; LV, left ventricular; LSys, longitudinal strain; GLSGE, global longitudinal strain; MCF, myocardial contraction fraction; MAPSE, mitral annular plane systolic excursion; n. a., not available; N number; RV, right ventricular; RRSR, relative regional strain ratio; TAPSE, tricuspid annular plane systolic excursion

Table 8. Key Literature Summarizing the Diagnostic Value of CMR for Cardiac Amyloidosis

First author	Year	N Patients	N Controls	Patient cohort	ATTR/ AL	Method	Criterion	Sensitivity	Specificity	Comments
Maceira ⁷⁴	2005	30	16	Systemic (extracardiac) amyloidosis confirmed by non-cardiac histology and cardiac involvement by echocardiographic criteria	AL/ ATTR	LGE, T1 mapping at 4 minutes after contrast	Subendocardium-blood T1 difference of 191 ms at 4 minutes after injection	90%	87%	Limitations of a cohort controlled study, small sample size, and selection bias
Vogelsberg ⁷⁷	2008	33	0	Suspected cardiac amyloidosis patients referred for to undergo both endomyocardial biopsy and CMR	Did not specify	LGE	Diffuse subendocardial LGE involvement	80%	94%	Small sample size and selection bias
Austin ⁸¹	2009	47	0	Suspected cardiac amyloidosis patients	AL/ATTR	LGE	Diffuse subendocardial LGE involvement	88%	90%	LGE more accurate than ECG and TTE parameters combined
Ruberg ⁸⁰	2009	28	0	Systemic (extracardiac) AL amyloidosis patients with variable cardiac involvement	AL	LGE	Presence and size of LGE (6 SD threshold)	86%	86%	
Syed ⁷⁸	2010	120	0	Systemic (extracardiac) AL amyloidosis confirmed either by cardiac histology (n = 35) or monoclonal protein/plasma cell abnormalities (n = 85)	AL/ ATTR	LGE	Any LGE abnormality	Cardiac histology group: LGE sensitivity 97% in detecting CA by EMB	NA	Non-cardiac histology group: LGE abnormality more prevalent than echocardiographic criteria of CA (69% vs 58%)
Karamitsos ⁸²	2013	53	53 (36 normal and 17 pts with aortic stenosis)	Systemic (extracardiac) AL amyloidosis patients with variable cardiac involvement	AL	Native T1 mapping (ShMOLLI)	Native T1 value 1020 ms	92%	91%	
White ⁷⁹	2014	90	64 pts with HHD	Suspected cardiac amyloidosis patients	AL/ ATTR	Visual T1 assessment	Myocardial T1 curve crosses the null point before blood T1 curve	100%	70%	
Kwong ⁶⁸	2015	22	37 pts with HHD and 22 pts with DCM	Systemic amyloidosis confirmed by cardiac histology	AL/ ATTR	Left atrial LGE > 1/3 all left atrial segments	Number of left atrial segments with abnormal LGE	76%	94%	Limitations of a cohort controlled study, small sample size, and selection bias
Zhao ⁸³	2016	257	0	Meta-analysis of 7 published studies	AL/ ATTR	LGE	Presence of a typical LGE pattern	85%	92%	Binary LGE classification, a lack of consideration of cardiac amyloidosis subtypes, lack of T1 mapping data, limitations of a meta-analysis
Martinez-Naharro ¹⁷²	2017	313	0	ATTR cardiac amyloidosis pts corroborated by ⁹⁹ Tc SPECT (n = 201) or TTR mutation (n = 12) and AL cardiac amyloidosis pts (n = 50)	AL/ ATTR	Asymmetric increase in LV septal thickness, typical LGE pattern		79%	NA	Asymmetrical increase in septal wall thickness is common and LGE typical in ATTR

AL, light chain amyloidosis; ATTR, transthyretin amyloidosis; CMR, cardiac magnetic resonance imaging; DCM, dilated cardiomyopathy; ECG, electrocardiogram; HHD, hypertensive heart disease; LGE, late gadolinium enhancement; LV, left ventricular; NA, not applicable; SD, standard deviation; ShMOLLI, Shortened Modified Look-Locker Inversion recovery; TTE, transthoracic echocardiography; TTR, transthyretin

Table 9. Key Literature Summarizing the Utility of CMR for Risk Assessment in Cardiac Amyloidosis

