Case report

HHV-8-negative multicentric Castleman disease presenting as a crescentic immune complexes membranoproliferative glomerulonephritis

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SUMMARY
Multicentric Castleman disease is a rare polyclonal lymphoproliferative disorder mainly associated with two renal manifestations: thrombotic microangiopathy and amyloidosis. Nevertheless, we report here a case of human herpes virus-8 negative multicentric Castleman disease with membranous proliferative glomerulonephritis and extracapillary proliferation. A patient was successfully treated with corticosteroids, anti-CD20 and cyclophosphamide therapy.

BACKGROUND
Castleman disease is a rare polyclonal lymphoproliferative disorder characterised by three histological patterns: hyaline vascular, plasma cell and plasmablastic.2 Clinical presentation is heterogeneous and classified into unicentric or multicentric disease.1,4–6 Unicentric Castleman disease is the most frequent type.1 It involves a single lymph nodal region, and symptoms are mainly due to local lymph node enlargement.5 Its surgical resection is curative.3 By contrast, multicentric Castleman disease involves multiple lymph node areas.4–6 It is subclassified into the human herpes virus-8 (HHV-8) related or not.5–7 Multicentric Castleman disease is a severe multisystemic disease, with nonspecific symptoms such as lymphadenopathy, fever, weight loss, fatigue, oedema and ascites.3,8 The high level of cytokines production, notably interleukin-6 (IL-6),7 can affect each system and eventually lead to multiorgan failure.6 IL-6 pathway is the therapeutic target.3 5 9 10 Nevertheless, recurrence is frequent and prognosis poor.10 Renal involvement is also common1,1 and various histological patterns have been described. A membranous proliferative glomerulonephritis has been previously reported, nevertheless extracapillary proliferation has not been described yet.

CASE PRESENTATION
A 48-year-old man from the Middle East presented to our hospital because of night sweats, lower extremity oedema and a weight loss of 12 kg in 3 months.

He had a medical history of dyslipidaemia, severe obesity (body mass index of 35 kg/m²) and 25 pack-year of previous smoking. 2.5 months before admission, he was admitted to another hospital because of an acute left limb pain. A multisegmental artery occlusion of the limb was then diagnosed which justified an anticoagulation by vitamin K antagonist.

The patient had no other complaints. Except for increased blood pressure and bilateral pitting oedema, clinical examination was normal.

INVESTIGATIONS
Blood test showed acute kidney injury (creatininaemia 200 µmol/L) with hypoalbuminaemia (26 g/L), an elevated sedimentation rate (>100 mm/hour) and elevated CRP (90 mg/L). Urinalysis showed erythrocytes cast. Twenty-four hour urine protein excretion was 9.24 g/day. Finally, a kidney ultrasound with Doppler showed no anomaly. Because of the nephritic syndrome, a renal biopsy was performed. On microscopic examination (23 glomeruli), we observed a membranous proliferative pattern with the presence of a crescent (figures 1 and 2).

Immunofluorescence was positive for IgA (+/–), rare deposits, IgG (+/+), IgM (+/–), C3 (+/+), C1q (+) and C5b9 (+/–) deposits (full house), but kappa and lambda light chains were negative (figures 3 and 4). The electron microscopy disclosed the presence of subendothelial and subepithelial deposits. We estimated between 20% and 30% of fibrosis. At this stage, a crescentic immune complex glomerulonephritis with a membranoproliferative pattern was suspected.

The immunological workup was negative: absence of monoclonal spike but a polyclonal hypergammaglobulinaemia on the immunofixation; negative antimicrobial antibodies and antineutrophil cytoplasmic antibodies, normal C3 and C4 as well as the absence of cryoglobulinaemia and antiphospholipid antibodies. An exhaustive infectious workup was also done with negative polymerase chain reaction (PCR) for cytomegalovirus (CMV), Epstein-Barr virus (EBV), hepatitis B, C, HIV and HHV-8, negative bacterial serologies for leptospiriosis, Brucella spp, Borrelia burgdorferi, Coxiella burnetii, Bartonella henselae, syphilis, as well as toxoplasma. Finally, interferon-gamma release assay was negative as well as urine culture for mycobacteria. Blood culture was also negative. Although, a high level of IL-6 (18.5 µg/L, n<1 µg/L) prompted us to search for an inflammatory process. A 2-fluoroxyglucose (2-FDG) positron emission tomography scan revealed supra and subdiaphragmatic hypercapitive adenopathies, a mediastinal mass hypercaptation.
and a splenic abnormal FDG uptake. A left axillary lymph node resection was performed. Pathology determined a Castleman-like pattern with hyperplastic adenopathy, associated with follicular hyperplasia, interfollicular expansion and significant polytypic plasma cells and circularly capillary penetrated some germinal centres. There was no evidence of lymphoma. HHV-8 and EBV immunohistochemistry were negative. Moreover, fewer than 10% of plasma cells were IgG4 positive.

