Superimposed Polymyositis in a Patient with Myeloperoxidase-Related Crescentic Glomerulonephritis

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Abstract

Renal involvement especially glomerulonephropathy in polymyositis or dermatomyositis is considered rare. Crescentic glomerulonephritis is even more so. Herein, we report a patient who was diagnosed with concurrent polymyositis and myeloperoxidase-related crescentic glomerulonephritis. We discuss the association between polymyositis or dermatomyositis and renal involvement, glomerulonephropathy, and crescentic glomerulonephritis. This is the first report of a case of polymyositis and myeloperoxidase-related crescentic glomerulonephritis with a clear temporal correlation.

Keywords: Myeloperoxidase-related crescentic glomerulonephritis; Polymyositis; Glomerulonephropathy

Introduction

The major presentation of polymyositis or dermatomyositis is the obvious muscle weakness and skin rash. The superimposed renal involvement had been previously considered as rare. However, the notion of its scarcity is incorrect after we performed this literature review focusing on renal involvement in polymyositis or dermatomyositis. The acute tubular injury is the most prevalent type of renal involvement followed by glomerular injury. Of all the glomerular injury, crescentic glomerulonephritis is the least common. This article will present the first case of pauci-immune related crescentic glomerulonephritis superimposed with polymyositis.

Case Report

A 72-year-old man was presented with kidney injury and gradual dysphagia for 2 months at the outpatient department. Uremia-related dysphagia was first excluded due to low blood urea nitrogen (BUN:59 mg/dL). Duodenoscopy revealed no mechanical obstruction. Patient later claimed about his muscle soreness. Biochemical data were as follows: creatinine kinase, 2472 U/L; lactic dehydrogenase, 450 U/L; aldolase, 21.7 U/L; Jo-1, 139 units and anti-nuclear antibody, 1:160. Esophageal transit time was prolonged and electromyography also suggested myopathy. Myositis scan revealed diffuse myositis at four extremities. Polymyositis was highly suspected even without muscle biopsy. Malignancy surveys including chest X-ray (CXR), enteroscopy, and abdominal sonography were negative. We prescribed 100 mg of hydrocortisone three times per day as his major treatment. The patient’s serum creatinine was increased from 0.8 mg/dL to 6.5 mg/dL with hematuria and subnephrotic range of proteinuria. His immunological data disclosed negative double-strand DNA (dsDNA), normal proteinase 3, and high myeloperoxidase (MPO, 109.2 units) in the absence of anti-glomerular basement membrane (GBM) antibody. Furthermore, his renal biopsy showered cellular crescent (Figure 1) and mesangial proliferation. The immunofluorescence microscopy was negative for IgA staining and there were no full-house or double-contour staining nor linear or granular staining over GBM, which suggest pauci-immune glomerulonephritis. Five days later, double-filtration plasmapheresis (DFP) was administered 5 times per day. We did not administer methylprednisolone or cyclophosphamide pulse therapy to the patient owing to concern of sepsis. After five-time-DFP, we initiated hemodialysis (HD) due to high BUN (107 mg/dL), and metabolic acidosis (13 meq/L of HCO3–). Soon after HD, the patient went into shock and Klebsiella pneumoniae was found in blood and sputum with pneumonia over right lower lung. Two days later, the patient was expired due to refractory signs of shock regardless of piperacillin and tazobactam use.

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Materials and Methods

We performed an extensive literature review from MEDLINE. Articles writing in all languages were included. There are only four cases with MPO-related crescentic glomerulonephritis in polymyositis/dermatomyositis. We had summarized the characteristics in Table 1.

Table 1: MPO-related crescentic glomerulonephritis in polymyositis/dermatomyositis

<table>
<thead>
<tr>
<th>Year reported</th>
<th>gender</th>
<th>age</th>
<th>Period (years) between PM/DM and crescentic glomerulonephritis</th>
<th>Level of MPO (EU)</th>
<th>outcome</th>
</tr>
</thead>
<tbody>
<tr>
<td>2013 (present study)</td>
<td>male</td>
<td>60</td>
<td>10</td>
<td>1280</td>
<td>uremia</td>
</tr>
</tbody>
</table>

Discussion

Renal involvement in polymyositis or dermatomyositis (PM/DM) had been considered as rare. However, according to a recent report of 14 PM/DM patients [1], its incidence rate is 25.1% (14 patients). These patients are categorized into myoglobin-related acute tubular injury (ATI) and glomerulonephropathy. Here raises the questions of why renal involvement is seem as uncommon. In order to understand the reasons behind the issue, first we acknowledge that PM/DM itself is a rare disease. Second, although ATI is common, it could easily be missed without meticulous checkup due to its self-limiting well prognosis. Third, most clinicians are reluctant to perform renal biopsy just to prove ATI. Therefore, if only we could precisely diagnose every ATI as renal involvement in PM/DM, the incidence rate will not be low. In the study by Yan et al. [1], 9 of the 14 PM/DM patients (64.3%) with renal involvement had ATI but only 2 patients (14.3%) had biopsy-proved glomerulonephropathy. The incident of glomerulonephropathy is still rare.

According to a literature [2], only 22 cases of PM/DM with glomerulonephropathy had been reported worldwide. Adding on our patient (23 patients), 12 patients had mesangial proliferative glomerulonephritis, 6 had membranous glomerulonephropathy, 4 had crescentic glomerulonephritis (CGN), and 1 patient had minimal change disease. We believe this scarcity is due to under diagnosis. Clinical manifestations of PM/DM are more dominant than those with glomerulonephropathy and clinicians would usually first prioritize treating the more dominant symptoms of PM/DM. Patients’ glomerulonephropathy would then be left partially treated and causing the status of its under diagnosis. Our patient had mesangial proliferation, but IgA nephropathy or lupus nephritis was unlikely due to negative stain for IgA and lack of full-house staining pattern or dsDNA. CGN were also detected and it is even more uncommon currently.

There are only 4 reported cases of PM/DM with CGN [3]. To the best of our knowledge, this patient in our study is the 5th case. They were all immune-complex related and what differs from our case to the others is that our patient’s report shows pauci-immune, which has never been previously described in literatures. Reaizlizing that complement deposits attack vascular endothelium in muscles, damage muscle capillaries, and attack glomerular membrane, we know they all act to compromise the protective mechanism of the capillaries. Therefore, the mechanism of the immune-complex-related CGN is reasonable and myoglobin-anti-myoglobin antibody-related IC has been reported. However, this case was diagnosed as MPO-related CGN. After extensive reviews, we found three more cases of MPO-related CGN [4-6], but DM occurred at least 3 years before CGN. Correlation between MPO-related CGN and PM/DM is not yet clear. To the best of our knowledge, our case is the first attempt to describe a clear temporal association between MPO-related CGN and PM/DM (Table 1). Beforehand, concurrence was considered coincidental because there are at least 3 years elapsed between the onset of these two diseases. Henceforth, the likelihood of temporal correlation should be considered strong and clinicians should be aware of the signs of PM/DM with renal function impairment. We highly suspect that the rarity of concurrence is mostly due to under-diagnosis, which may be caused by partial treatment of PM/DM with steroids.

Conclusion

Renal function involvement in PM/DM is not as uncommon as most clinicians believe in the past. Most of the reported cases have involved ATI, and we should always keep in mind the possibility of concurrent MPO-related CGN and PM/DM.

Learning Point for Clinicians

Renal involvement in polymyositis/dermatomyositis is rare due to partial treatment and under diagnosis. Clinicians should take heed about crescentic glomerulonephritis and myeloperoxidase-related crescentic glomerulonephritis regarding its underestimated prevalence and treatment methods.

References