CASE REPORT

Cast nephropathy: an extremely rare renal presentation of Waldenström’s macroglobulinaemia

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SUMMARY

Renal involvement in Waldenström’s macroglobulinaemia (WM) is very unusual when compared to multiple myeloma. We report a case of a patient who developed anuric acute kidney injury secondary to cast nephropathy, dependent on high-flux haemodialysis. Complementary study revealed the presence of blood IgM monoclonal gammopathy and a massive bone marrow lymphoplasmacytic infiltration. There were no osteolytic lesions and no clinical signs/symptoms of hyperviscosity syndrome. The diagnosis of WM was established and a dexamethasone plus cyclophosphamide regimen was started, in addition to plasmapheresis. The patient partially recovered renal function allowing haemodialysis and plasmapheresis withdrawal. He remained asymptomatic with a good response to chemotherapy and 12 months after his renal function remained stable. This is a rare clinical case in which WM presented as an IgM cast nephropathy, which in turn is an extremely rare renal presentation of this equally rare haematological disorder.

BACKGROUND

Waldenström’s macroglobulinaemia (WM) is a rare, low-grade malignant lymphoplasmacytic proliferative disorder accounting for <5% of all non-Hodgkin lymphomas and for approximately 2% of all haematological malignancies.1 2 It is a distinct clinicopathological entity characterised by bone marrow (BM) infiltration of a lymphoplasmacytic lymphoma (LPL), responsible for immunoglobulin M (IgM) monoclonal gammopathy in the blood.2 Other organs can be infiltrated, especially liver and spleen. Renal involvement in WM is unusual when compared to multiple myeloma (MM) and results in monoclonal IgM deposition in the glomerular basement membrane (GBM) (±cryoglobulinaemia), interstitial infiltration of neoplastic lymphoplasmacytic cells and/or amyloid light-chain (AL)-amyloidosis typically causing nephrotic syndrome.3 Cast nephropathy can be seen more frequently in MM (commonly referred to as ‘myeloma kidney’, a myeloma-defining event) and is otherwise extremely rare in WM.4 4 We report a case of WM where cast nephropathy with acute kidney injury (AKI) was not only its renal manifestation but also its initial presentation.

CASE PRESENTATION

An 80-year-old Caucasian man was referred to our nephrology department after a 24 h history of anuric AKI. Concurrently, he presented with an acute respiratory infection and was treated with antibiotics, bronchodilators and oxygen.

The patient’s medical history was notable for diabetes mellitus (for 20 years, without evidence of diabetic retinopathy), arterial hypertension, glaucoma and asymptomatic cholelithiasis. There was no known renal disease and his last known serum creatinine, 3 years earlier, was 1.1 mg/dL. His outpatient medication was composed exclusively of metformin, glibenclamide, enalapril, clopidogrel and ophthalmic drops.

On admission, he was afebrile with a blood pressure of 139/56 mm Hg, bilateral pulmonary ronchi but no respiratory distress. Cardiac and abdominal examinations were unremarkable and there were no cutaneous lesions, peripheral oedema or adenopathies.

Initial laboratory studies yielded the following: normocytic and normochromic anaemia (Hb=9.3 g/dL) with normal leucocyte and platelet blood count, elevated C reactive protein (6.48 mg/dL, normal range (NR) <0.5 mg/dL), serum creatinine 7.91 mg/dL, blood urea nitrogen 89 mg/dL, hyperkalaemia 5.8 mmol/L, hyperglycaemia 263 mg/dL with borderline hyponatraemia of 135 mmol/L and non-compensated metabolic acidosis (pH=7.22, bicarbonate 15.1 mmol/L, carbon dioxide pressure 37 mm Hg, oxygen pressure 69 mm Hg and normal serum lactate). Chest X-ray was negative for pneumonic foci or pleural effusions. Renal ultrasound revealed normal kidneys with no signs of obstruction or nephrolithiasis.

During the first few hours, the patient remained anuric and was started on high-flux haemodialysis. Additional and relevant studies included: total protein/albumin of 7.3/2.9 g/dL, normal lactic acid dehydrogenase, corrected calcium 9.2 mg/dL, hyperphosphoremia of 5.3 mg/dL and parathormone of 120 pg/mL (NR 9–72 pg/mL). Glycated haemoglobin was 6%, uric acid 3.9 mg/dL (NR 3.5–7.2 mg/dL) and folic acid 3.6 ng/mL (NR >5.4 ng/mL). The levels of rheumatoid factor, complement C3 and C4 and anti-streptolysin O titre were normal. Antinuclear antibodies were also normal and cryoglobulins were absent. Viral serologies were negative for hepatitis B surface antigen, hepatitis C antibody, HIV 1 and 2 antibodies, Epstein-Barr virus, cytomegalovirus and herpes simplex virus. Peripheral blood cultures were sterile. Serum electrophoresis revealed a monoclonal spike in the β region, which was identified by immunoelectrophoresis and immunofixation as an IgM M component bearing κ light chains: IgM 35.40 g/L (NR 0.40–2.30 g/L), κ light chains


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16.78 g/L (NR 6.66–14.65 g/L), κ/λ relation 3.94 (NR 1.35–2.65), IgG 6.56 g/L (NR 7.0–16.0 g/L), IgA 0.65 g/L (NR 0.70–4.00 g/L) and IgE 24 UI/mL (NR 0–100 UI/mL). Serum viscosity was not performed, as signs/symptoms of hyperviscosity were absent. Serum β2-microglobulin was 16.70 mg/L (NR 1.09–2.53 mg/L).