First author	Year	N patients	Design	Follow-up period (months)	CMR sequence	Outcome	Event rate		Odds ratio or Hazard ratio	Comments
Late gadolinium enhancement (LGE)										
Maceira ²⁰⁹	2008	29 (25 AL, 4 ATTR)	Prospective	20	2D, Segmented IR GE sequence (LGE); IR segmented RSP cine 30 ms/frame	All-cause mortality	LGE + 0.35	LGE - 0.37	OR 0.90*	LGE not predictive, mortality was predicted by gadolinium kinetics
Migrino ²¹⁰	2009	29 (all AL)	Prospective	29	PSIR (LGE)	All-cause mortality	0.25	0	OR 19.84*	LGE predicted mortality
Austin ⁸¹	2009	25 (15 AL, 9 ATTR, 1 miscellaneous)	Retrospective	12	PSIR (LGE)	All-cause mortality	0.48	0.25	OR 2.73*	LGE predicted mortality
Ruberg ⁸⁰	2009	28 (all AL)	Prospective	29	2D, spoiled segmented IR GE sequence (LGE)	All-cause mortality	0.09	0.05	OR 2.13*	LGE did not predict mortality
Mekinian ¹⁶⁷	2010	29 (All AL)	Retrospective	32	IR GE (LGE). Used the LL sequence to define CMR+ and CMR-	All-cause mortality	0.38	0.04	OR 132.60*	Abnormal nulling on the LL sequence predicted mortality
White ⁷⁹	2014	46 (41 AL, 2 ATTR, three miscellaneous)	Prospective]	29	2D, segmented IR GE sequence (LGE), T1 scout (T1 visual assessment)	All-cause mortality	0.32	0.10	OR 9.75*	Diffuse enhancement by visual T1 assessment predicted mortality
Fontana ⁹⁰	2015	250 (119 AL, 122 ATTR, 9 mutation carriers)	Prospective	24	PSIR (43% of patients), magnitude-IR LGE. Post contrast T1 at equilibrium of contrast used to offset errors in nulling before adoption of PSIR	All-cause mortality	0.16	0.05	OR 4.44*	Transmural LGE predicted mortality
T1, T2 mapping										
Kotecha ⁸⁹	2018	286 (100 AL, 163 ATTR, 12 suspected ATTR, 11 mutation carriers)	Prospective	23	T2 mapping	All-cause mortality	0.26		HR 1.48 for 3 ms change	T2 is a predictor of prognosis in AL amyloidosis, not in ATTR
Martinez-Naharro ⁸⁹	2018	227 (215 ATTR, 12 mutation carriers)	Prospective	32	T1 mapping ShMOLLI	All-cause mortality	0.42		HR Native T1 (59 ms change) 1.22 HR ECV (0.03 change) 1.16	Native T1 and ECV predicted mortality, only ECV remained independent after adjusting for known predictors
Lin ²¹¹	2018	82 (all AL)	Prospective	8	T1 mapping MOLLI	All-cause mortality	0.26		HR ECV (> 0.44) 7.25 HR LGE+ 4.80	ECV and LGE predicted mortality. Native T1 did not predict mortality
Banyersad ¹⁷¹	2015	100 (all AL)	Prospective	23	T1 mapping ShMOLLI	All-cause mortality	0.25		HR ECV (> 0.45) 3.84 HR native Native T1 (> 1044 ms) 5.39	Native T1 and ECV predicted mortality
Martinez-Naharro ¹⁷²	2017	292 (263 ATTR, 17 suspected ATTR, 12 mutation carriers)	Prospective	19	T1 mapping ShMOLLI or MOLLI	All-cause mortality	0.22		HR ECV (0.03 increase) 1.16	ECV predicted mortality

AL, light-chain amyloidosis; ATTR, transthyretin-related amyloidosis; LGE, late gadolinium enhancement; PSIR, phase sensitive inversion recovery; LL, look locker; IR, inversion recovery; GE, gradient echo

*OR from Raina et al ¹⁷⁰

AL, light-chain amyloidosis; ATTR, transthyretin-related amyloidosis; LGE, late gadolinium enhancement; PSIR, phase sensitive inversion recovery; LL, look locker; IR, inversion recovery; GE, gradient echo

* OR from Raina et al ¹⁷⁰

Table 10. Key Literature Summarizing the Diagnostic Value of ^{99m}Tc -PYP Radionuclide Imaging for Cardiac Amyloidosis

