Two cases of multiple myeloma, presenting with light-chain tubulopathy

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Abstract
Multiple myeloma (MM) is a plasma-cell dyscrasia presenting with generalized neoplastic changes in bones, accompanied by impaired haematopoiesis, susceptibility to infections, and end-organ damage. However, the clinical picture of myeloma might be quite different from the classic manifestation, patients can present with renal disease, dominating or precluding multiple myeloma features. Diagnosis and differential diagnosis demand renal tissue pathology evaluation, most often kidney pathology demonstrate cast-nephropathy, AL amyloidosis, and light-chain deposition disease. Light-chain proximal tubulopathy is less frequently reported variant of paraproteincenic kidney damage in patients mostly not demonstrating overt clinical features of multiple myeloma. This lesion is almost always induced by the excess of kappa light chains, excreted through the kidney and reabsorbed in the proximal tubule cells, which manifests with moderate CKD, and, rarely, with acquired adult Fanconi syndrome. We present two cases of myeloma with light-chain proximal tubulopathy. None of our two patients met multiple myeloma criteria at admission, and only kidney biopsy, which discovered light-chain proximal tubulopathy, prompted further work-up, in turn resulted in the diagnosis of multiple myeloma. Our cases demonstrate some peculiarities, making them rare even in the setting of light-chain proximal tubulopathy, which is rare itself. Case one is an extremely infrequent example of light chain lambda deposition in the proximal tubules, case two demonstrates acute kidney injury on the top of chronic kidney disease, unusual for the light-chain proximal tubulopathy. We conclude that kidney biopsy should be performed, unless contraindicated, in all patients with proteinuria and/or impaired kidney function of unknown origin; and that the chemotherapy with bortezomib-based regimens is effective for treatment of myeloma with light-chain proximal tubulopathy.

Introduction
Multiple myeloma (MM) is a plasma-cell dyscrasia presenting with generalized neoplastic changes in bones, accompanied by impaired haematopoiesis and susceptibility to infections. To distinguish MM from other plasma cell dyscrasias, the diagnosis is based on histologic, serologic, and radiographic features: bone marrow clonal plasma cells; monoclonal protein in the serum or urine; and end-organ damage, evidenced by renal impairment, hypercalcemia, anemia, or lytic bone lesions. Sometimes, however, the clinical picture of MM is quite different from the classic manifestation, which can cause diagnostic difficulties, thereby delaying treatment. Importantly, symptoms of kidney damage may dominate over MM symptoms, and even preclude overt MM, and the diagnosis of MM often results from the workup of unexplained renal disease. Renal disease clinical presentation in MM includes acute kidney injury (AKI), nephrotic syndrome, proteinuria and/or haematuria, arterial hypertension and chronic kidney disease (CKD), occasionally, patients with MM present with renal tubular dysfunction [1-8].

Renal involvement is mainly caused by the deposition of monoclonal immunoglobulin or fragments thereof, mostly light chains (LC), as organized (crystals, fibrils, microtubules) and non-organized deposits, involving all compartments of the renal parenchyma: glomeruli, tubules, interstitial space and vessels. Organized deposits induce cast-nephropathy, light-chain proximal tubulopathy (LCPT), AL amyloidosis, glomerulonephritis with organized microtubular monoclonal deposits, and cryoglobulinemic glomerulonephritis, while non-organized deposits lead to the light chain deposition disease or proliferative glomerulonephritis with monoclonal deposits of IgG/IgA. 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. Differential diagnostics is the major challenge, demanding pathology evaluation of kidney tissue, as the above mentioned lesions cannot be differentiated solely on the base of clinical features. Most often kidney pathology in patients with MM demonstrate cast-nephropathy, AL amyloidosis, and light-chain deposition disease, less frequently cryoglobulinemic glomerulonephritis proliferative glomerulonephritis can be seen [9-18].