Urine sediment (when the patient recovered urine output) contained many erythrocytes, leucocytes and pathological cylinders. Urine microbiological examination was negative. Proteinuria was 4058 mg/g. Urine immunoelectrophoresis revealed albumin and κ light chains (0.98 g/1 L urine/24 h).

BM aspirate revealed 75% of lymphoplasmacytic cells with 7% of plasmocytes and BM biopsy showed a massive infiltration (90%) by lymphoplasmacytic cells. Flow cytometry demonstrated 35% of IgM and κ light chain clonal B cells with the following immunophenotype: CD19+, CD20+, CD79b+, CD38+ (heterogeneous expression), IgM and κ light chains, CD10–, CD5– and CD23–. This immunophenotype was compatible with LPL/WM and no chromosomal abnormalities were explored. Skeletal X-ray was negative for osteolytic lesions, and a CT scan revealed multiple thoracic and abdominal lymphadenopathies without hepatosplenomegaly.

During the first week, the patient remained dependent on haemodialysis and a renal biopsy was performed (figures 1–6).

Light microscopy showed mild thickening of the capillary wall and a slight mesangial expansion, PAS positive, without hypercellularity. Double contoured basement membranes were absent. There was mild interstitial fibrosis, signs of ischaemic tubulopathy and mild multifocal tubulointerstitial nephritis. Many renal tubules showed PAS-positive casts, which became polychromatic on Masson’s Trichrome stain. Immunohistochemical stain revealed IgM deposits along the GBM and mesangium. Tubular casts and peritubular capillaries also stained intensively for IgM and moderately for κ light chains. Congo red and Wright staining were negative. Immunofluorescence study revealed a strong staining of casts and glomeruli for IgM, at this level granular, mesangial and subepithelial. Electron microscopy was not performed.

The patient was diagnosed with an AKI secondary to cast nephropathy due to WM and was started on intravenous cyclophosphamide 750 mg/m² every 21 days, plus oral dexamethasone 10 mg twice a week. Also, plasmapheresis guided by serum para-protein levels was performed on alternate days (therapeutic plasma exchange using a centrifuge, with an exchange of 1.0 plasma volume and using 5% human albumin plus fresh-frozen plasma as replacement fluids) as well as MESNA protocol and antibiotic prophylaxis. During hospitalisation, the patient was
started on insulin therapy and submitted to 14 plasmapheresis treatments plus three cycles of chemotherapy. There was a significant reduction in serum IgM and κ light chains to 7.74 and 5.56 g/L, respectively. The patient’s renal function slowly recovered, becoming independent of haemodialysis, with a stable serum creatinine of 4.5 mg/dL. His hospital admission was otherwise uneventful and he was discharged after 56 days, dialysis free. He was maintained on regular surveillance and a year later remained asymptomatic with a serum creatinine of 2.68 mg/dL, a bland urine sediment and proteinuria of 463 mg/g.

DISCUSSION

Our patient presented with a serum IgM monoclonal gammopathy and an AKI secondary to a myeloma-like cast nephropathy, with unusual and strong staining for IgM in addition to κ light chains. In this case, and in general, the differential diagnosis between rare diseases such as WM and classical MM with an IgM paraprotein (IgM-MM) is challenging. The distinction is made on clinical and cellular grounds. Classical MM is associated with lytic bone lesions and AKI. Symptoms of hyperviscosity, the presence of lymphadenopathy and/or splenomegaly, and the absence of osteolytic lesions support the diagnosis of WM. Moreover, lymphoplasmacytic cells in the BM of patients with WM are immunophenotypically different from plasma cells seen in the BM of patients with MM. The clonal population in WM appears to derive from a B-cell arrest before terminal differentiation into a plasma cell and a surface IgM-positive CD5–CD10–CD19+CD20+CD23– immunophenotype in association with an intra-trabecular monoclonal lymphoplasmacytic infiltrate is diagnostic of WM.2 Cytogenetic studies are optional in the diagnostic approach of a suspected case of WM but sometimes helpful in patients with MM who may have lymphoplasmacytic or small mature plasma cell morphology, also expressing CD20. In these cases, cytogenetic analysis to search for t(11;14) chromosomal translocation may help establishing the diagnosis, and its presence is in favour of an IgM-MM as WM is not associated with specific chromosomal abnormalities. In the present case, there was no cytogenetic analysis as the clonal population had a characteristic immunophenotype that, along with the clinical criteria, led to the diagnosis of WM.