First author	Year	N patients	N controls	Planar/SPECT	Patient cohort	Amyloidosis type	Criterion	Sensitivity	Specificity	Comments
Wizenberg ⁴⁷	1982	10	0	Planar	Biopsy proven	Not defined	$\geq 2+$	100%	-	
Falk ⁴³	1983	20		Planar	Biopsy proven	AL	$\geq 2+$	91%	90%	
Lee ²¹²	1983	7	10	Planar	Biopsy proven	AL	$\geq 2+$	86%	100%	MDP lower sensitivity vs PYP
Eriksson ²¹³	1984	12	0		FAP	ATTR	$\geq 2+$	33%	-	
Leinonen ²¹⁴	1984	6	0	Planar	Systemic amyloid	5 AA	$\geq 2+$	0	-	
Falk ²¹⁵	1984	9	0	Planar	FAP	ATTR	$\geq 2+$	77.7%	-	
Gertz ⁴⁴	1987	34		Planar/SPECT	Biopsy proven	Not defined	$\geq 1+$	21% (Retro)	25%	
Hongo ²¹⁶	1987	15		Planar/SPECT	FAP	ATTR	$\geq 2+$	85%* (Prosp)	95%	
Goldstein ²¹⁷	1989	32	0	Planar	Elderly ≥ 85 yrs screening for ATTRwt	ATTR	$\geq 2+$	67%	-	
Hartmann ⁴⁵	1990	7		Planar/first pass		Not defined	$\geq 2+$	71.4%		
Foumier ²¹⁸	1993	9	6	Planar	FAP	Majority ATTR	Scintigraphic index		-	
Yamamoto ⁴⁸	2012	13	37	Planar/SPECT	CHF, LVH, Suspected cardiac amyloidosis	Not defined	PYP score	85%	95%	
Bokhari ¹¹³	2013	45		Planar/SPECT	Biopsy proven	AL/ATTR	$\geq 2+$	97% (ATTR)	100%	ATTR vs AL
Castano ¹¹⁴	2016	121	16	Planar	Retrospective	ATTR/AL	H/CL; visual	91%	92%	Multicenter
Gillmore ³	2016	1217	360	Planar	Retrospective, referral centers	ATTR/AL	H/CL; visual	74%	100%	+ PYP and absence of monoclonal gammopathy

AL, light chain amyloidosis; ATTR, transthyretin amyloidosis; AA, Apo serum amyloid A; CHF, congestive heart failure; CL, contralateral; DPD, 3,3-diphosphono-1,2-propanodicarboxylic acid; HMDP, hydroxymethylene diphosphonate; H, heart; H/CL, heart/contralateral lung; LVH, left ventricular hypertrophy; FAP, familial amyloid polyneuropathy; PYP, ^{99m}Tc pyrophosphate; DPD, ^{99m}Tc -3,3-diphosphono-1,2-propanodicarboxylic acid (DPD); HMDP, ^{99m}Tc -hydroxymethylene diphosphonate; MDP, ^{99m}Tc -methylene diphosphonate

Table 11. Key Literature Summarizing the Diagnostic Value of ^{99m}Tc -DPD/HMDP Radionuclide Imaging for Cardiac Amyloidosis