LCPT is a unique entity which is less frequently reported as case reports and small case series. In LCPT the excessive LC, almost always of kappa type, are excreted through the kidney and reabsorbed in the proximal tubule cells, leading to tubular damage, sometimes resulting in the defects in acidification and concentration and, rarely, the features of complete acquired Fanconi syndrome (FS). Of interest, in many of the LCPT cases renal manifestations dominated in the clinical

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presentation, patients did not have typical features of MM, and the
diagnosis was predicated upon thorough immunopathology assessment
of renal biopsies material. Most of these patients present with moderate
CKD, slowly progressing towards ESRD and rarely – to symptomatic
myeloma, and according to the International Kidney and Monoclonal
Gammapathy Research Group (IKMGRG) definition regarded as
monoclonal gammapathy of renal significance (MGRS) cases [19-31].

As only very few series of LCPT have been published, the evidence
for the treatment efficasy is very weak, and the results of the so-called
novel anti-myeloma agents has not been evaluated in these patients.
According to the recommendations of the IKMGRG treatment choices
should be adapted to the degree of renal failure. Thus in patients with
stages 1 to 3 CKD, chemotherapy should be considered to try to slow
progression to ESRD, cyclophosphamide-, bortezomib-, or thalidomide-
based regimens are the best options [31]. Treatment of LCPT patients,
presenting with AKI is not described in the literature, however the study
of LC-induced AKI in the patients with cast-nephropathy demonstrated
substantial improvement of renal function, mainly in cases with less
severe renal impairment [32].

Here we present two cases of MM, manifested with kidney disease,
and diagnosed on the basis of immunopathology evaluation of renal
tissue.

**Case 1**

Caucasian male, 61 y.o., admitted to our unit January 18 2016.

**Main complaints:** general weakness, dizziness, back pain.

**Previous medical history:** inguinal hernia, prostate benign
hyperplasia, mild arterial hypertension, myocardial infarction in 2005,
benign colon neoplasm with laparoscopic sigmoid resection in 2010.

**History of present illness:** February 2015 during outpatient annual
work-up proteinuria 1.8 g/L was found. December 2015 proteinuria
increased to 5 g/L and he was referred to our clinic.

**At admission:** conscious, alert, oriented. Body temperature 36.7°C,
RR 17 per minute, pulse regular 78 per minute, BP 115/70 mm Hg.
Slightly obese (BMI 29). Skin clear, normally colored, no edema. Eyes
clear. HEENT and neck otherwise normal. Peripheral lymph nodes not
palpable. Joints: no swelling, movements not restricted. Vertebral and
flat bones percussion non painful. Lungs: clear . Heart : regular rate  and
rhythm, no murmur . Abdomen soft, non-tender, bowel sounds normal.
Liver, spleen and kidneys not felt. Urine output 1500 ml/24 hours.

**Work-up:** total blood count, blood chemistry, serum osmolality and
pH, serum electrophoresis, serum immunoglobulinis and coagulation
tests - within normal range, infectious screening negative. Urine specific
gravity 1017-1030, urine pH 5.5, proteinuria 5.8 g/24 hours, glycosuria
not detected, urinary sediment unremarkable. Electrocardiogram -
post-infarction cicatricle changes. Plain chest X-ray otherwise normal.

**Kidney biopsy**

**Light microscopy:** sections of formalin fixed paraffin-embedded
tissue were stained with H&E, Masson’s trichrome, periodic acid-Shiff
and Congo red. 43 glomeruli, 4 out of them totally sclerosed. The rest
appear otherwise normal, without proliferative changes, capillary walls
thin, single-contour (Figure 1). Mild interstitial fibrosis and tubular
atrophy, up to 10% of parenchyma, diffuse-focal interstitial infiltration
with inflammatory cells in the areas of interstitial fibrosis. Tubular
chain tubulopathy with focal global glomerulosclerosis (CKD stage 2), secondary immunodeficiency

Treatment and follow-up: patient was referred to the hematology unit for chemotherapy and received 6 courses of bortezomib-based chemotherapy. At the latest follow-up visit in August 2016 he was doing well, his Bence-Jones lambda protein excretion decreased to 0.9 g/24 hours, serum creatinine remained normal.

