IgD Myeloma with Two Types of Paraproteinemic Kidney Damage: Case Report

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Abstract

Clinical presentation of multiple myeloma and other lymphoproliferative disorders is characterised by specific haematological signs and symptoms, by a broad spectrum of kidney lesions. Differential diagnostics is the major challenge, demanding pathology evaluation of kidney tissue, as different types of lesions, mainly paraproteinemic, cannot be differentiated solely on the basis of clinical features. Importantly, symptoms of kidney damage may dominate over LPD symptoms, and even preclude overt LPD’s.

IgD myeloma is rare, but aggressive tumour, diagnosed approximately in 2-2.5% cases of MM. Diagnostic problems arise from the fact, that the routine test does not detect the M-spike in 60% of patients, and when it is detected, the concentration is usually smaller than 2g/dl. An overproduction of light chains (LC) is found in 90-96% of patients. The disease course is often accompanied by AL amyloidosis and light chain proteinuria, renal failure is observed in 33% of cases at the moment of diagnosis.

The pathology pattern of renal damage in patients with LPD, showing a combination of cast-nephropathy and light chain deposition disease is also rare and reported in literature like single cases or small series.

We present here a case of IgD myeloma, manifested with combined kidney injury – light chain deposition disease and cast nephropathy.

Keywords: Myeloma; Immunoglobulin D; Cast Nephropathy; Light Chain Deposition Disease

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Clinical presentation of Multiple Myeloma (MM) and other Lymphoproliferative Disorders (LPD) is characterized, beyond specific haematological signs and symptoms, by a broad spectrum of kidney lesions. Renal involvement is defined by numerous mechanisms, but mainly by the deposition of secreted paraproteins. Paraproteinemic lesions, in turn, may be represented by organized (crystals, fibrils, microtubules), and non-organized deposition of monoclonal immunoglobulin’s or fragments thereof, mostly light chains (LC). They may involve the four components of the kidney parenchyma: glomeruli, tubules, interstitium and blood vessels. Organized depositions include Cast Nephropathy (CN), light-chain tubulopathy, AL/AH amyloidosis, glomerulonephritides with an organized microtubular monoclonal deposits, and cryoglobulinic Glomerulonephrites (GN). Non-organized deposition group comprise Light Chain/Heavy Chain Deposition Disease (LCDD/HCD), proliferative GN with monoclonal deposits of IgG/IgA, and non-proliferative GN with monoclonal deposits of IgM. It appears that amino acid sequence of the monoclonal LC and other monoclonal proteins, defining inherent biochemical properties, is the primary determinant of the specific pattern of renal parenchymal deposition and clinical disease. Patients may present with Acute Kidney Injury (AKI), Nephrotic Syndrome (NS), proteinuria and/or haematuria, arterial hypertension or chronic kidney disease. Differential diagnostics is the major challenge, demanding pathology evaluation of kidney tissue, as the above mentioned lesions cannot be differentiated solely on the basis of clinical features. Importantly, symptoms of kidney damage may dominate over LPD symptoms, and even preclude overt LPD’s [1-13].

MM presents with end organ damage manifested as one or more of the following – bone pain, kidney damage, impaired haematopoiesis, hypercalcemia, and susceptibility to infections. In laboratory tests MM typically manifests itself by the presence of high paraprotein (mainly IgG, IgA and Bence-Jones) levels in serum and urine. Sometimes, however, the clinical picture of MM is quite different from the classic manifestation, which can cause diagnostic difficulties, thereby delaying the treatment. One of these rare and unusual forms of MM is IgD myeloma.

Plasma cell leukemia with IgD paraprotein was firstly described in 1968, and since that time IgD myeloma remains rare, but aggressive tumour, diagnosed approximately in 2-2.5% cases of MM and affecting young people. Diagnostic problems arise from the fact, that the routine test does not detect the M-spike in 60% of patients, and when it is detected, the concentration is usually smaller than 2g/dl. An overproduction of LC (Bence-Jones paraprotein), usually λ, is found in 90-96% of patients.
The disease course is often accompanied by AL amyloidosis and LC proteinuria, renal failure is observed in 33% of cases at the moment of diagnosis. Lymphadenopathy is seen in 10% of patients [14-21].