First author	Year	N patients	N controls	Isotope	Planar/SPECT	Patient cohort/diagnostic standard	ATTR/AL/ others	Crite- rion	Sensi- tivity	Speci- ficity	Comments
Puille ²¹⁹	2002	8	10	DPD	Planar/SPECT	Prospective/rectal biopsy with IHC + genotyping	ATTRv	HR WBR	-	-	HR/WBR >> in ATTR pts (vs controls) ($p < 0.001$)
Perugini ¹¹²	2005	25	10	DPD	Planar/SPECT	Prospective/cardiac biopsy with IHC + genotyping + echo	ATTR AL	(a) HR: H/WB (b) Visual score ≥ 1	(a) > in ATTR pts (b) 100% for ATTR	(b) 100%	Evaluation of diagnostic accuracy in the etiological diagnosis (identification of ATTR pts)
Rapezzi ¹⁰⁵	2011	40	23	DPD	Planar/SPECT	Retrospective/FPA or cardiac biopsy + genotyping + echo	ATTRv	Moderate/intense cardiac uptake (visual score ≥ 2)	100% in pts with CA	-	4/23 pts without cardiac amyloidosis had a positive bone-scan (early diagnosis)
Rapezzi ²²⁰	2011	79	15	DPD	Planar/SPECT	Retrospective/FPA or cardiac biopsy + genotyping + echo	ATTR AL	(a) HR (b) H/WB (c) Visual score ≥ 1	(a) > in ATTR (b) 71% in AL+ (c) 100% in ATTR pts	(a) 100% (b) 100% (c) -	Evaluation of diagnostic accuracy in the etiological diagnosis (identification of ATTR pts)
Quarta ¹⁰³	2012	46	16	DPD	Planar/SPECT	Prospective/Cardiac biopsy + genotyping	ATTRwt	Visual score ≥ 2	100%	100%	Evaluation of the diagnostic accuracy in the differential diagnosis with other cardiomyopathies mimicking cardiac amyloidosis
De Haro ²²¹	2012	19	-	DPD	Planar/SPECT	Retrospective/various organ biopsy + genotyping + echo	ATTR AL	Visual score ≥ 2	100%	100%	Evaluation of the diagnostic accuracy in the etiological diagnosis (identification of ATTR pts)
Hutt ¹⁰⁶	2014	321	-	DPD	Planar/SPECT	Prospective/suspected or proven (biopsy driven) amyloidosis—cardiac amyloidosis diagnosed by echo/CMR	ATTR AL AA Other	Visual score ≥ 1	100% (for detecting ATTR pts with cardiac amyloidosis)	-	In 85 pts amyloidosis was ultimately excluded
Ferreira ²²²	2015	19	-	DPD/ MDP	Planar/ SPECT	Prospective/echo + geno-typing	ATTR	Visual score ≥ 1 (DPD)	50%	94%	Evaluated accuracy in detecting CA
Galat ¹⁰²	2015	69	52 (other CMP [37 with LVH; 15 no LVH])	HMDP	Planar/SPECT	Prospective/various organ biopsy with IHC + genotyping + echo	ATTR AL	(a) Visual score ≥ 1 (amyloidosis vs control) (b) Visual score ≥ 2 (identification of ATTR pts with cardiac amyloidosis)	(a) 75% (b) 83%	(a) 100% (b) 100%	Evaluation of diagnostic accuracy in the etiological diagnosis (in identification of ATTR pts) All ATTRv without cardiac amyloidosis and carriers had no cardiac uptake
Pilebro ²²³	2016	55	-	DPD	Planar/SPECT	Retrospective/FPA or cardiac biopsy	ATTRv	Visual score	-	-	Evaluation of different cardiac uptake according to amyloid fibril type (strong association between DPD uptake and fibrils type)
Abulizi ²²⁴	2016	6	-	HMDP vs DPD	Planar/SPECT	Prospective/various organ biopsy + genotyping	ATTRv	H/medi-astinum	-	-	HMDP and DPD showed comparable cardiac uptake
Galat ²²⁵	2016	135	31 (pts with LVH) vs late phase)	HMDP	Planar/SPECT	Prospective/various organ biopsy with IHC + genotyping + echo	ATTR AL	a) H/ Mediastinum >1.210 (early phase)	100% (for detecting ATTR pts vs AL)	100% (for detecting ATTR pts vs AL)	All controls had no cardiac uptake
Cappelli ¹⁰¹	2017	65	20	HMDP	Planar/SPECT	Retrospective/various organ biopsy with IHC + genotyping + echo	ATTR AL	Visual score ≥ 1	100% (for detecting ATTR pts vs AL)	100%	Evaluation of diagnostic accuracy in the etiological diagnosis (identification of ATTR pts)
Van Der Gucht ²²⁶	2017	61	-	HMDP	Planar/SPECT	Prospective/various organ biopsy with IHC + genotyping + echo	ATTR	Cardiac uptake in the early phase	100%	-	Evaluation of LV distribution of early phase uptake
Moore ²²⁷	2017	21	-	DPD	Planar/SPECT	Prospective/various organ biopsy with IHC + genotyping + echo	ATTR AL	Visual score ≥ 1	100% (for detecting ATTR pts vs AL)	87%	Evaluation of diagnostic accuracy in the etiological diagnosis (identification of TTR pts)

AL, light chain amyloidosis; ATTRv, transthyretin amyloidosis variant; ATTRwt, transthyretin amyloidosis wild type; AF, atrial fibrillation; CMR, cardiac magnetic resonance imaging; DPD, 3,3'-diphosphono-1,2-propanediacetic acid; FPA, fat pad aspiration; IHC, immunohistochemistry; HMDP, hydroxymethylene diphosphonate; HCM, hypertrophic cardiomyopathy; HR, heart retention; H, heart; N, number; TTR, transthyretin; WBR, whole-body retention; WB, whole body.