Case 2

Caucasian female, 57 y.o., admitted to our unit April 1 2016

Main complains: general weakness, fatigue.

Previous medical history: tonsillectomy; tuberculosis as a child, received specific therapy and never recurred; cystitis; two uncomplicated pregnancies; uterine fibroid, severe menstrual pains for many years, for which she was regularly taking pain-killers, metrectomy in 2007.

History of present illness: in 2007 she developed pedal edema, outpatient work-up showed mild proteinuria and serum creatinine 150 µmol/L, which was considered as analgesic nephropathy and urinary tract infection. After metrectomy she stopped taking pain-killers, edema resolved spontaneously.

December 2014 she complained for back pain, outpatient work-up revealed serum creatinine 169 µmol/L, total calcium and urinalysis were normal. Serum electrophoresis showed total protein 90 g/l, M-spike in gamma zone. Other blood chemistry tests, serum osmolality and pH within normal range. IgA and IgM within normal range, IgG 37.4 g/L. Infectious screening negative, coagulation tests otherwise normal. Urine specific gravity 1010-1011, urine pH 5.0, proteinuria 0.5-0.01 g/24 hours, glycosuria not detected, urinary sediment unremarkable. Electrocardiogram and plain chest X-ray unremarkable. Kidneys and abdomen ultrasound – nothing but mild hepatomegaly.

Kidney biopsy

Light microscopy: sections of formalin fixed paraffin-embedded tissue were stained with H&E, Masson's trichrome, periodic acid-Shiff and Congo red. 20 glomeruli, 4 out of them totally sclerosed (Figure 4). The rest appear enlarged, without any proliferative changes, capillary walls thin, single-contour (Figure 5). Mild interstitial fibrosis and tubular atrophy, up to 15% of parenchyma, diffuse-focal interstitial infiltration with inflammatory cells in the areas of interstitial fibrosis. Few tubules with signs of epithelial dystrophy without any casts in the tubular lumen (Figure 6). Arteries and arterioles otherwise normal. Congo red staining was negative.

Immunofluorescence on unfixed cryo-sections: with fluorescein
Final diagnosis: multiple myeloma IgG kappa and Bence-Jones kappa, light-chain tubulopathy with focal global glomerulosclerosis, AKI on the top of CKD 2, secondary immunodeficiency. Hematological and organ remission after 4 courses of chemotherapy.

Discussion

In both cases typical clinical features of MM, like ossalgia, anemia, hypercalcemia and bone lytic lesions were absent, patients presented with kidney disease, and only kidney biopsy immunopathology study gave a clue to the diagnosis. This is in agreement with the literature data, suggesting that the renal manifestations, mostly with moderate CKD and not obligatory complete FS, dominated as the clinical presentation in the majority of LCPT cases [19,24-26,29,30]. However, our cases demonstrate some peculiarities, making them rare even in the setting of LCPT, which is rare itself.

In the first case presented with unexplained new onset of severe proteinuria, routine work-up was not suggestive for myeloma: his total blood count, serum calcium, creatinine and total protein were normal, and serum electrophoresis did not show M-spike. Thus he could be suspected with non-glomerular proteinuria only because of the combination of massive proteinuria and normal serum albumin level. Therefore, kidney biopsy was performed to unmask the cause of proteinuria, and found tubular atrophy with LC lambda expression within tubular epithelial cells and very mild and unspecific focal glomerulosclerosis. That lead to the diagnosis of LCPT, and given the lack of MM signs and symptoms we considered the patient as MGRS case rather than MM. Surprisingly, his bone marrow smear showed 63.5% of plasma cells, and immunochemistry detected massive Bence-Jones protein lambda almost completely excreted in urine, which explained his normal serum protein and the absence of M-spike on standard serum electrophoresis. Severe plasma cell bone marrow infiltration, significant Bence-Jones protein secretion with biopsy-proven end-organ damage, and secondary immunodeficiency altogether fulfilled the criteria of MM. The absence of bone lytic lesions and anemia, as well as the absence of FS clinical and laboratory features in this case

Pathologists conclusion: Focal global glomerulosclerosis (20%), immunofluorescence data suggestive for light-chain tubulopathy.