According to the literature, there are few case reports describing patients with WM and cast nephropathy.6–9 Ours was unique in two ways: (1) AKI was the clinical manifestation that led to the diagnosis of WM, which is in itself a rare event in WM and (2) renal biopsy documented the presence of cast nephropathy with tubular casts expressing IgM and κ light chains on immunohistochemical staining, which is also very unusual in WM.

It is widely known that light-chain cast nephropathy, commonly referred to as myeloma cast nephropathy or myeloma kidney, is the most common cause of AKI in MM, and is related to a high tumoural burden.9 In MM, monoclonal free light chains (FLC) are easily filtered through the glomerular filtration barrier (Bence-Jones proteinuria). However, the high FLC concentration overwhelms proximal tubule reabsorption capacity, allowing for large amounts of FLC to reach the loop of Henle, where Tamm-Horsfall protein (THP) is produced. Here, they form obstructing tubular casts, which are the result of FLC’s binding and aggregation to THP. The tubulopathic FLC typically incite accompanying tubulointerstitial nephritis.

Urinary monoclonal light chains can be detected by immunofixation in 80–90% of patients with WM, but significant Bence-Jones proteinuria (>1.0 g/day) only occurs in a small fraction of those patients (<5%).3 This may help to explain why cast nephropathy (and AKI) is so rare in WM.

Possible pathological patterns of renal involvement in WM are: (1) glomerular lesions and (2) tubulointerstitial lesions. The first, which are most frequently encountered, consist of intracapillary monoclonal deposit disease (intraglomerular periodic acid-Schiff (PAS)-positive hyaline pseudothrombi occluding the capillary lumens corresponding to monoclonal IgM aggregates), membranoproliferative glomerulonephritis (cryoglobulinemia), light chain amyloidosis, light-chain deposition disease (LCDD) and minimal change disease. The second are related to lymphomatous infiltrate, acute tubular necrosis, cast nephropathy, Fanconi syndrome and, also, LCDD.9–13

In our patient, glomerular and tubulointerstitial compartments were both affected. Immunohistochemical stain and immunofluorescence pattern may suggest a deposit disease. However, a definite diagnosis is not possible, as electronic microscopy was not performed. Had it been confirmed, there would be a combination of cast nephropathy and deposit disease, an even rarer histopathology in WM.11 But it is important to stress that patients’ staining patterns may represent non-specific ‘trapping’ of the circulating M component in basement membrane, which might be a reflection of high serum levels as opposed to real deposits.11

**Figure 5** The κ light chain immunohistochemical expression in the casts, some of them showing lamination (arrow). (κ light chain, ×400).

**Figure 6** IgM 3+ deposits in casts (arrows) (IgM immunofluorescence, ×400).
It was also very interesting to find IgM staining in tubular casts. At first sight, this might be difficult to understand due to its pentameric form and high molecular mass. However, IgM has important physiochemical properties, such as aggregability, polymerisation and cryoprecipitation, which may predispose to its GBM trapping particularly where protein concentration is especially elevated by glomerular ultrafiltration. It may somehow induce glomerular damage (without triggering glomerular proliferation) and non-selective proteinuria, explaining particularly elevated protein excretion in AKI.

Another relevant issue in our patient’s renal dysfunction was his long history of type 2 diabetes mellitus. However, renal histopathology excluded co-existing diabetic nephropathy, which could otherwise justify his residual renal dysfunction after cast nephropathy treatment.

WM’s immunosuppressive treatment is the same as for MM, and includes alkylating agents, purine analogues and monoclonal antibody anti-CD20 (rituximab). The latter may be regarded as a reasonable option in patients with IgM autoantibody-related neuropathies (anti-myelin-associated glycoprotein antibodies), which were absent in our patient. Moreover, plasmapheresis is indicated for the acute management of symptoms of hyperviscosity (generally present when IgM>50 g/L), and, in cast nephropathy, as an adjunct to chemotherapy, as it helps remove the toxic circulating paraproteins (while their production is being impaired by chemotherapy). However, there are conflicting data regarding its effectiveness in patients with AKI secondary to cast nephropathy.

Despite having cast nephropathy, renal prognosis in this patient with WM was better than for most patients with MM with similar renal involvement. Even so, median survival of patients with WM, from the time of diagnosis, is approximately 5 years and the majority die of disease progression (eg, transformation to large cell lymphoma). The remaining succumb to unrelated causes, as they are generally elderly.

Learning points

- Compared to multiple myeloma (MM), renal involvement in Waldenström’s macroglobulinaemia (WM) is very unusual, with glomerular lesions being the most frequently observed abnormality.
- Cast nephropathy is extremely rare in WM since significant Bence-Jones proteinuria occurs in <5% of these patients.
- In rare cases, IgM trapping in glomerular basement membrane may induce glomerular damage with the appearance of IgM in tubular casts.
- WM treatment is the same as that for MM.
- Management of cast nephropathy with acute kidney injury includes chemotherapy with or without plasmapheresis, according to the current literature.

Competing interests None declared.

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REFERENCES