Table 12. Key Literature Summarizing the Role of Radionuclide Imaging in Screening for Cardiac Amyloidosis in Selected Settings

First author	Year	Clinical setting	N Pts.	Iso- tope	Planar/ SPECT	Type of study and patient selection	N Pts. with CA	Preva- lence of CA	Crite- rion for CA	Reference diagnosis of CA	Com- ments
Longhi ²²⁸	2014	Bone-scan for non- cardiac reasons	12,400	DPD	Planar/ SPECT	Retrospective/patients with a positive bone scan were contacted for a cardiac evaluation	14 ATTR CA	1.4% in men in the 9th decade	Visual score ≥ 2	Echo + EMB + genotyping	
Gonzalez ¹³	2015	Hosp. pts HFpEF	120	DPD	Planar/ SPECT	Prospective-consecutive pts/ Pts with HFpEF and LVH (echo)	16 ATTRwt	13% (16/ 120)	Visual score ≥ 2	Genotyping + EMB or extra cardiac tissues	
Bennani Smires ¹⁵	2016	Hosp. pts HFpEF	49	DPD	Planar/ SPECT	Prospective-consecutive pts/Pts with HFpEF and no CAD	9 ATTRwt 5 AL 10% (5/49)	18% (9/49)	Visual score = 3; H/WB	CMR + genotyping + various organ biopsy (for AL diagnosis)	Selected pts underwent a cardiac screening
Longhi ¹¹⁷	2016	Severe AS evaluated for AVR	43	DPD	Planar/ SPECT	Prospective/ Pts with ≥ 1 of the following echo "red flags"	5 ATTRwt	12% (5/43)	Visual score ≥ 2	EMB + genotyping	2/5 low flow-low gradient, reduced LVEF 2/5 low flow-low gradient, preserved LVEF

Table 13. Key Literature Summarizing the Prognostic Value of ^{99m}Tc-DPD/HMDP Radionuclide Imaging for Cardiac Amyloidosis

First author	Year	N patients	Follow- up duration	Isotope	Planar/ SPECT	Type of study and patient population	ATTRv/ ATTRwt	Prognostic scintigraphic criterion	Prognostic role confirmed at multivariable analysis (Yes/No)
Hutt ³¹	2017	602	29.6 months	DPD	Planar/ SPECT	Retrospective/all pts who received a bone-scan to evaluate for cardiac amyloidosis	225 ATTRv; 377 ATTRwt	Visual score (N pts):	No
Univariable predictors: age; 6MWT; LVEF; troponinT; NT-proBNP; NYHA class; Echo performance status; GFR; SBP; Multivariable predictors: echo performance status; GFR; evaluation of the prognostic role of the scintigraphic visual score; survival was significantly longer in score 0 pts (absence of cardiac amyloidosis) compared to score 1-2-3 pts.									
Galat ¹⁰²	2015	121	111 [50-343] days	HMDP	Planar/ SPECT	Prospective/ Consecutive pts with suspected cardiac amyloidosis	55 ATTR (m+wt) cardiac amyloidosis	H/Skull	a.None b.NYHA III-IV c.NYHA III-IV
Comments: Cox's regression uni- and multivariate analysis were performed and prognosis was assessed in terms of predictors of MACE. a) model with scintigraphic + echo variables; (b) model with scintigraphic + echo + clinical variables; c) model with scintigraphic + echo + clinical + biological variables									
Univariable predictors: NYHA class; LVEF; E/e'; NT-proBNP; TroponinT									
Multivariable predictors: (a) none; (b) class NYHA III-IV; (c) class NYHA III-IV									
Kristen ¹⁷⁴	2013	36	27.4 [0.1- 106.2] months	DPD	Planar/ SPECT	Retrospective/consecutive pts with ATTRwt (20/36 underwent bone-scan)	ATTRwt	HR: Visual score	N/A
Comments: diastolic dysfunction; univariate Kaplan-Meier analysis was performed									
Rapezzi ¹⁰⁵	2011	63	14 [6.2-32] months	DPD	Planar/ SPECT	Retrospective/consecutive pts with ATTRv with/without cardiac amyloidosis	ATTRv	HR: H/WB	No
Univariable: age; left atrial diameter; LV wall thickness; LV M/V; NYHA class; low QRS; voltage; restrictive filling pattern; HR; H/WB									
Multivariable: age; restrictive filling pattern									
Cox's regression uni- and multivariate analysis were performed; prognosis was assessed in terms of predictors of MACE									