Additional work-up

Bone marrow biopsy from March 2016 was re-evaluated: Immunohistochemistry with CD3, CD19, CD20, CD56, CD138, Cyclin D1, IgA, IgG, lambda and kappa light chains and was ordered and found large number of mature plasma cells CD138+, located inter- and para-trabecular discretely and in clusters up to 10 cells. These cells were kappa-positive and express CD56 (membrane reaction) and IgG (cytoplasmic reaction). Small B-cells (CD19+, CD20+) and small T-cells (CD3+) are dispersed. Histocytes were Cyclin D1-positive.

Clinical diagnosis: Multiple myeloma IgG kappa and Bence-Jones kappa, light-chain tubulopathy with focal global glomerulosclerosis (CKD stage 3b), secondary immunodeficiency

Treatment and follow-up: Patient received 4 courses of bortezomib-dexamethasone chemotherapy in our unit. At the latest evaluation October 2016 she is doing well, her serum creatinine decreased from 213 to 98 µmol/L, and immunochemistry showed the decrease of IgG-kappa secretion from 24.6 to 7.1 g/L (Figures 6 and 7), Bence-Jonce protein not detected neither in serum nor in urine. Hypogammaglobulinemia persisted.
might be explained by the early diagnosis. The most unusual in the case 1 is the type of LC, as the association of the LCPT with lambda LC is exceptionally rare described in the literature so far.

The second case presented with 9-years history of moderate, slowly progressive renal dysfunction without significant proteinuria. Already 1.5 years before the kidney biopsy she was found with elevated total serum protein and M-spike, confirmed by detection of significant IgG kappa and mild Bence-Jones protein kappa secretion without urinary excretion of the latter. As far as she had no bone lytic lesions, anemia and hypercalcaemia, and her bone marrow biopsy didn’t find plasma cell infiltration, MM diagnosis was not proved, even though she had monoclonal gammapathy, impaired kidney function and immunodeficiency. She was suspected with MGRS, and kidney biopsy indeed revealed paraprotein-induced renal damage with LCPT and focal glomerulosclerosis. To differentiate MGRS and MM we ordered bone marrow immunohistochemistry, which provided the evidence for MM. Given FS clinical features were absent we regarded her longstanding kidney function impairment as a result of glomerulosclerosis of 20% of glomeruli secondary to the tubular damage. However, her kidney function completely recovered under chemotherapy, meaning that she had AKI on the top of her CKD, which is most unusual. Our interpretation is that the patient had a long course of smoldering myeloma with LCPT and recent evolution towards MM with exacerbation of kidney damage. The latter is suggestive for cast-nephropathy rather than for LCPT, but tubular casts were not found in her biopsy. There were no other reasons for AKI beyond LCPT, as she was not taking analgesics for many years before biopsy, didn’t receive contrast-media, antibiotics or any other nephrotoxic agents and had no signs of volume depletion.

Finally, in both cases treatment with the bortezomib-based chemotherapy regimens was successful and lead to hematologic and organ remission.

Conclusions

Light-chain proximal tubulopathy is an infrequent variant of paraproteinemnic kidney damage in patients mostly not demonstrating clinical features of multiple myeloma. This lesion is almost always induced by kappa light chains, presenting with moderate CKD, and, rarely, with Fanconi syndrome. None of our two patients met multiple myeloma criteria at admission, and only kidney biopsy, which discovered light-chain proximal tubulopathy, prompted further work-up in turn resulted in the diagnosis of multiple myeloma. Case one is an extremely rare example of light chain lambda deposition in the proximal tubules, case two demonstrates acute kidney injury on the top of chronic kidney disease, unusual for the light-chain proximal tubulopathy. We conclude that kidney biopsy should be performed, unless contraindicated, in all patients with proteinuria and/or impaired kidney function of unknown origin; and that the chemotherapy with bortezomib-based regimens is effective for treatment of myeloma with light-chain proximal tubulopathy.

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