AL Amyloidosis for Cardiologists
Awareness, Diagnosis, and Future Prospects:
JACC: CardioOncology State-of-the-Art Review

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ABSTRACT

Amyloid light chain (AL) amyloidosis is a rare, debilitating, often fatal disease. Symptoms of cardiomyopathy are common presenting features, and patients often are referred to cardiologists. Cardiac amyloid infiltration is the leading predictor of death. However, the variable presentation and perceived rarity of the disease frequently lead to delay in suspecting amyloidosis as a cause of heart failure, leading to misdiagnoses and a marked delay in diagnosis, with devastating consequences for the patient. A median time from symptom onset to correct diagnosis of about 2 years is often too long when median survival from diagnosis for patients with AL amyloidosis and cardiomyopathy is 4 months to 2 years. The authors highlight the challenges to diagnosis, identify gaps in the current knowledge, and summarize novel treatments on the horizon to raise awareness about the critical need for early recognition of symptoms and diagnosis of AL amyloidosis aimed at accelerating treatment and improving outcomes for patients. (J Am Coll Cardiol CardioOnc 2022;4:427–441)

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Cardiac amyloid infiltration is the leading cause of death, accounting for >61% of fatalities and the primary determinant of survival in patients with AL amyloidosis.\textsuperscript{13,19} Cardiovascular events leading to death include progressive heart failure or arrhythmias, although the exact cause is often unclear.\textsuperscript{19} Common cardiovascular complications include atrial or ventricular (less common, but often fatal) arrhythmia, heart failure, embolism, stroke, and conduction delays, including advanced degrees of atrioventricular block. Patients with cardiac involvement experience the greatest impact on all aspects of quality of life (QoL), including physical health, work productivity, and emotional health, which includes depression and role limitation, compared with patients with involvement of other organs.\textsuperscript{20,21}

Cardiac amyloidosis is often listed as a cause of heart failure with preserved ejection fraction, but this does not fully characterize the functional phenotype that typically shows both systolic and diastolic impairment.\textsuperscript{13-15} Patients with AL amyloidosis, amyloid infiltration results in symmetrical biventricular wall thickening with nondilated or small ventricles. Ejection fraction is typically normal until late disease stage.\textsuperscript{22} The reduction in systolic function disproportionally affects the longitudinal rather than the radial axes.\textsuperscript{22} This is much more marked than other causes of thick-walled heart failure, and the reduction in longitudinal strain typically spares the apex, giving the characteristic relatively apical sparing picture (“bull’s-eye pattern”) on parametric longitudinal strain polar maps.\textsuperscript{23} Initial infiltration in the left ventricle is characterized by impaired relaxation, which then invariably progresses to typical restrictive physiology. Nonspecific extensively described findings include thickening of the valves and the interatrial septum and a speckled appearance of the myocardium. Pericardial and pleural effusions are relatively common findings.\textsuperscript{22}

AL amyloidosis is difficult to diagnose because many of the presenting symptoms are not specific to this disease.\textsuperscript{13,15,24,25} Delays in diagnosis and misdiagnoses are common and result in increased symptom burden, cumulative organ damage, poorer prognosis, and higher early mortality.\textsuperscript{25-28} Rapid accurate diagnosis within 6 months of symptom onset correlated positively with survival, whereas diagnostic delay (>6 months) correlated with poor survival.\textsuperscript{20} Worsening organ damage due to diagnostic delay limits treatment options, such as making patients ineligible for autologous stem cell transplantation (ASCT) or having poorer tolerance to standard-of-care chemotherapy. As many patients present with cardiovascular symptoms or complications, they are often referred to cardiologists by their primary care physicians.\textsuperscript{26,29} A survey of patients and caregivers indicated that about 44% of patients initially were misdiagnosed and treated as having unrelated cardiomyopathy, with 3 of 4 cardiologists surveyed reporting that lack of disease awareness was the most common cause of misdiagnosis.\textsuperscript{26} Delays in diagnosis can increase the risk for death 3- to 5-fold.\textsuperscript{20} Thus, it is critical to increase awareness among health care providers to ensure timely diagnosis and improved outcomes.