AA, amyloid A amyloidosis; AF disease, Anderson-Fabry disease; AL, light-chain amyloidosis; ATTR, transthyretin-related amyloidosis; ATTRv, amyloidosis due to transthyretin gene mutations; ATTRwt, amyloidosis due to deposition of wild-type transthyretin (senile amyloidosis); AV, atrio-ventricular; AVR, aortic valve replacement; CMP, cardiomyopathies; echo, echocardiogram; FPA, fat pad aspiration; GFR, glomerular filtration rate; HR, heart retention; H/WB, heart/whole body; HCM, hypertrophic cardiomyopathy; HFpEF, heart failure with preserved ejection fraction; IHC, immunohistochemistry; LVH, left ventricle hypertrophy; LVEF, left ventricular ejection fraction; LVM/V, left ventricle mass/volume ratio; mBMI, modified body mass index; MS, mass spectrometry; NYHA, New York Heart Association; MACE, major adverse cardiac events; RV, right ventricle; SBP, systolic blood pressure; TAVR, transcatheter aortic valve replacement; WBR, whole-body retention

Table 14. Key Literature Summarizing the Utility of ^{99m}Tc -PYP Radionuclide Imaging for Risk Assessment in Cardiac Amyloidosis

First author	Year	N patients	N controls	Design	Follow-up period	Outcome	Hazard ratio	Comments
Vranian ¹⁷⁵	2017	75	27	Retrospective	Not specified	Correlation to echo/ biomarkers	Not given	PYP predicted mortality in suspected ATTR, but not confirmed ATTR
Sperry ²³⁰	2017	54	0	Retrospective	Up to 4 years (median 1.8 years)	Regional PYP uptake by SPECT	0.73	
Castano ¹¹⁴	2016	121	16	Retrospective	5 years	Mortality	3.91	Multicenter registry

AS, aortic stenosis; ATTR, transthyretin amyloidosis; PYP, pyrophosphate; SPECT, single photon emission computed tomography

Table 15. Key Literature Summarizing the Diagnostic Value of ^{123}I -MIBG Radionuclide Imaging for Cardiac Amyloidosis

