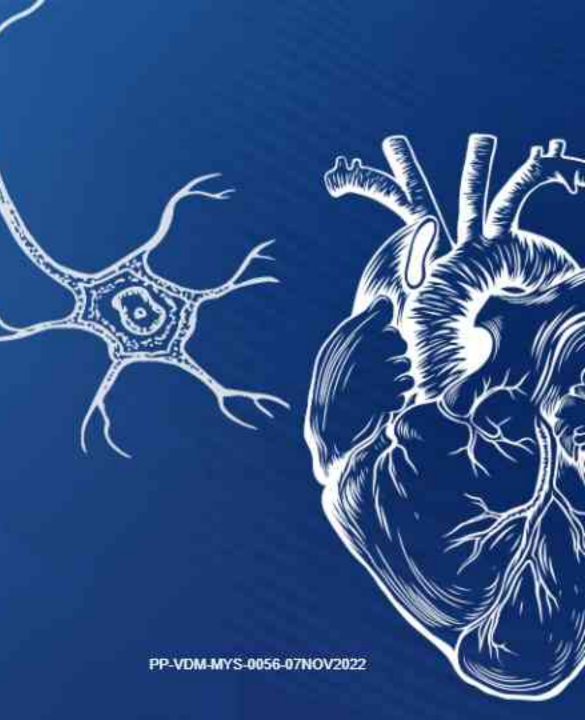


# Ruling out Light-Chain (AL) amyloidosis in ATTR-CM

Transthyretin amyloid cardiomyopathy (ATTR-CM)





# ATTR-CM and Light Chain (AL) amyloidosis have similar symptom presentation which may make diagnosis challenging

It is important to clinically differentiate between cardiac manifestations of ATTR and AL amyloidosis, as they have different clinical courses and treatment options, and AL amyloidosis is considered a hematologic urgency 1,2

# ATTR-CM and AL amyloidosis (CA) 95% ATTR-CM and AL amyloidosis account for 95% of all CA diagnoses Other types of CA ATTR-CM is under recognized, may present with symptoms similar to AL and may be misdiagnosed due to incomplete testing<sup>4,5</sup>

### Symptoms that may present in both ATTR-CM and AL amyloidosis

#### Cardiovascular<sup>3,6-8</sup>

- Heart failure with potential intolerance to standard therapy
- Cardiac arrhythmia
- Aortic stenosis
- Low voltage relative to left ventricular (LV) mass
- Echocardiography showing increased LV wall thickness

Ocular4,11,12

#### Nervous System4,8-10

- Automatic neuropathy
- Orthostatic hypotension
- Peripheral sensory motor dysfunction
- · Peripheral neuropathy
- · Unexplained weight loss
- Sexual impotence

#### Musculoskeletal/Orthopedic<sup>4</sup>

Carpal tunnel syndrome

#### Renal10,13,14

Renal impairment

Periorbital purpura\*

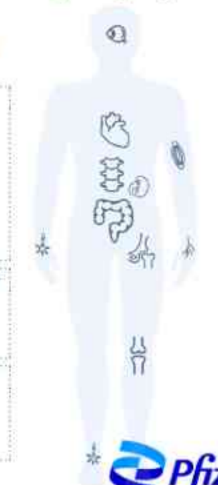
· Vitreous opacity

Glaucoma

- Nephrotic syndrome\*
- Cardiorenal symptoms

#### Other 4,10,15

- · Gastrointestinal complaints
- Macroglossia (large tongue)\*
- Nail dystrophy\*



ore commonly seen in AL

## Ruling out light chain amyloidosis (AL) is a key step in achieving a definitive ATTR-CM diagnosis 16

#### SERUM AND URINE TESTS FOR RULING OUT AL AMYLOIDOSIS\*

	Serum protein electrophoresis (SPEP) with immunofixation†	Urine protein electrophoresis (UPEP) with immunofixation <sup>†</sup>	Serum free light chain assay
What does it detect?	Clonal immunoglobulin and/or clonal light chain	Clonal immunoglobulin and/or clonal light chain	Free kappa chain Free lambda chain kappa: lambda ratio
Most sensitive test for	Confirming clonal immunoglobulin production	Confirming clonal light chain production	Detecting low-level clonal light chain production; clonality assumed if ratio is far from 1:1
Normal range	No M-spike present	No M-spike present No proteinura	Kappa lambda ratio=0,26- 1,65 <sup>‡</sup>

- AL amyloidosis is a hematological urgency and requires immediate treatment
- Survival of untreated patients with AL amyloidosis with cardiac involvement may be <6 months</li>
- Cardiac localization of radiotracer by bone scintigraphy can occur in approximately 30% of patients with AL amyloidosis
- Specific disease-modifying therapies for AL amyloidosis are available



<sup>&</sup>quot;If any of these tests are abnormal, bone scintigraphy should not be used to make the diagnosis of transthyrefin amyloldosis, and a blopsy is recommended. Adapted from Witteles et al., 2019 Table 1, which uses SPIE (serum protein electrophoresis with immunofixation) and UPIE (unine protein electrophoresis with immunofixation).

<sup>+</sup>SPEP and UPEP are more sensitive than protein electrophorests without immunofitiation and should be ordered as preferred test.

In patients with kinney disease, mild elevations in the kappa lambda ratio are frequently encountered, in the setting of a normal SPEP/UPEP, a kappa lambda ratio up to 2.5 can typically be considered normal.

AL rule out is an important step in the diagnostic pathway. All three tests should be sent to evaluate for AL in order to maximize sensitivity and specificity<sup>17</sup>

Order Studies to Test for Abnormal Monoclonal Protein\*

Serum Kappa/Lambda Free Light Chain Ratio

Abnormal result:

Ratio < 0.26 or > 1.65

Serum Protein Immunofixation
Abnormal result:
Monoclonal protein is detected

Urine Protein Immunofixation
Abnormal result:
Monoclonal protein is detected

#### Key Considerations:

- It is recommended that all three tests be sent to evaluate for AL amyloidosis. However, monoclonal protein presence does not rule out the
  possibility of ATTR-CM
- Amyloid subtyping is the only way to effectively differentiate between ATTR-CM and AL Amyloidosis

Adapted and reprinted with permission from Maurer MS, et al. Expert consensus recommendations for the suspicion and diagnosis of transitiyretin cardiac amyloidosis. Circ Heart Fall. 2019;12:e205075. doi:10.1161/ CIRCHEARTFAILURE 119.006075 © 2019 American Heart Association, inc. All rights reserved.

"If any of these tests are abnormal, bone scintigraphy should not be used to make the diagnosis of transthyretin amyloidosis, and a bloopy is recommended.

Immunofixation of the serum and urine is a much more sensitive test and usually demonstrates a monoclonal band. Thus, immunofixation and serum-free light chains should always be measured when amyloidosis is suspected. "Turine protein electrophoresis with immunofixation can be performed on soot or 24-h urine collection."





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