Amyloid Myopathy as an Inclusion Body Myositis Mimic

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ABSTRACT

Introduction: Amyloid myopathy is a rare presentation of systemic amyloidosis. Amyloid myopathy can be initially misdiagnosed as sporadic inclusion body myositis (IBM). Methods: We report 4 cases of amyloid myopathy clinically mimicking inclusion body myositis and initially thought to be phenotypically IBM by neuromuscular experts. Results: Case 1: An 81-year-old female was evaluated for progressive generalized weakness and dysphagia for two years. There was no similar family history. On examination, she had tongue enlargement (Supplementary Fig. 1), MP4 with asymmetric weakness (myopathy pattern 4). Case 2 is a 76-year-old man with primary systemic amyloidosis who presented with myopathy pattern 4 and progressive dysphagia for four years. Case 3 is an 82-year-old man with progressive myopathy pattern 4 weakness and swallowing difficulty. Case 4 is a 62-year-old man with progressive bilateral finger flexor weakness. Muscle biopsies in all 4 cases showed perivascular amyloid deposits. Discussion: Amyloid myopathy may be clinically indistinguishable from IBM. Muscle biopsy is of critical importance in the evaluation of patients suspected to have IBM. Keywords: amyloid, myopathy, amyloidosis, IBM, myositis.

Introduction

Primary systemic amyloidosis is a disease resulting from deposition of amyloid in tissues causing organ dysfunction. Kidneys, heart, peripheral nerves and liver are commonly affected organs.1 Amyloid deposits are formed from monoclonal serum proteins in plasma cell dyscrasia2 and may result from deposition of heavy chain, monoclonal light chains or its N-terminal fragment.3

Amyloid myopathy is a rare presentation of primary systemic amyloidosis. Typically, it presents with proximal muscle weakness and elevated creatine kinase level.4,5

Sporadic inclusion body myositis (IBM) is the most common idiopathic inflammatory myopathy (IIM) after age 50,6 characterized by distal upper extremity and proximal lower extremity weakness, also known as myopathy pattern 4 (MP4).7

A literature search yielded two case reports of amyloid myopathy patients initially presenting with an IBM phenotype.5,8 We present four more cases from four large academic centers.

Methods

We report four cases of amyloid myopathy clinically mimicking inclusion body myositis and initially thought to be phenotypically IBM by neuromuscular experts.

Results

Case 1: An 81-year-old female was evaluated for progressive generalized weakness and dysphagia for two years. There was no similar family history. On examination, she had tongue enlargement (Supplementary Fig. 1), MP4 with asymmetric weakness right and left respectively in knee extension 3/4, hip flexion 2/5, finger flexion 4/3 and shoulder abduction 4/4+. There was decreased pinprick below ankles, and ankle reflexes were absent. She was initially suspected to have IBM. Laboratory testing showed cre-

Supplementary Figure 1. Enlargement of the tongue of case 1.
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Case 1: A 76-year-old female presented with progressive dysphagia for four years and thigh predominant leg weakness for four to five years. Around dysphagia onset, she was diagnosed with heart and renal failure due to primary amyloidosis. Despite chemotherapy, cardiac function declined. Neuromuscular examination demonstrated mild flaccid dysarthria and symmetric limb weakness. Leg weakness was mostly proximal, barely antigravity in quadriceps muscles. Arm weakness was most prominent distally with finger flexors involvement. There was no sensory loss or tongue enlargement. Initially, the patient was suspected to have IBM, despite known systemic amyloidosis diagnosis. EMG revealed myopathy with increased insertional activity. Vastus lateralis muscle biopsy revealed endomyosial and perivascular amyloid deposits (Supplementary Fig. 2). There were no rimmed vacuoles nor inflammation. He died of presumed cardiac arrhythmia or aspiration one month following myopathy diagnosis.