First author	Year	N patients	N controls	Planar/ SPECT	Patient cohort	ATTR/ AL	Criterion	Comments/Pt outcome
Tanaka et al ⁹⁸	1997	12	15	Planar/ SPECT	Prospective	ATTRv	Cardiac tracer accumulation	No cardiac tracer accumulation in 8 of 12 Mean FU 15.5 $\pm$ 5.8 months: no lethal arrhythmia, no cardiac death
Delahaye et al ¹³⁰	1999	17	12	Planar/ SPECT	Consecutive	ATTRv	Late HMR	Mean late HMR in patients 1.36 $\pm$ 0.26 vs in healthy controls 1.98 $\pm$ 0.35 ($p < 0.001$), no difference in wash-out
Delahaye et al ²³¹	2001	21	12	Planar/ PET	Consecutive	ATTRv	Muscarinic receptor density Late HMP	Mean muscarinic receptor density was higher in patients than in control subjects: B _{max} , 35.5 $\pm$ 8.9 vs 26.1 $\pm$ 6.7 pmol/mL ($p = 0.003$) Mean late HMR in patients 1.43 $\pm$ 0.28 vs in healthy controls 1.98 $\pm$ 0.35 ($p < 0.001$), mean wash-out 29 $\pm$ 6.8% vs 21 $\pm$ 6% ($p = 0.003$). Individual muscarinic receptor density did not correlate with late HMR
Watanabe et al ¹⁴¹	2001	4	10	Planar/ SPECT	Prospective	ATTRv	Late HMR	Mean late HMR in patients 1.1 $\pm$ 0.2, vs 2.4 $\pm$ 0.2 in healthy controls (p value N/A)
Hongo et al ¹³⁹	2002	25	12	Planar/ SPECT	Prospective	AL	Late HMR	Mean late HMR in patients without autonomic neuropathy 1.53 $\pm$ 0.06 vs in with autonomic neuropathy 1.29 $\pm$ 0.05 ($p < 0.001$), mean wash-out 42 $\pm$ 4.8% vs 31 $\pm$ 4.0% ($p < 0.001$)
Lekakis et al ¹⁴⁰	2003	3	23	Planar	Retrospective	AL	Late HMR	Mean late HMR 1.33 $\pm$ 0.1 vs in 2.13 $\pm$ 0.2 healthy controls (p value N/A)
Coutinho et al ²³²	2004	34	-	Planar	Prospective	ATTRv	Late HMR	Mean late HMR 1.75 $\pm$ 0.5 in all patients. Mean late HMR in patients without neuropathy 2.2 $\pm$ 0.5 vs patients with neuropathy 1.5 $\pm$ 0.4 ($p = 0.001$)
Delahaye et al ¹³⁸	2006	31	12	Planar/ SPECT	Prospective	ATTRv	Late HMR	Mean late HMR 6 months before liver transplantation 1.45 $\pm$ 0.29, vs 1.98 $\pm$ 0.35 of controls ($p < 0.001$) No cardiac death or lethal arrhythmia reported. Neuronal worsening in FAP patients after liver transplantation
Algalarrondo et al ²³³	2012	32	-	Planar	Retrospective	ATTRv	Late HMR	Late HMR $\leq$ 1.6 in 26 out of 32 patients No cardiac death or lethal arrhythmia reported
Noordzij et al ¹³⁵	2012	61	9	Planar	Consecutive	AL AA ATTR	Late HMR	Mean late HMR in all patients 2.3 $\pm$ 0.75 vs healthy control subjects 2.9 $\pm$ 0.58 ($p < 0.005$). Mean late HMR in ATTR patients 1.7 $\pm$ 0.75 vs AL patients 2.4 $\pm$ 0.75 ($p < 0.05$) Mean wash-out in patients 8.6 $\pm$ 14% vs in healthy control subjects - 2.1 $\pm$ 10% ($p < 0.05$)
Coutinho et al ¹²⁹	2013	143	-	Planar	Longitudinal consecutive	ATTRv	Late HMR	Mean late HMR 1.83 $\pm$ 0.43, and mean wash-out 47 $\pm$ 11% Mean FU 5.5 years: hazard ratio all-cause mortality 7 if HMR $<$ 1.6, progressive increase in 5-year mortality with decrease in late HMR
Takahashi et al ²³⁴	2014	6	-	Planar	Prospective	ATTRv	Late HMR	Mean late HMR at baseline 1.7 $\pm$ 0.9 ($p = 0.004$) No cardiac death or lethal arrhythmia reported
Algalarrondo et al ¹³¹	2016	215	-	Planar	Retrospective	ATTRv	Late HMR	Median late HMR 1.49 (inter-quartile range 1.24-1.74, range 0.97-2.52) Median FU 5.9 years after liver transplantation: 5-year survival 64% if late HMR $\leq$ 1.43, vs 93% if HMR $>$ 1.43 ($p < 0.0001$)
Azevedo Coutinho et al ¹⁸⁶	2017	232	-	Planar	Prospective	ATTRv	Late HMR	Initial assessment: mean late HMR 1.83 $\pm$ 0.03, median wash-out 2.5 (inter-quartile range - 2.3 to 8.5) Median FU 4.5 years (inter-quartile range 2.1-7.7 years) Initial HMR $<$ 1.55: HR mortality 9.36 (95% CI 4.27-20.56, $p < 0.001$) Initial HMR 1.55-1.83: HR mortality 4.27 (95% CI 1.68-9.05, $p = 0.002$).

AA, amyloid A; AL, amyloid light chain; ATTRv, hereditary transthyretin amyloidosis; FU, followup; HMR, heart-to-mediastinal ratio; HR, hazard ratio; Pt, patient

Table 16. Key Literature Summarizing the Utility of ^{123}I -MIBG Radionuclide Imaging for Risk Assessment in Cardiac Amyloidosis

