

AL-Amyloidosis Quick Reference Guide

Disease Overview

AL-Amyloidosis (Light Chain) is a rare but life-threatening disease caused by abnormal plasma cells producing misfolded light chain proteins. These proteins form amyloid fibrils that deposit in organs - most commonly affecting: Heart, Kidneys, Liver, Gastrointestinal system, Nervous system. **This leads to progressive and often irreversible organ dysfunction.**

Early suspicion and testing are critical. AL-Amyloidosis often presents with vague symptoms, and delays in diagnosis can result in severe organ failure.

Diagnosis should be treated as a medical emergency as rapid intervention is essential to prevent further damage. Treatment focuses on halting light chain production through: Chemotherapy, Monoclonal antibodies, Stem cell transplantation.

A high index of suspicion and prompt action can significantly improve patient outcomes.

Key Diagnostic Testing

Initial Screening Tests

Serum free light chains (kappa & lambda)

Serum electrophoresis with immunofixation

Urine electrophoresis with immunofixation

Troponin T & NT-proBNP

Electrocardiogram

Echo (with strain imaging)/Cardiac MRI (with and without contrast)

Further Diagnostic Testing

Abdominal fat pad biopsy

Bone marrow biopsy

Diagnosis confirmation requires a tissue biopsy with Congo red staining to detect amyloid deposits:

- Abdominal fat pad biopsy
- Bone marrow biopsy

Note: If both biopsies are negative but clinical suspicion for AL-Amyloidosis remains high, consider biopsy of the affected organ.

Different Diagnosis

Abnormal Monoclonal Testing Diagnosis Considerations

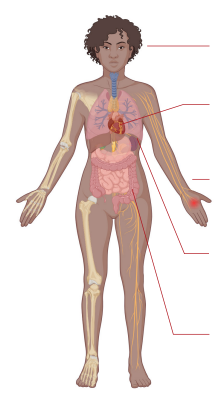
Monoclonal Gammopathy of undetermined significance

Multiple Myeloma, Smoldering Multiple Myeloma, or Light Chain Smoldering Multiple Myeloma

Waldenström Macroglobulinemia

POEMS Syndrome

Clinical Clues



General:

Weakness, unexplained fatigue, macroglossia

Hematologic:

Easy bruising, periorbital purpura (raccoon eyes)

Cardiac:

Dyspnea, hypotension, edema, arrhythmias, increased for L-Ventricularwall thickness

Neurologic:

Peripheral neuropathy, carpal tunnel syndrome, autonomic dysfunction

Renal:

Proteinuria, nephrotic syndrome, kidney dysfunction

Gastrointestinal:

Significant unintentional weight loss, diarrhea/ constipation, malabsorption, unexplained GI bleeding, hepatomegaly

AL-Amyloidosis

Quick Reference Guide



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Diagnosing Amyloidosis

Disease Suspicion See Clinical Clues

Initial Screening Tests

- Serum free light chains (kappa & lambda)
- Serum electrophoresis with immunofixation
- Urine electrophoresis with immunofixation

Abnormal Monoclonal Protein

AL Pathway Yes

Urgent Referral to Hematology

Additional Testing:

- Abdominal fat pad biopsu
- Bone marrow biopsy
- Tissue biopsy confirmation requires detection of amyloid deposits using Congo red staining, with amyloid typing determined by mass spectrometry

Pathway Confirmation of AL- Amyloidosis:

- Tissue biopsy (e.g., abdominal fat pad) Congo red-positive
- Amyloid typing confirms AL (via mass spectrometry)
- Clonal plasma cell disorder identified (bone marrow)

Initiate Hematology-directed

ATTR Pathway

Cardiac Scintigraphy Uptake- Grade 2/3

Yes No

ATTR Cardiac Amyloidosis Unlikely

> Yes **ATTRwt** Wild Type

No **ATTR**_V Variant

Genetic

Testing

Therapy Immediately



Referral Guidance

- Refer suspected AL-Amyloidosis patients to an amyloidosis center when possible.
- If an amyloidosis center is unavailable, prompt consultation with hematology and cardiology is essential.
- A diagnosis of AL-Amyloidosis is a medical emergency that demands timely evaluation and multidisciplinary collaboration to ensure the best possible outcomes.

Patient Advocacy and Support Resources



mm713.ora



AMYLOIDOSIS SUPPORT GROUPS You Are Not Alone

amyloidosissupport.org