Supplementary Figure 2. Biopsy of the vastus lateralis muscle of case 3. A) H&E stain showing muscle fiber size variability. B) Congo red stain showing salmon-pink amyloid deposits in blood vessel wall and muscle fibers. C) Congo red stain under polarized light showing apple-green birefringent amyloid deposits in blood vessel and muscle fibers.

Case 2: A 76-year-old male presented with progressive dysphagia for four years and thigh predominant leg weakness for four to five years. Around dysphagia onset, she was diagnosed with heart and renal failure due to primary amyloidosis. Despite chemotherapy, cardiac function declined. Neuromuscular examination demonstrated mild flaccid dysarthria and symmetric limb weakness. Leg weakness was mostly proximal, barely antigravity in quadriceps muscles. Arm weakness was most prominent distally with finger flexors involvement. There was no sensory loss or tongue enlargement. Initially, the patient was suspected to have IBM, despite known systemic amyloidosis diagnosis. EMG revealed myopathy with increased insertional activity. Vastus lateralis muscle biopsy revealed endomyosial and perivascular amyloid deposits (Supplementary Fig. 2). There were no rimmed vacuoles nor inflammation. He died of presumed cardiac arrhythmia or aspiration one month following myopathy diagnosis.

Case 3: An 82-year-old male presented with 3 years of progressive proximal weakness. He had difficulty initially lifting arms above his head, followed by difficulty standing up and keeping his head up, then difficulty with buttoning and imbalance leading to falls. He also experienced swallowing difficulties and chronic constipation. Weakness was noted on neck flexion and extension, right greater than left proximal arms, and finger flexors greater than extensors, as well as antigravity strength in proximal legs and knee extensors. Reflexes were absent in the legs with reduced vibration. Lambda free light chain was elevated. EMG showed an irritative myopathy. Deltoid muscle biopsy showed myofiber necrosis, type II atrophy, mild fibrosis and upregulated HLA class I expression (Supplementary Fig. 3). Congo red stains revealed amyloid within arteriolar walls and endomyosial connective tissues. Bone marrow biopsy showed presence of Lambda Bence Jones protein without evidence of amyloid deposition on Congo red stain.

Case 4: A 62-year-old male presented with a one and a half-year history of progressive bilateral hand weakness, worse on the left. Examination showed weakness of flexor digitorum profundus 4+/3 (digits 3, 4 and 5) and 3+/2 (digits 1 and 2), right and left respectively. CK was 280 IU/L. He had normal serum and urine electrophoresis, immunofixation electrophoresis, and kappa to lambda free light chain ratio. EMG showed irritative myopathy. He was initially thought...
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to have IBM. Left biceps muscle biopsy revealed prominent perivascular deposition of amyloid leading to the diagnosis of amyloid myopathy. There was also rare rimmed vacuoles, fiber muscle necrosis and perivascular inflammation. There was HLA Class I sarcolemmal expression, CD3+ T-cells with rare invasion of non-necrotic muscle fibers, and variable expression of phosphorylated neurofilaments on SMI-31 antibody staining (Fig. 2). Immunochemistry for transthyretin was weakly positive but negative for beta-

**Supplementary Figure 3.** Biopsy of the right deltoid muscle. A) H&E stain showing muscle fiber size variability and subsarcolemmal aggregates. B) Congo red stain showing muscle fiber size variability and interstitial thickening/amyloid. C) Congo red stain showing vascular (right) and interstitial (center) amyloid. D) Electron microscopy (EM) 30,000x with interstitial amyloid fibrils, 8-10 nm. E) EM 4800x showing sarcomere Z line disruption. F) EM 2900x of affected vessel.

**Figure 2.** Biopsy of left biceps muscle. A) H&E stain showing endomysial inflammation. B) H&E stain showing perivascular inflammation. C) IHC showing predominantly CD3+ T-cells. D) Non-polarized Congo red stain showing amyloid deposits in blood vessel wall. E) Polarized Congo Red stain confirming the presence of amyloid in thickened blood vessel walls. F) IHC SMI-31 Ab showing phosphorylated neurofilament deposition.