Previously published state-of-the art reviews focused on the utility of supportive therapy for cardiac involvement, chemotherapy, or immunotherapy.\textsuperscript{23,24} This review, focusing on cardiac aspects of AL amyloidosis, builds on that foundation, with the aim of improving awareness about suspicion, diagnosis, gaps in treatment and response evaluations, and recent advances in the management of this disease (Central Illustration).

### Diagnosis

Most of the symptoms and clinical manifestations of AL amyloidosis are nonspecific.\textsuperscript{33,34} Diagnosis of AL amyloidosis is a multistep process (Figure 2).\textsuperscript{25,35} The first step is suspicion of amyloidosis, which initiates specific diagnostic tests to verify the diagnosis.\textsuperscript{25,35} This is followed by tests to identify the amyloid fibril type, which is critical to differentiate AL amyloidosis from other subtypes.\textsuperscript{25,35} The final step of diagnosis is to identify the organs involved and the extent to which they are affected by the deposition of
amyloid fibrils. This forms the basis of accurate diagnosis and a risk-adapted treatment plan for patients with AL amyloidosis.

**STEP 1: SUSPICION OF AMYLOIDOSIS.** Suspicion of amyloidosis should be very high when a patient presents with heart failure combined with a constellation of unexplained extracardiac symptoms such as neuropathy, bleeding, carpal tunnel syndrome, nephrotic syndrome, proteinuria, diarrhea, hepatomegaly, peripheral and autonomic neuropathy, macroglossia, and periorbital purpura. Another key cardiovascular presentation that should trigger suspicion of amyloidosis is an increase in left ventricular mass or wall thickness in the absence of hypertension or the presence of hypotension, especially when associated with increased right ventricular mass. Cardiovascular findings that suggest amyloidosis include restrictive cardiomyopathy with disproportionately impaired longitudinal function compared with radial contraction (apical sparing on longitudinal strain imaging) and low-voltage electrocardiogram or pseudoinfarct pattern. Elevated serum cardiac troponin T and/or N-terminal pro-brain natriuretic peptide (NT-proBNP) are common in patients with cardiac AL amyloidosis. Although macroglossia and periorbital purpura are considered pathognomonic for AL amyloidosis, they occur in only about 15% of patients. Although no single symptom is
This figure summarizes (A) the current treatment paradigm, (B) the gaps that still exist in treatment and in relevant endpoints, and (C) emerging therapies that target removal of existing amyloid fibril deposits from organs and tissues. AL = amyloid light chain; ASCT = autologous stem cell transplantation; CR = complete response; IgG = immunoglobulin G; PCD = plasma cell dyscrasia; QoL = quality of life.
Specific for AL amyloidosis, it should be suspected when there is a combination of cardiomyopathy and/or other aforementioned nonspecific symptoms.

