A Rare Case of Hydronephrosis Secondary to an Intra-abdominal Haematoma in a Patient with Marfan’s Syndrome

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Abstract
A patient with Marfan’s Syndrome and type B aortic dissection presented to an academic teaching hospital with sudden onset left loin to groin pain and haemodynamic instability. Her INR was raised at 9.3 and haemoglobin had dropped from 126 to 106 g/L. CT abdomen revealed unilateral left hydronephrosis secondary to ureteric compression by an 11.8×7.0×8.6 cm retroperitoneal haematoma originating from the left iliacus. The haematoma was managed conservatively with transfusion of fresh frozen plasma and strict bed rest. A left retrograde uretric JJ stent was placed without complication and she was discharged from hospital eight days after presentation.

Keywords: Hydronephrosis; Marfan’s; Iliopsoas rupture; Haematoma

Case Report
A 56 year-old lady presented with a 24 hour history of sudden onset intense burning left loin to groin pain radiating down her left leg, with associated nausea and vomiting. She was passing good amounts urine with no urinary symptoms of note. Bowel habit was regular. She was known to have Marfan’s syndrome, and a complex history of cardiothoracic surgery including a type A aneurysm repair and aortic arch repair 25 years previously; mechanical aortic valve replacement requiring warfarinisation 5 years previously; and a type B aortic aneurysm requiring distal aortic arch and descending thoracic aorta replacement 2 years previously.

On examination she had left loin tenderness with no peritonism and no masses palpable. All peripheral pulses were present and equal. Chest was clear. Blood pressure was 114/71, heart rate 80, regular and no masses palpable. All peripheral pulses were present and equal.

Urinalysis showed only trace of blood.

Nitrites, protein and leucocytes were negative. Her INR was raised at 9.3 and haemoglobin 12.6 g/L. White cell count was 20.2×10^9/L. Creatinine was 55 µmol/L, urea 3.4 mmol/L. A repeat blood test after 4 hours showed a drop in haemoglobin to 106 g/L.

A leaking aortic dissection was initially suspected and CT aortic angiogram performed. In the left retroperitoneum extending down to the left groin around the common femoral and superficial femoral artery, a large heterogenous mass was visualised, consistent with acute haematoma. Dilatation of the left renal pelvis and upper ureter was seen related to the left retroperitoneal haematoma.

The patient was initially transferred to a high dependency unit for strict monitoring of blood pressure, haemoglobin and renal function. Her raised INR was reversed with three units of fresh frozen plasma.

Following discussion at uro-radiology multi-disciplinary team meeting it was decided that the haematoma was caused by acute bleeding from a blood vessel within the iliopsoas muscle with secondary ureteric impingement, resulting in hydronephrosis (Figure 1). There was no concomitant tracking of haematoma from previous aortic dissection (Figures 2 and 3).

Decision was made to manage the haematoma conservatively. Once her INR was adequately reduced to 1.5, a left retrograde uretric double J stent was placed without complication. The patient was discharged eight days following admission. The patient remained asymptomatic with no flank pain, indicative of non expanding haematoma. Haemoglobin and renal function were measured and continued to be within normal limits. Due to multiple CT scans on admission it was decided that follow-up CT scan would be delayed until 6 months to minimise further exposure to radiation.

Discussion
Marfan’s syndrome is an autosomal dominant connective tissue disorder with many associations including aortic aneurysm or dissection, aortic or mitral valve prolapse and spontaneous pneumothorax. Marfan’s syndrome is characterised by mutations in gene FBN-1. Glycoprotein fibrillin-1 is encoded by FBN-1 and contributes to formation of connective protein elastin and extracellular matrix. Tissue in which elastin is abundant such as blood vessels are commonly affected by Marfan’s syndrome due to altered elasticity, resulting in aneurysm and rupture.

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Diagnosis should be made by combining clinical correlation with CT radiological imaging. Causes of intrabdominal haematoma include retroperitoneal haematoma should include rectus sheath haematoma, inferior epigastric artery rupture and iliopsoas rupture. Risk factors for intrabdominal haematoma include anticoagulation, coagulation disorders, acute increase in intrabdominal pressure, trauma and invasive procedures [6].

Management of this condition involves correcting any haemodynamic instability, followed by reversal of anticoagulation or treatment of bleeding diatheses. Once this is accomplished, decompression of the obstructed urinary system is necessary. The haematoma may be treated conservatively by regular monitoring for improvement in size, or surgically via evacuation of the clot [5,7].

**Conclusion**

Intra-abdominal haematomas causing hydronephrosis are very uncommon and more so in Marfan’s Syndrome. Due to the higher incidence of aortic aneurysms in Marfan’s Syndrome, leaking or ruptured aneurysms are important causes of intra-abdominal haematomas that need to be rapidly diagnosed and treated. A conservative approach to management of iliopsoas haematomas is highly effective in combination with double J stenting to relieve hydronephrosis.

**References**