Discussion

IBM classically causes distal arm and proximal leg weakness also known as MP4 but can have a pleomorphic presentation (Table 1). The differential diagnosis of IBM includes inflammatory, autoimmune and genetic diseases, and in some cases, degenerative or even metabolic disorders. Table 1 lists confounders of IBM by phenotypic presentation.

Recognition and description of the clinical pattern of weakness remains an essential part of the diagnostic approach to myopathies. The patients described here, however, illustrate that tissue diagnosis is of key importance. While our patients had clinical presentations consistent with IBM as determined by neuromuscular experts, muscle histopathology revealed a different diagnosis with quite different management.

The clinical scenario was slightly different in our cases. Cases 1, 3 and 4 presented for evaluation of myopathy and were ultimately diagnosed with amyloidosis. Case 2 had an established diagnosis of systemic amyloidosis and only later amyloid myopathy. In cases 1, 2 and 3, the distribution of weakness was striking and notable for distal arm and proximal leg weakness. In case 4, there was weakness more notable in the finger flexors when compared to finger extensors without lower extremity weakness. Therefore, all patients required muscle biopsy to disprove the initial erroneous diagnosis of IBM.

Our study has several limitations. The number of cases is small. The data was retrospectively abstracted. We did not measure antibody titers to cyclic nucleotidase-1A as patients were evaluated before the availability of these autoantibodies. Furthermore, these autoantibodies have been increasingly described to be non-specifically elevated in various inflammatory myopathies and autoimmune rheumatologic disorders. For example, these are found in Sjögren’s syndrome (23-36%), systemic lupus erythematosus (14-20%) and dermatomyositis (15%).

For case 4, follow up duration was short. Though histopathologic evidence of inflammation was described previously in amyloid myopathy leading to an erroneous diagnosis of polymyositis, rare rimmed vacuoles would be quite an unusual finding. The rarity of these vacuoles renders them...
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Table 1: IBM phenotypes and differential diagnosis according to clinical patterns.

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<tr>
<th>IBM Phenotypes</th>
<th>Differential diagnosis</th>
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<tr>
<td>Distal Arm/Proximal Leg (MP4)</td>
<td>Myotonic MD, Sarcoidosis, Amyloid Myopathy</td>
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<tr>
<td>Distal Weakness (MP2)</td>
<td>Myotonic MD, ALS</td>
</tr>
<tr>
<td>Quadriceps Atrophy</td>
<td>Becker MD, LGMD, Emery-Dreifuss, PM, SMA</td>
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<tr>
<td>Limb-girdle pattern (MP1)</td>
<td>Polymyositis, LGMD, Pompe, Amyloid Myopathy, Sarcoidosis</td>
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<tr>
<td>Bulbar (MP7) - dysphagia</td>
<td>MG, LEMS, ALS, LGMD</td>
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non-diagnostic. Furthermore, these have been non-specifically observed in other disorders such as polymyositis, hereditary IBM, and neurogenic atrophy.16

All of our cases fulfilled the 2011 ENMC clinical and laboratory criteria for IBM (Supplementary Table 1). Cases 3 and 4 even fulfilled criteria for clinically defined IBM and probable IBM respectively. This is given pathologic evidence of HLA class 1 expression in Case 3 and inflammation with myofiber invasion in Case 4. This highlights the importance of obtaining muscle biopsy in patients with IBM and that Congo Red staining should be part of the routine evaluation of IBM muscle. Beyond that, it raises questions about the specificity of the ENMC 2011 IBM criteria. For instance, if Congo Red stain was not done in cases 3 and 4, the diagnosis would have been IBM based on these criteria (Supplementary Table 1).

Conclusion
Amyloid myopathy may clinically mimic IBM and manifest as an initial or delayed presentation of primary systemic amyloidosis. Despite the availability of antibody testing, muscle biopsy remains critically important in the evaluation of patients suspected to have IBM and serves to distinguish IBM from its many mimics.

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Bibliography