**STEP 2: DIAGNOSIS OF AMYLOIDOSIS.** Once amyloidosis is suspected, there are several tests that can be undertaken to verify the diagnosis, detect the presence of an underlying PCD, and determine which organs are involved (Figure 3). Free light chain (FLC) assays can detect abnormal levels of Ig light chain in serum. Serum and urine immunofixation electrophoresis assays are used to quantify and determine the type of abnormal Ig light chains or monoclonal protein. Positive results in these assays would indicate the presence of a monoclonal gammopathy of undetermined significance, multiple myeloma, indolent B cell lymphoma, Waldenström macroglobulinemia, or AL amyloidosis. The combination of these assays can be very effective in detecting amyloidogenic light chain in most patients with AL amyloidosis. The absence of a detectable monoclonal protein in the serum and urine by immunofixation electrophoresis and serum FLC assays makes AL amyloidosis less likely. Congo red staining of a tissue biopsy sample is required for confirmation of the diagnosis. Abdominal fat pad biopsies are a fast and safe option to detect amyloid deposits. Although the presence of amyloid deposits by Congo red staining in fat pad confirms the diagnosis of amyloidosis, their absence does not rule out disease. Bone marrow examination is an essential component of AL amyloidosis diagnosis. Combination of abdominal fat and bone marrow biopsies will identify 85% of patients with AL amyloidosis. In case of a negative biopsy result, alternative sites (labial salivary glands, gut) or a biopsy of the affected organ (eg, heart, kidney) should be considered. There are structural and functional differences between cardiac AL amyloidosis and other hypertrophic cardiomyopathies, but significant overlap exists. Echocardiography can provide an assessment of the likelihood of cardiac amyloid infiltration vs other hypertrophic cardiomyopathies and can assess the severity of cardiac involvement. Although echocardiography is useful in patients with proven amyloidosis for both diagnosis and monitoring of heart involvement, the overlap of echocardiographic features among multiple diseases makes echocardiography poorly suited to rule out the diagnosis definitively. Cardiac magnetic resonance imaging (CMR) offers accurate information regarding the heart’s structure and function with precision...
advantages over echocardiography; the key is its unique ability to give information about the tissue composition by “myocardial tissue characterization,” making it a highly sensitive and specific imaging modality to distinguish cardiac amyloidosis from other hypertrophic phenocopies.\textsuperscript{38,50-59} CMR with late gadolinium enhancement (LGE) can visualize the continuum of cardiac infiltration, from subendocardial LGE to increasing transmurality as the disease progresses.\textsuperscript{52,60,61} When combined with T1 mapping, CMR with LGE can be used to characterize and measure the degree of cardiac amyloid infiltration with 70% to 80% accuracy in patients with known systemic AL amyloidosis.\textsuperscript{62-69} It should be mentioned that although CMR is sensitive for distinguishing AL from other cardiomyopathies, it may not be suggestive in patients with early involvement of the heart by AL amyloidosis (ie, the absence of LGE on CMR does not rule out early cardiac involvement). Extracellular volume mapping should be considered in the CMR protocol to exclude early cardiac amyloid infiltration.

STEP 3: TYPING AND CONFIRMING DIAGNOSIS OF AL AMYLOIDOSIS. It is critical to identify the type of amyloid fibril to avoid misdiagnosis and initiation of incorrect or inappropriate treatment, which could have disastrous consequences for a patient. This has become even more important with increasing recognition of cardiac amyloidosis in older patients (differential diagnosis between wild-type transthyretin [ATTR] and AL amyloidosis). The most common methods of amyloid fibril typing include immunohistochemistry or laser capture, followed by mass
spectrometry to identify the amyloid subtype. As the accuracy of immunohistochemistry is dependent on the expertise of the laboratory and needs an extensive panel of antibodies for accurate reporting, laser capture with mass spectrometry has become the method of choice for amyloid fibril typing.

Biopsy has been an essential step in diagnosis of amyloidosis until recent advances, showing accurate diagnosis of cardiac ATTR amyloidosis using 99mTc-labeled bone scintigraphy tracers, a noninvasive procedure. Specifically, in the absence of a detectable monoclonal protein in blood or urine by immunofixation and normal serum FLCs, grade 2 or greater uptake on 99mTc-pyrophosphate or 99mTc-3-diphosphono-1,2-propanodicarboxylic acid scintigraphy is considered diagnostic of cardiac ATTR amyloidosis. Conversely, among patients who present with abnormal results on serum protein electrophoresis or FLC assays, no cardiac uptake on 99mTc-pyrophosphate or 99mTc-3-diphosphono-1,2-propanodicarboxylic acid scan can only exclude cardiac ATTR amyloidosis but crucially does not rule out cardiac AL amyloidosis. Indeed, in AL amyloidosis, a full nonbiopsy path to diagnosis does not exist.