DIFFERENTIAL DIAGNOSIS

We concluded that this patient suffered from HHV-8 negative multicentric Castleman’s disease. Treatment was initiated with corticosteroids, anti-CD 20 therapy with rituximab and cyclophosphamide.

In our case, due to the absence of monotypic deposits, a monoclonal gammopathy of renal significance is easily excluded. In the absence of a predominant C3 deposits, a C3 glomerulopathy seems unlikely. The full-house pattern points to two hypotheses: lupus nephritis or an infectious glomerulonephritis.

In the absence of signs or symptoms of lupus and antinuclear antibodies, a diagnosis of lupus nephritis is excluded. Finally, an infection-related glomerulonephritis is unlikely because of the negativity of the full infectious workup. Moreover, improvement of the clinical status under a strong immunosuppressive therapy does not suggest an infectious process and supports that Castleman disease is the only possible aetiology.

Therefore, we concluded that the crescentic immune complex glomerulonephritis with a membranoproliferative pattern was secondary to multicentric Castleman disease.

OUTCOME AND FOLLOW-UP

The patient quickly improved: a few days after corticoids administration the fever and inflammatory markers decreased. A 2-FDG positron emission tomography scan was repeated 3 months later, which showed regression of the lymphadenopathies without hypercaption. In parallel, proteinuria level decreased completely.
after 2 months and renal function improved slowly. One year later, creatinaemia was 144 μmol/L.

DISCUSSION
Clinicians investigating membranoproliferative glomerulonephritis must do a comprehensive search to find the right aetiology and the nature of the immune deposit is the principal clue.

Up to 54% of multisystemic Castleman disease, mostly in plasma cell or mixed cellular type, are associated with nephropathy, defined as haematuria, proteinuria or renal insufficiency.11–14 The two main histological patterns are thrombotic microangiopathy (60%) and amyloidosis (20%).15–16 Those histological patterns could be explained by the dysregulation of IL-6 production and vascular endothelial growth factors in the lymph node mantle.15–19 Indeed, IL-6 proinflammatory cytokine stimulates the precursor of the AA protein and vascular endothelial growth factors, leading to thrombotic microangiopathy.20–21

Nevertheless, our patient had proliferative immune complex glomerulonephritis. We hypothesised that the high level of IL-6 promoted B cell activation, which in turn induced immune complexes. Those immune complexes get ‘trapped’ in the subendothelial space of the glomeruli and lead to a proliferative glomerulonephritis.

In the recent years, new treatments for HHV-8-negative/idiopathic multicentric Castleman disease (iMCD) have been developed, such as anti-IL-6 monoclonal antibody (anti-IL-6 mAb) siltuximab or tocilizumab.9,22 They are now considered as preferred first-line therapy,22 sometimes in association with corticosteroid.22 Anti-CD20 mAb (rituximab) is an alternative for anti-IL-6 mAb in the case of non-severe iMCD (defined as a good performance status without organ dysfunction), however, with a lower proportion of complete responses and progression-free survival compared with siltuximab.23

Learning points

► Castleman disease is a rare non-neoplastic lymphoproliferative disorder characterised by various renal manifestations with two main histological patterns: thrombotic microangiopathy and amyloidosis.

► Castleman disease is also a possible aetiology for a proliferative immune complex glomerulonephritis after exclusion of other frequent causes such as infection-related glomerulonephritis.

► The anti-interleukin-6 monoclonal antibody, siltuximab or tocilizumab are considered as first-line treatment for human herpes virus-8-negative/idiopathic multicentric Castleman disease.

Patient’s perspective

The experience of sickness, the lose of strength, especially when you are a man of 48 years old, is hard to accept. I did not understand what was wrong with my body. I was afraid to die. During hospitalisation doctors had to performed a lot of tests to find out why I was losing weight and why my kidneys were failing. Then, I was shocked to learn that I had a rare disease called Castleman disease. I asked myself several times: why me? With the help of the medical team I understood better my disease and I started a treatment. At the end, I feel lucky because the treatment improved my condition and my kidneys function, and I have the support of my family. Currently I have almost a normal life.

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