First author	Year	N patients	N controls	Design	Follow-up period	Outcome	Event rate (annualized)	Hazard ratio (95% CI)	Comments
Azevedo Coutinho et al ¹⁸⁶	2017	232	-	Prospective observational	4.5 years	All-cause mortality	38 (11 71.5 patient years)	HR 0.183 (95% CI 0.075-0.450)	Late H/M multivariable predictor Liver transplantation patients (n = 70)
Algarrondo et al ¹³¹	2016	215	82 (no MIBG)	Consecutive	5.9 years	All-cause mortality	84 pts in total	HR 0.049 (95% CI 0.014-0.170)	Late H/M univariable predictor Liver transplantation patients (n = 65)
Coutinho et al ¹²⁹	2013	143	-	Prospective longitudinal consecutive	5.5 years	All-cause mortality	32 total (22.4%)	HR, 0.18 (95% CI 0.06-0.57)	Liver transplantation patients (n = 53)

CI, confidence interval; HR, hazard ratio; H/M, heart to mediastinal ratio

Table 17. Recommendations for Standardized Acquisition of ^{123}I -MIBG for Cardiac Amyloidosis

Imaging procedures	Parameters	Recommendation
Preparation	No fasting required Withdrawal of certain drugs: readers are referred to Ref. 235 Oral thyroid blockage 30 min before administration ^{123}I -MIBG: Lugol solution (130 mg adults, body weight adjusted children) OR potassium perchlorate (500 mg adults, body weight adjusted children)	Preferred
Scan	Rest scan	Preferred
Dose of ^{123}I -MIBG	370 MBq (10 mCi) intravenously	Preferred
Time between injection and acquisition	15 minutes 3-4 hours	Preferred Preferred
General imaging parameters		
Field of view	Heart Chest	Preferred Preferred
Image type	SPECT	Preferred
Position	Planar Supine	Preferred Standard
Energy window	Upright	Optional
Collimators	159 keV, 15-20% Medium energy, high resolution	Standard Preferred
Matrix	Low energy, high resolution	Optional
Pixel size	128 × 128 (maximum 256 × 256) 4.5-6.4 mm	Standard Standard
Planar imaging specific parameters		
Number of views*	Anterior	Standard
Detector configuration	Planar	Standard
Image duration (count based)	10 min	Standard
Magnification	1.45	Standard
SPECT imaging specific parameters		
Angular range	180°	Standard
Detector configuration	90°	Standard
Angular range	360°	Optional
Detector configuration	180°	Optional
ECG gating	Off; Nongated imaging	Standard
Number of views/detector	64 over 180°	Standard
Time per stop	20 seconds	Standard
Magnification	1.0	Standard

For details on ^{123}I -MIBG imaging, readers are referred to Ref. 235

Table 18. Recommendations for ^{123}I -MIBG Imaging Reporting

Parameters	Elements
Demographics	Patient name, age, sex, reason for the test, date of study, prior imaging procedures, biopsy results if available (required)
Methods	Imaging technique, radiotracer dose and mode of administration, interval between injection and scan, scan technique (planar and SPECT) (required)
Findings	Image quality Thyroid uptake Visual scan interpretation (required) Semi-quantitative interpretation heart-to-mediastinum (H/M) (required): Regions of interest (ROIs) are drawn over the heart (including the cavity) and the upper mediastinum (avoiding the thyroid gland) in the planar anterior view. Average counts per pixel (CPP) in the myocardium are divided by average counts per pixel in the mediastinum. The myocardial washout rate (WR) from initial to late images is also calculated, and expressed as a percentage, as the rate of decrease in myocardial counts over time between early and late imaging (normalized to mediastinal activity)
Ancillary findings	On planar imaging and SPECT (optional) Interpret CT for attenuation correction if SPECT/CT scanners are used (recommended)
Conclusions	The appearance of images should be described succinctly, including a statement on quality if suboptimal. Sympathetic activity SPECT defects should be classified in terms of location relative to myocardial walls, extent and severity. Other abnormalities that should be mentioned are LV dilatation, increased lung uptake of tracer, or significant noncardiopulmonary tracer uptake. The findings should be integrated with the clinical data to reach a final interpretation. A comparison with any previous study should be included Normal values for late H/M ratio and WR vary in relation to age (inversely for the late H/M ratio, directly for the WR) and image acquisition (LEHR vs ME collimation and acquisition time) In general: A WR > 20% between early and late imaging is considered as abnormal A late H/M ratio < 1.60 is abnormal, between 1.60 and 1.85 equivocal, and > 1.85 as normal (LEHR collimator)

H, heart; *LEHR*, low energy high resolution; *M*, mediastinum; *ME*, medium energy; *LV*, left ventricular; *SPECT*, single photon emission computed tomography; *WR*, wash out rate

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