For cardiac AL amyloidosis, different centers may have different approaches as it pertains to cardiac biopsy, given the associated risks of cardiac biopsy and that noninvasive cardiac imaging confirming cardiac involvement along with Congo red staining and typing of a noncardiac biopsy can diagnose AL amyloidosis with high sensitivity. However, if there is high clinical suspicion of cardiac AL amyloidosis, cardiac biopsy should be performed if biopsy of noncardiac sites is negative.

**CURRENT STANDARD OF CARE**

The current standard of care is based on observations of improved overall survival and organ function when synthesis of amyloidogenic light chain is suppressed or stopped, along with supportive care to address specific organ dysfunction. The response to treatment requires dual assessment: 1) primary assessment of hematologic response of the underlying clone to chemotherapy, which can manifest within a few weeks of treatment initiation; and 2) assessment of change in organ function for organ response or progression. Improved organ function is usually delayed by several months (median 10.4 months), and only 25% to 50% of all patients treated with current chemotherapy regimens achieve an organ response. The criteria for evaluating hematologic and organ responses are given in Table 1.

At diagnosis, patients are first evaluated to determine whether they are eligible to receive ASCT (~20% of patients) or will be receiving combination chemotherapy (~80% of patients) (Figure 3). The criteria for ASCT are very strict to minimize treatment-related mortality, and many patients opt to not undergo the procedure even when eligible. Data suggest better outcomes for patients who undergo induction chemotherapy prior to ASCT, with the
caveat that a small proportion of patients may become ASCT ineligible during induction treatment. Among patients receiving ASCT as first-line therapy to treat AL amyloidosis, 73% exhibited complete response (CR) or very good partial response (VGPR) as determined by hematologic response criteria (Figure 4). Patients not eligible for ASCT (≈80%) and those who have opted not to receive it are treated with one of several options of anti-PCD therapy. Current guidelines recommend a combination of cyclophosphamide, bortezomib, and dexamethasone (CyBorD) and daratumumab as the preferred first-line anti-PCD therapy, with CyBorD and bortezomib-melphalan-dexamethasone as alternatives when there is limited access to daratumumab. Efficacy of the most commonly used first-line treatments with respect to hematologic response are shown in Figure 4. The hematologic efficacy of daratumumab-CyBorD is very high, with 78% of patients achieving VGPR or better. More than 20% of patients discontinue at least 1 therapy, and >30% of patients reduce treatments because of adverse events. Patients who present with significant neuropathy can be treated with regimens that avoid bortezomib and use daratumumab monotherapy. Ixazomib may be useful in patients with relapsed disease who have not been previously exposed to a proteasome inhibitor.

Orthostatic hypotension, a highly debilitating symptom experienced by patients with AL amyloidosis as a consequence of autonomic neuropathy, can be managed by adjunct supportive treatment,
including wearing compression garments and medications such as midodrine or droxidopa.\textsuperscript{2,12,97} Although fludrocortisone may help some patients, it often causes fluid retention and is often not well tolerated.\textsuperscript{2}

\section*{ANTIFIBRIL TREATMENTS IN DEVELOPMENT}

At the present time, 2 antibodies, birtamimab and CAEL-101, are being investigated as antifibril agents.\textsuperscript{29-107} Dezamizumab, another such agent, is no longer being investigated. Antifibril antibodies have the potential to remove existing amyloid fibrils from organs by activating immune cells to cause chemical and enzymatic degradation of existing amyloid fibrils and inducing antibody-dependent phagocytosis of the fibrils.\textsuperscript{107,108} In addition, although this utility has not yet been adequately investigated, antifibril antibodies have the potential to be used as diagnostic agents to image amyloid deposits and for amyloid typing on tissue biopsies.

