Polyarteriitis Nodosa with Real Arteries and Vein Involvement and Urological Manifestations: Case Report

Abstract
Current Nomenclature defines polyarteriitis nodosa (PAN) as a medium vessels vasculitis - a necrotizing arteritis of medium or small arteries without glomerulonephritis or vasculitis in arterioles, capillaries, or venules; and not associated with ANCA. PAN typically affects skin, joints, peripheral nervous system, digestive tract and kidneys. Here we present a case of PAN with extremely severe combined kidney damage: renal artery aneurism and occlusion with peri-vasal hematoma, multiple kidney infarctions, renal vein thrombosis, necrotizing ureteritis with periureteritis, with left orchitis and with skin and joints involvement, successfully treated by nephrectomy and immunosuppression.

Keywords: Necrotizing vasculitis; Renal artery aneurism; Kidney infarction; Renal vein thrombosis; Ureteritis; Orchitis

Introduction
Current Nomenclature defines polyarteriitis nodosa (PAN) as a medium vessels vasculitis - a necrotizing arteritis of medium or small arteries without glomerulonephritis or vasculitis in arterioles, capillaries, or venules; and not associated with ANCA [1]. PAN typically affects skin, joints, peripheral nervous system, digestive tract and kidneys [2]. Here we present a case of PAN with extremely severe combined kidney damage.

Case presentation
28 years old Caucasian man, admitted May 31 2017.

Main complaints
General weakness, fever, back pain, muscle pain, weight loss.
was suspected with left testicle neoplasm; however his LH, FSH, estradiol, progesterone, prolactin, testosterone, ACTH, cortisol, aldosterone, AFP, PSA and free beta HCG were all normal. Antiviral treatment was discontinued, and he underwent contrast-enhanced abdomen CT and CT-angiography, which showed pseudo-aneurism of the left artery, left artery proximal occlusion and reduced left kidney excretory function (Figure 1 & 2). Eleven months after his first presentation he was referred to our hospital and admitted to the vascular surgery.

**Work up**

**TBC:** Hb 7.1 g/L, WBC 14.5x10⁹/L, PLT 612x10⁹/L, ESR 52 mm/h.

**Blood chemistry:** Creatinine 163 µmol/L, urea 13.1 mmol/L, uric acid 378 µmol/L, total protein 81 g/L, serum iron 10.7 µmol/L; CRP 165 mg/L; electrolytes, glucose, albumin, cholesterol, bilirubin, liver enzymes - within normal range.

**Urinalysis:** SG 1020, protein 1.0 g/L, RBC 3-4, WBC 75-80 hpf.

**Infections screening:** RPR-test for Treponema Pallidum, HBSAg, anti-HCV and anti-HIV-antibodies negative, tuberculin skin test - all negative.

**ECG; plain chest X-Ray and ECHO-CG:** Unremarkable.

**Abdomen and kidneys ultrasound:** Liver slightly enlarged; pancreas and spleen otherwise normal. Right kidney 110x45 mm, with normal parenchyma 17 mm. Left kidney 84x50 mm, with hyperechoic parenchyma 14 mm. Space-occupying mass in the projection of the main stem of the left renal artery - most likely renal artery aneurism with aneurism thrombosis and paravascular hematoma. Pelvis not dilated.

**Urologist consults:** Left-side epididymitis and orchitis.

**Neurologist consults:** No signs or symptoms of any neurologic disease.

**Diagnostic considerations and treatment:** At that point left nephrectomy was considered and nephrologist was invited to see the patient. Given the combination of weight loss, orchitis, myalgia/arthritis, arterial hypertension, elevation of serum creatinine and renal artery aneurism, the patient was diagnosed with polyarteritis nodosa. June 2 2017 nephrectomy was performed.

**Excised kidney pathology:** Light microscopy sections of formalin fixed paraffin embedded tissue were stained with H&E, PAS and Masson’s trichrome.

**Renal artery:** Figures 3&4 Fragment of artery of elastic type, severely thickened due to irregular sclerosis and hyalinosis with impaired histoarchitectonics of the wall layers and integrity. Internal elastic membrane segmentally preserved. Severe subendothelial fibrosis with focal lympho-histiocytic infiltration and hemosiderophages clusters. Vessel lumen and internal surface of the vessel are covered with thrombotic masses with secondary changes, presented by organization, cell infiltration and recanalization (neo-formed vessels), and fresh thrombotic masses without cell reaction. Perivascular lymph node with prominent sclerosis, and lymphoid tissue hypoplasia presented by few lymphoid follicles with attenuated centers. Muscular type arteries changes are the same as in the renal tissue and ureter wall arteries.