The pathology pattern of renal damage in patients with LPD, showing a combination of cast nephropathy and LCDD is also rare – such coexistence is reported in literature like single cases or small series. At least two possible explanations for these findings suggested so far. It was observed, that both fibrillar and non-fibrillar monoclonal LC deposits may coexist in the same patient, and the identity of the amino acid sequence of the deposited protein has been reported. On the other hand, cases with more than one pattern of LC deposition may be explained by the bicalon proliferative process with more than one pathogenic LC, causing damage. Again, only rare MM (2%) result in biclonal gammopathy with the production of two different heavy chains and/or light chains, with the combinations of IgG/IgM, IgA/IgG, k/ALC and IgD/IgM described in few reports, unfortunately no data concerning influence of coexistence of two and more patterns on renal outcome is available [22-30].

Here we present a case of IgD myeloma, manifested with the combination of LCDD and cast nephropathy.

Caucasian Male, 52 Years Old, Admitted July 14, 2015.

Main Complaints: Weakness, dizziness, breath shortage, loin pain.

Previous Medical History: In 1988 he was diagnosed with hepatitis B, no treatment, no follow-up for many years. In 2010, his nose basalioma was removed in the outpatient clinic, preoperative work-up was unremarkable.

History of Present Illness: March 2015 he developed unexplained high grade fever, spontaneously resolved a week later, since that time felt weakness, palpitations, dyspnea, and diffuse abdominal pain. His BP was 120/80mm Hg, HR 90bpm. Work-up in outpatient clinic (May 2015) showed Hb 13.1g/dL, Plt 222x10^9/L, ESR 14mm/h, creatinine 43.9 μmol/L (4.96mg/dL), urea 20.8mmol/L(125mg/dL) K+ 5.9mmol/L, AsAT 35U/L, AlAT 50U/L, GGTP 135U/L, AP 113U/L, proteinuria 1.03g/L with normal urinary sediment. Gastroscopy found 2 gastric and duodenal ulcers. He was taking omeprazole and beta-blockers without any improvement.

June 2015 he was admitted to the nephrology unit of local hospital with anemia (Hb 9.4 g/dL), thrombocytopenia (Plt 106x10^9/L) and progressive decline of kidney function with hyperkalemia and acidosis (creatinine 634 μmol/L [7.17md/dL], urea 26mmol/L [156mg/dL], uric acid 520μmol/L [8.7mg/dL], phosphate 2.2mmol/L, K+ 6.5mmol/L, pH 7.27, BE -12,9mmol/L). Proteinuria was still moderate (1.8% of plasma cells and 22.9% of lymphocytes (probably as peripheral blood admixture). Immunophenotyping of bone marrow cells found plasma cell phenotype CD45+dim, CD117-, CD20+.

Bone marrow aspiration in search for LPD was performed. Bone marrow smear showed 1.0% of plasma cells and 22.9% of lymphocytes (probably due to peripheral blood admixture). Immunophenotyping of bone marrow cells found plasma cell phenotype CD45+dim, CD38+[high expression], CD138+, CD56-, CD28+, CD200-, CD117-, CD20+.

He was suspected with rapidly-progressive GN, possibly cryoglobulinemic, and underwent kidney biopsy. According to the pathology report: Light Microscopy (LM) with standard staining’s showed 8 hypertrophied glomeruli with marked hypercellularity (capillary WBC infiltration); interstitial fibrosis with prominent lymphohistiocytic infiltration and tubular atrophy about 30%; massive hyaline casts in preserved tubular lumen; and oedematous arterial walls. Additional Congo red staining for amyloid was negative. Immune staining’s were negative for IgA, IgG, IgM, C3 and k LC, and strongly positive for λ LC – linear deposits along Glomerular Basement Membrane (GBM) and Tubular Basement Membrane (TBM) and also λ LC deposition in casts. Electron microscopy (EM) confirmed dense fine granular GBM and TBM deposits.

That was interpreted as the LCDD, and bone marrow aspiration in search for LPD was performed. Bone marrow smear showed 1.0% of plasma cells and 22.9% of lymphocytes (probably due to peripheral blood admixture). Immunophenotyping of bone marrow cells found plasma cell phenotype CD45+dim, CD38+[high expression], CD138+, CD56-, CD28+, CD200-, CD117-, CD20+.

It was referred to our clinic for the second opinion.

At Admission: Conscious, alert, slightly depressed. Body temperature 37.2°C, RR 18 per minute, pulse regular 98 per minute, BP 140/90mm Hg, Pale, well nourished, low extremities mild oedema. HEENT and neck otherwise normal. No palpable peripheral lymph nodes. Lungs: no dullness to percussion, any rhonchi, wheezes or rubs. Heart: regular rhythm, no murmur. Abdomen soft, non-tender, slightly tympanic, bowel sounds normal. Liver +2 cm below rib arch, non-painful, spleen and kidneys not felt. Urination is free, urine normally coloured, and urine output was about 1000ml/day.