\textbf{BIRTAMIMAB.} Birtamimab (NEOD0001) is a fully humanized monoclonal antibody that targets a cryptic epitope on serum amyloid A protein that is revealed when misfolded.\textsuperscript{109} It cross-reacts with Ig light chain amyloid fibrils and reportedly activates macrophage-mediated degradation and clearance of light chain fibrils. A first-in-human phase 1/2 study (NCT01707264) of intravenous birtamimab given every 4 weeks was conducted in 27 patients who had experienced at least a partial hematologic response to prior light chain-suppressive chemotherapy but had persistent organ dysfunction. The treatment was well tolerated, with no dose-limiting toxicities or discontinuations due to drug-related adverse events. Organ responses were observed in 57% of cardiac-evaluable patients and 60% of renal-evaluable patients.\textsuperscript{110} Birtamimab, given as an intravenous infusion every 28 days, was well tolerated up to 24 mg/kg and demonstrated cardiac or renal response in the majority of patients, with the remainder having stable disease.\textsuperscript{110} In some cases, organ response was delayed by several months relative to hematologic response.\textsuperscript{111} On the basis of these promising results, the phase 3 VITAL study (NCT02312206) randomized 260 patients with newly diagnosed AL amyloidosis with cardiac involvement to receive standard-of-care chemotherapy plus either birtamimab or placebo. A futility analysis performed after 103 primary endpoint events (time to all-cause mortality or cardiac hospitalization) showed no advantage for the birtamimab arm, and the study was therefore terminated. However, a post hoc analysis identified a benefit for both primary endpoints in patients with Mayo stage IV cardiac disease, with median survival of 8.3 months in the placebo group compared with not reached (>12 months) in the birtamimab group, leading to the initiation of AFFIRM-AL (A Study to Evaluate the Efficacy and Safety of Birtamimab in Mayo Stage IV Patients With AL Amyloidosis; NCT04973137), a new phase 3 trial in patients with advanced cardiac disease.

\textbf{CAEL-101.} CAEL-101 is a chimeric monoclonal antibody that targets a cryptic epitope on Ig light chains that is exposed when the light chains are misfolded.\textsuperscript{107,112} It binds to misfolded free Ig light chains as well as amyloid fibrils deposited in organs.\textsuperscript{107,113,114} It is hypothesized that CAEL-101 opsonizes the amyloid fibrils and misfolded light chains, thereby attracting and activating macrophages that degrade the complex by phagocytosis and/or enzymatic and chemical proteolysis.\textsuperscript{107} Preclinical studies further demonstrated that treatment with CAEL-101 resulted in regression of transplanted human amyloidomas in a mouse model.\textsuperscript{113,115,116} Phase 1 studies with CAEL-101 (NCT00807872 and NCT02245867) demonstrated that it recognized amyloid fibrils deposited in tissues. Furthermore, CAEL-101 had a dose-proportional pharmacokinetic profile without any dose-limiting toxicities.\textsuperscript{114,117,118} Preliminary results also showed that there were reductions in biomarkers of cardiomyopathy and nephropathy.\textsuperscript{117,118} The phase 2 study (NCT04304144) confirmed that there were no dose-limiting toxicities up to 1,000 mg/m\textsuperscript{2} and that it could be administered with CyBorD or CyBorD plus daratumumab.\textsuperscript{100,101} Furthermore, the exposure to CAEL-101 was not affected by adding daratumumab to the anti-PCD treatment regimen. Initial evaluations also confirmed the improvements in biomarkers of cardiomyopathy and nephropathy.\textsuperscript{100} On the basis of these promising results, 2 randomized, double-blind phase 3 studies are currently enrolling patients at more than 70 sites in several countries to evaluate the efficacy and safety of coadministering CAEL-101 with standard-of-care anti-PCD therapy in patients with AL amyloidosis and severe cardiomyopathy in stages IIIa (NCT04512235) and IIIb (NCT04504825). These are the first time-to-event studies focusing exclusively on patients with severe cardiomyopathy.