**Kidney:** Arterioles and small arteries walls otherwise normal. Middle arteries walls are diffuse thickened due to the fibrous connective tissue with impaired histoarchitectonics of the wall layers and subtotal lumen obstruction. Large arteries walls are thickened due to edema, massive transmural and perivascular infiltration with neutrophils and histiocytes, and irregular fibrous connective tissue with impaired histoarchitectonics of the wall layers. Multiple pseudo-aneurisms in the zones of large arteries

**At admission**

Alert, oriented. Body temperature 37.8° C, RR 19 per minute, pulse regular 92 per minute, BP 160/95 mm Hg. Undernourished. Skin pale, moist, with few slightly painful nodular erythematous eruptions on the arms and shoulders. No peripheral oedema. HEENT and neck otherwise normal. No palpable peripheral lymph nodes. Joints: no swelling, ROM normal. Lungs: clear. Heart: rhythm regular, no murmurs. Abdomen soft, slightly tender in the upper left quadrant, bowel sounds normal. Liver +2 cm below costal margin, non-painful, spleen and kidneys not felt. Urine output 850 mL/day, urine normally coloured.
bifurcation with the severe edema and prominent walls infiltration. Multiple cortical macrofocal ischemic and hemorrhagic renal tissue necrosis. Preserved renal tissue presented by normal glomeruli. Total acute tubulonecrosis without tubular atrophy or interstitial fibrosis (Figure 5 & 6).

Ureter: Erosive-ulcerous lesions of mucosa with massive fibrino-leucocyte excaudate. Ureter wall is thickened due to the severe edema and massive leucocyte infiltration of all layers; with multiple intramural necrosis and prominent neutrophil karyorhexis. Small, medium and large arteries walls are thickened due to the irregular neutrophil infiltration, with circular fibrinoid necrosis of small and medium arteries walls; and focal impaired histoarchitecture and integrity of large arteries walls due to severe infiltration; with multiple pseudo-aneurism formation. Diffuse severe fibrous periureteritis with massive fibrin deposition and leucocyte infiltration with prominent karyorhexis (Figure 7-9).

Renal vein: With total obstruction of the lumen by the sludged thrombi.

Pathologists conclusion: Polyarteritis nodosa - diffuse segmental transmural necrotizing and sclerosing vasculitis with renal artery, small, medium and large intrarenal and ureter wall arteries, with pseudo-aneurism formation and thrombosis; macrofocal ischemic and hemorrhagic renal infarctions; renal vein thrombosis. Phlegmonous-ulcerous ureteritis, fibrinous periureteritis.
Final diagnosis and further treatment: Pathology evaluation confirmed the diagnosis of polyarteritis nodosa. Within 2 weeks after the surgery patient was moved to nephrology and started on oral prednisone 50 mg/day and cyclophosphamide i.v. “pulses” 500 mg every 2 weeks.

Treatment results: At the last evaluation after 6 weeks of treatment patient is doing well. His body temperature is normal, BP is 130/80 mm Hg, his skin lesions resolved, and he gained 2 kilos of weight. Work-up showed Hb 13.7 g/dL, WBC 10.5x10^9/L, Plt 264x10^9/L, ESR 17 mm/hour, CRP 1.0 mg/L, serum creatinine 109 µmol/L, his urinalysis are normal (proteinuria and microhematuria not detected). He already received 1500 mg of cyclophosphamide and continued on cyclophosphamide “pulses” 500 mg every other week and oral prednisone 45 mg/day with slow tapering to 30 mg/day.

Discussion

Our patient presented with fever, erythema nodosum, muscle and joint pain, orchitis, weight loss, elevated ESR and CRP, thrombocytosis, leukocytosis, anemia, proteinuria, arterial hypertension, impaired kidney function, renal artery pseudoaneurism and occlusion, and peri-renal mass. Matching clinical presentation over ACR 1990 criteria [3] he had 6 out of 10 most characteristic disease features:

i. Weight loss of 4 kg or more - present
ii. Livedo reticularis - absent
iii. Testicular pain/tenderness - present
iv. Myalgia or leg weakness/tenderness - present
v. Mononeuropathy or polyneuropathy - absent
vi. Diastolic blood pressure greater than 90 mm/Hg - present
vii. Elevated blood urea nitrogen or creatinine level unrelated to dehydration or obstruction - present
viii. Presence of hepatitis B surface antigen or antibody in serum - absent
ix. Arteriogram demonstrating aneurysms or occlusions of the visceral arteries - present
x. Biopsy of small- or medium-sized artery containing polymorphonuclear neutrophils - not evaluated at the time of diagnosis