Express Work-Up: Creatinine 1100μmol/L [12.4mg/dL], pH 7.25, K+ 5.8mmol/L, urine Bence-Jones protein positive. Ultrasound: significantly enlarged kidneys (148x66 and 142x68mm, parenchyma 25mm), hepatosplenomegaly, periportal lymphadenopathy, enlarged (12, 8 and 6mm) right axillary lymph nodes with coexistence of amyloidosis and compromised differentiation.
Diagnostics Considerations, Treatment and Further Work-Up: At that point we diagnosed AKI, and concluded that the cause of AKI could be nothing but cast nephropathy on top of LCDD in a patient with myeloma. Patient was urgently started with hemodialysis and normal saline infusions, skeletal X-ray, peripheral lymph nodes ultrasound, and bone marrow biopsy were performed, and immunochemistry results ordered from external lab, and kidney biopsy paraffin blocks were reprocessed and re-evaluated.

Skeletal X-ray: Did not found any lesions.

Immunohistochemistry: Showed traces of paraprotein D-λ and Bence-Jones-λ in serum, and urinary excretion of Bence-Jones-λ 1.22g/day. Cryoglobulins were not found IgG, IgA and κ/λ coefficients were below the normal range, CRP and β2-microglobulin significantly elevated.

Kidney biopsy re-processing and re-assessment (Figures 1-6): Sections of formalin fixed paraffin-embedded tissue were stained with Masson’s trichrome, periodic acid-Schiff, and Jones’ silver for LM. 12 glomeruli, slightly enlarged, with normal capillary wall, mild mesangial widening without mesangial or endocapillary proliferation. Total acute tubular necrosis with multiple large tubular casts, PAS- and Jones-negative, and fuxin-positive on Masson’s staining, surrounded by giant polynuclear cells. Severe tubulo-interstitial infiltration with lymphocytes, plasma cells and neutrophils, most prominent in zones of casts accumulation. Diffuse tubular atrophy and interstitial fibrosis about 40%. Arterioles and small arterial walls thickened due to muscular layer hypertrophy. Immunofluorescence on formalin fixed/paraffin embedded sections with FITC-conjugated anti IgA, IgG, IgM, C1q, C3, fibrinogen, λ and κ LC antibodies showed diffuse linear expression 4+ for λ LC along all basal membranes – glomerular, tubular, arteriolar and arterial, and also 4+ λ LC expression in the casts. All other immune stainings were negative. EM: toluidine-blue semi-thin sections showed 1 otherwise normal
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glomerulus, and tubule containing large cast with peripheral cell reaction. Electron microphotographs demonstrated diffuse fine granular intramembranous dense deposits in GBM and TBM. Conclusion: combined paraproteinemic nephropathy – cast-nephropathy plus LCDD.

Bone marrow biopsy: Revealed enhanced number of plasma cells by LM Immunohistochemistry found most plasma cells expressing CD138+, λ LC restriction, focal expression of CD56+, and few plasma cells expressing CD19 (aberrant immunophenotype of plasma cell population).

Final Diagnosis: Multiple myeloma IgDλ and BJλ, cast nephropathy combined with LCDD; AKI, treated with HD; anaemia, thrombocytopenia, hepatosplenomegaly, peripheral, mediastinal and periportal lymphadenopathy, secondary immunodeficiency.

Treatment and Follow-Up: Patient was seen by haematologist and referred to haematology unit for chemotherapy. At the latest evaluation (September 15, 2015) patient receiving standard chemotherapy per BCD (Bortezomib-Cyclophosphamide-Dexamethasone) protocol, doing well, but still on hemodialysis.

Conclusions

Presented case demonstrates characteristic features if IgD myeloma, like relatively young age, low grade IgD paraproteinemia, overproduction of LC with LC-proteinuria, lymphopenopathy and renal failure at presentation. Pathology findings with typical picture of LCDD were misleading, and only rapidly-progressive renal failure with kidney enlargement, not findings with typical picture of LCDD were misleading, and only finally gave a clue to the diagnosis. The pathology pattern of renal damage: a combination of cast nephropathy and LCDD confirms the leading role of cast-nephropathy in the clinical presentation with AKI, dominating over other symptoms. According to our experience such combined renal damage is rare, but not unique – in our cohort of 139 patients with LPD and renal damage, confirmed by pathology, we also have patients with AL amyloidosis and LCDD, and with combination of cast nephropathy, AL amyloidosis and LCDD. This case also illustrates the importance of renal damage pattern, typical for MM, for diagnostics of this disease in patients without clinical MM features.

References


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