\section*{GAPS}

\textbf{TREATMENT GAPS.} At present, although treatment guidelines exist, there is no consensus on which regimen should be used as first-line treatment.\textsuperscript{87} The anti-PCD treatment recommended in current National Comprehensive Cancer Network guidelines is CyBorD with daratumumab.\textsuperscript{87} The components of this
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or refractory AL amyloidosis.129,130 However, this and pomalidomide, usually in combination with treatment gap. 

function, especially that of the heart, remains a huge existing amyloid deposits to repair and/or restore organ overall survival.121 However, the reasons for this occurred prior to treatment initiation or the damage that can continue to be caused by these preexisting deposits.24,38,124,127,128 Reducing or eliminating amyloid burden by partial or complete removal of preexisting amyloid deposits to repair and/or restore organ function, especially that of the heart, remains a huge treatment gap.

Immunomodulatory agents, such as lenalidomide and pomalidomide, usually in combination with dexamethasone, are commonly used to treat relapsed or refractory AL amyloidosis.129,130 However, this combination has a low hematologic response rate and can result in hematological toxicity, increased NT-proBNP, and sometimes nephrotoxicity.129,130 Most of the current chemotherapy regimens include dexamethasone (or other corticosteroids) as a key component. However, it is poorly tolerated, often causing worsening of heart failure, fatigue, increased skin fragility, and risk for sudden death in patients with advanced cardiac AL amyloidosis.129,120,130 Careful monitoring for complications and using the lowest doses possible are crucial. Results from a retrospective analysis show that venetoclax may be effective among patients with AL amyloidosis with a t(11:14) translocation and refractory to standard anti-PCD treatment, suggesting that genetics may play a role in response.131 Thus, there is a need to better understand resistance to treatment and more effective and safer treatment options for patients with relapsed or refractory AL amyloidosis. This has become especially important with use of daratumumab as a frontline agent in treatment of AL amyloidosis.

Because AL amyloidosis is very heterogeneous in its presentation, supportive care to help manage the many symptoms experienced by patients is critical.132 However, the heterogeneous nature of AL amyloidosis also makes standardization of supportive care very difficult. At present, there are no consensus standards for supportive care. Diuretic agents, β-blockers, and angiotensin-converting enzyme inhibitors are the principal drugs used to prevent cardiac failure among patients with nonamyloid cardiomyopathy.7,135,136 However, in patients with AL amyloidosis with advanced cardiomyopathy, hypotension is common, and β-blockers and angiotensin-converting enzyme inhibitors are poorly tolerated and can be dangerous.7,135,136 These drugs should not be used for this group of patients except in very particular circumstances, determined on an individual case basis.133 Bradyarrhythmia may require implantation of pacemakers, but the role of cardiac resynchronization devices remains unclear. Hypotension can also severely limit hemodialysis, which is essential for patients with renal involvement.132,133 Management of autonomic neuropathy remains challenging. In addition, there is a huge unmet need for supportive care among patients with comorbid gastrointestinal symptoms, for whom the current drugs and dietary modifications are not sufficient.132

GAPS IN RELEVANT ENDPOINTS. The primary endpoint preferred by regulatory agencies is overall survival.134 However, as many patients, especially those with minimal or no cardiomyopathy, can live for 5 years or longer after diagnosis, determining median overall survival requires very long trials, which proves difficult with a rare disease because of the paucity of patients available and delaying trial readouts. Better understanding from regulators is needed to accept surrogate endpoints such as the established criteria for hematologic response and NT-proBNP for cardiac response.134 However, outcomes that reduce comorbidities, reduce amyloid burden, improve organ function, or improve patient QoL are less likely to be accepted as primary endpoints for trials, even if they are clinically relevant. It is
essential that this gap be addressed, and preferably eliminated, to ensure that the best treatments are available to patients.