According to the ACR criteria, the presence of at least 3 of these features is required for classification of disease as PAN with 82% sensitivity and 87% specificity rate; therefore, the diagnosis of PAN in our patient was certain. In addition, fever, skin nodules, arthralgia’s, elevated ESR and CRP, thrombocytosis, leukocytosis...
and moderate proteinuria, which our patient demonstrated, are quite characteristic, even though non-specific features of PAN [4]. Moreover, male gender also matching with typical for PAN male-to-female ratio, which is 1.6-2.1 [5]. The absence of hepatitis B, which is known to be strongly related with PAN, does not exclude the diagnosis. It was estimated that hepatitis B is causative for 30% and more recently - for 7-10% of PAN cases [6,7]. Given that, he met ACR criteria for PAN even before the kidney pathology became available.

Renal manifestations in our patient were quite severe. Proteinuria and serum creatinine > 1.58 mg/dL (139.7 µmol/L), which both he demonstrated at presentation, are 2 out of 5 negative prognostic factors for PAN; presence of 2 and more factors is related to 45.9% 5-years mortality according to the prognostic Five Factor Score (FFS) [8]. Aneurismatic ruptures are one of the most serious complications of PAN, and may occur in many involved organs, including kidneys; in such cases aneurismatic ruptures are causative for peri-renal hemorrhage. The typical clinical presentation is a loin pain, significant Hb fall, hypotension and a peri-renal mass, which was also found in our patient. In some cases surgical evacuation of the hematoma or partial/complete nephrectomy is necessary [9-11].

In general, histology in PAN reveals a focal necrotizing arteritis of mixed cellular infiltrate within the vessel wall. Necrotizing arteritis begins as a segmental necrotizing inflammation of arteries with conspicuous infiltration of neutrophils and monocytes. This typically induces fibrinoid necrosis characterized by accumulation of plasma proteins in injured tissue, including coagulation factors that are converted to fibrin [1]. Kidney pathology data for PAN are scarce; kidney biopsy is performed only in exceptional cases due to the risk of aneurysmal rupture and bleeding [12]. However, there are few reports, confirming renal involvement with typical necrotizing vasculitis in patients with PAN [13,14]. We also found some old papers, describing single patients with PAN and renal vein thrombosis [15,16]. Possible explanation is that obliteration of the arterioles that supply the walls of the veins may be causative for the venous thrombosis.

Ureteric stricture, along with orchitis and peri-renal hemorrhage is listed among urogenital complications of PAN [1,11]. There are several case reports, describing ureteral strictures in PAN, always as unusual presentation of the disease [17-25]; thus, phlegmonous-ulcerous ureteritis with fibroin perireteritis found in our patient represent a very rare type of lesion. In our case the pathology findings, demonstrating the specific damage of renal artery, intra-renal vasculature, renal vein and ureter, were confirmative for PAN. With those findings our patient had altogether 7 out of 10 characteristic features of PAN. No crescents or any other kind of glomerular lesions were found, which, along with negative ANCA, completely ruled out the diagnosis of ANCA-associated vasculitis.

Treatment options for PAN include corticosteroids and immunosuppressants. Clinical trials suggest that corticosteroids alone are adequate first line therapy for patients with PAN with a FFS of zero, however 40% of them may require additional immunosuppression. Patients with a FFS ≥ 1 should receive intermittent intravenous cyclophosphamide in addition to corticosteroids [26,27]. In the steroid-refractory and recurrent PAN rituximab and infliximab or plasma exchanges has been used occasionally [28-30]. As our patient had FFS 2 (proteinuria and elevated serum creatinine), after nephrectomy, which was indicated given the combination of renal artery aneurism and stenosis with peri-vascular hematomata, kidney shrinking and absence of kidney function, he was started on the combination of prednizone and cyclophosphamide. Almost 2 months after nephrectomy and 6 weeks under the immunosuppression all his symptoms resolved, and he is continued on the immunosuppression with intermittent i.v. cyclophosphamide and slow tapering of the oral prednisone dosage. We conclude that this case represent a rare combination of PAN manifestations, demanding multidisciplinary approach. Indeed, he had renal artery aneurism and occlusion with peri-vascular hematomata, multiple kidney infarctions, renal vein thrombosis, necrotizing ureteritis with periureteritis, and orchitis, and a team including radiologist, vascular surgeon, urologist, nephrologist and nephropathologist participated in the diagnostics and treatment.

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