The goal for hematologic response has been for patients to achieve VGPR or better (Table 1).16,77,79 Recently, the goal changed to achieving CR because it correlated better with improved patient survival and QoL. However, the current criteria for CR do not include measurement of the difference between involved and uninvolved FLC (<10 mg/dL) or involved FLC (<20 mg/dL) in the serum, both of which are significant predictors of survival and organ response.121 There also is no consensus on the timing of hematologic response assessment. Hematologic response is variously reported 1, 3, or 6 months after treatment initiation, making it difficult to objectively evaluate the relative benefits of different treatment regimens as also is the lack of validated hematologic progression criteria. Time to retreatment often is used as a surrogate endpoint. These limitations in current hematologic response and progression criteria highlight the need for better means to measure initial and longitudinal response to treatment.

Organ response is typically determined only for a few organs, mainly the heart, kidneys, and liver, as these are considered the most relevant for survival.16,122 There are no established criteria for response to treatment for other organs, such as the gastrointestinal system, or for the neuropathy often experienced by patients. Although the hematologic and cardiac response criteria may reflect the level of cardiomyopathy, they do not inform the patients or those treating them about the disease state of other organs and tissues. Furthermore, neither the hematologic nor the organ response criteria measure the amyloid burden remaining in the body and continuing to have an adverse impact. Typically, organ response is observed much later than hematologic response, sometimes being delayed by months.79,135 Although achieving organ response within 1 year is predictive of longer survival, there are no standardized criteria on when organ response should be reported.79

Functional changes are often measured in clinical trials using the 6-minute walk test and various QoL instruments.21,136-138 The 6-minute walk test is a simple method to determine improvements in physical function and requires only an even flat surface at least 30 m in length and a timer.136 It is an easy test to implement and can reflect a patient’s level of cardiac impairment and the ability to perform an activity of daily living.136 The most commonly used instruments to measure QoL among patients with AL amyloidosis are the Short Form-36 (SF-36) and the Kansas City Cardiomyopathy Questionnaire.21,68,137-139 Both are measures of patient-reported outcomes. While the Short Form 36 is a robust, generic instrument that provides a reasonable snapshot of the physical and mental health of the patient, the Kansas City Cardiomyopathy Questionnaire is more disease specific and provides an estimation of the health of the heart of the patient at the time the test is administered. Both instruments have been validated for use with patients with AL amyloidosis.21,137-139 Although it is understandable that QoL instruments are focused on general and cardiac health because survival is most closely associated with degree of cardiomyopathy, there are gaps in understanding the health of other organs or systems from a patient’s perspective, despite instruments’ being available to determine the degree of nephropathy or hepatopathy.140,141

Structural improvements in the heart and decrease in myocardial amyloid deposits can be determined with CMR.50 Functional improvements of the heart can be determined with echocardiography and global longitudinal strain measurements, which can help with predicting survival.142-145 There are also several studies under way to validate positron emission tomography-based scintigraphy as a relatively noninvasive method to determine health of and the amount of amyloid deposited in the heart.75,106,146-151

Given the spectrum of endpoints used to evaluate response to treatment, there is no consensus on what defines a clinically meaningful response that translates to patient benefit in survival and QoL. Thus, there is a lack of consensus on what defines disease progression and, consequently, how to define progression-free survival.

It must be noted that the endpoints measured and the criteria for response to treatment were developed exclusively on the basis of the effectiveness of anti-PCD therapy, including ASCT. There are no generally accepted endpoints for reduction in amyloid burden or functional changes in organs other than the heart. Thus, there is a huge informational gap that needs to be addressed.

CONCLUSIONS

AL amyloidosis is a very devastating disease with continuing high early mortality. A significant majority of patients present with cardiomyopathy and are usually referred to cardiologists for diagnosis. However, for a substantial number of these patients, accurate diagnosis is either delayed or completely missed. These delays lead to poor prognosis and survival. It is therefore critical to increase awareness, knowledge, and understanding of AL amyloidosis
among all health care providers, cardiologists in particular. In this review, we have attempted to provide a concise overview of this very complex rare disease. Furthermore, we have summarized the promising novel interventions under investigation to address removal of amyloid fibrils already deposited in the organs, an important treatment gap that currently exists. This review is but one step toward improving the diagnostic and treatment odysseys experienced by patients with AL amyloidosis.

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