Abstract

Introduction: Retroperitoneal Fibrosis (RPF) is a clinicopathological condition characterized by inflammatory fibrotic reaction around infra renal aorta, iliac vessels and surrounding retroperitoneum with myriad presentations [1].

Major advancements have been made since then towards understanding the disease process, though lack of definite diagnostic criteria and treatment protocol still remains a challenge.

RPF has estimated incidence of 1-2 per 100,000 [2]. Given the rarity of the condition the diagnosis may often be missed if not included in differential diagnosis in appropriate clinical background. It is imperative to identify and treat RPF early on in its course in order to avoid potentially devastating complications including Deep Vein Thrombosis, Pulmonary Embolism, reno-vascular hypertension, ureteric obstruction, Chronic Kidney Disease (CKD) [3]. Here we describe a case of RPF in a patient on chronic beta blocker therapy who later presented with unilateral DVT and PE. Our objectives are to consider RPF as one of the potential complication of chronic beta-blocker therapy and to recognize DVT/PE as a potential complication of RPF.

Case Presentation

A 62-year-old Caucasian male with history of hypertension on chronic beta-blocker therapy for 2 years (Metoprolol 50 mg twice daily) presented with sub-acute left sided non-radiating lower back pain of 5-day duration. Computerized Tomography (CT) scan with contrast of the abdomen and pelvis revealed large segment of inflammatory stranding involving the periaortic retroperitoneum extending from the level of the kidneys upto the pelvis and incasing the left iliac veins. CT guided retroperitoneal core biopsy was done which was consistent with retroperitoneal fibrosis. Two months later, patient presented with complain of acute onset shortness of breath and increasing lower extremity edema, on the left side. Electrocardiogram was consistent with a finding of new onset Atrial fibrillation (A fib), Ultra Sono Gram (USG) Doppler study of lower extremities unveiled occlusive left sided DVT of the popliteal vein. A CT Angiography demonstrated segmental and sub-segmental pulmonary emboli of the right lower lobe with no evident pulmonary edema. He was treated with steroids, digoxin and warfarin. On follow up a repeat CT scan three months later of the abdomen and pelvis showed stable retroperitoneal mass with no further progression.

Conclusion: Our patient presented with lower extremity edema and imaging revealed extension of RPF to involve common iliac vessels. With beta-blockers as a possible inciting event, RPF causing venous stasis, iliac vein compression and thus DVT/PE is the most plausible explanation. This case reports add to the medical literature how DVT/PE can be cause by an underlying disease entity not related to the usual causes and if not worked up patients may be labeled as having unprovoked events. Any relationship between beta blockers and RPF is questionable and has not been proven in any randomized trials, but should be thought of by the physician if such clinical situation is encountered.
diagnosis included mesenteric infarct and lymphoma, and metastatic mass. Magnetic Resonance Imaging (MRI) showed abnormal soft tissue mass in left para-aortic region without any particular signal characteristics to help differentiation of the pathology. CT guided retroperitoneal core biopsy was done, which showed fibrofatty tissue admixed with lymphoplasmacytic infiltrate with scattered lymphoid aggregates accompanied by infiltration of eosinophils and numerous foamy macrophages. Immunohistochemistry demonstrated normal distribution of CD20-positive B-cells and CD3-positive B-cells without coexpression of CD5. A diagnosis of RPF was made given the clinical, imaging and pathological presentation. Patient was started on prednisone 40 mg daily. His beta-blocker therapy was stopped.

Two months later, patient presented with complain of acute onset shortness of breath and increasing lower extremity edema, on the left side. There was no history of trauma, long travel, infection or any other immediate precipitating event. Electrocardiogram was consistent with a finding of new onset Atrial fibrillation (A fib), which should be considered a red flag for possible new onset PE. Ultra Sound Gram (USG) Doppler study of lower extremities unveiled occlusive left sided DVT of the popliteal vein. A CT angiography demonstrated segmental and sub-segmental pulmonary embolism of the right lower lobe with no evident pulmonary edema (Figure 2). Coagulation panel was within normal range. Patient was treated accordingly for PE/DVT with heparin which was eventually bridged to warfarin and A fib with digoxin. His condition improved and he was discharged after 5 days. Patient is being followed up and a repeat CT scan three months later of the Abdomen and pelvis showed stable retroperitoneal mass with no further progression.

Discussion

RPF most commonly presents with dull, aching low back or flank pain. Systemic features of inflammation like weight loss, anorexia, fever, easy fatigability are common presenting symptoms. Patients can also present with complications secondary to local effects like new onset reno-vascular hypertension, lower extremity edema, hydrocoele, DVT [4]. Idiopathic cases account for nearly two third of the presentations.

Beta-blockers, including metoprolol and eye drops containing timolol have been reported to associate with RPF [5] others include Ergot derivatives [6], hydralazine [7], bromocriptine [8] and methyldopa [9], pergolide [10]. Certain malignancies like lymphomas, gastric and pancreatic cancers have been found to be associated with RPF [11]. Radiation therapy [3] has been found to be a significant risk factor associated with RPF. Certain infections like tuberculosis [12] actinomyces [13] and schistosomiasis [14] have been reported as causative factors of RPF. Previous trauma in form of surgery has also been reported [15].

Although the etiopathogenesis is poorly understood, recent research studies points to the role of autoimmune mechanisms. Hypothesis proposed include excessive immune reaction in response to steroid and oxidized LDL molecules in local aortic atherosclerotic process [16]. Role of IgG4 bearing plasma cell is also proposed [17]. RPF has been described in association certain autoimmune diseases like thyroiditis and association with HLA-DRB1*03 allele has been described [1].

Investigators have proposed a staging system based on the extent of the disease on imaging. Stage I RPF has fibrosis limited to infra-renal aorta or iliac vessels; stage II involves progression of fibrosis to inferior vena cava. Stage III further lateral extension of fibrosis occurs causing compression of the ureters and in stage IV the renal vessels are also involved. The same group of investigators has also proposed diagnostic criteria for idiopathic RPF with absence of any aneurismal dilatation of the aorta or absence of any mass in pelvic and abdominal cavity, in association with age specific cancer screening with normal outcomes [3].

Imaging remains the mainstay in diagnosing RPF. CT and MRI are most preferred modalities to identify and stage the disease process [18]. On MRI, RPF appears hypointense on T1 weighted images and hyperdense on T2 weighted images [1].

Fluorodeoxyglucose Positron Emission Technology (PET) is a non-invasive, safe, whole body imaging method with potential to identify other diseased sites, any associated autoimmune or neoplastic disorder and also can be used for surveillance of disease activity [19]. Biopsy can be done to identify histopathology of the disease [20]. It is generally required when infection or malignancy is strongly suspected, or location of lesion is atypical raising doubt about the diagnosis or surgical intervention is being planned.

Treatment of RPF can include both medical and surgical measures depending on the presentation. Stopping any possible inciting agent is the first step. Our patient had been on chronic beta-blocker...
therapy. It was stopped and in conjunction with steroid therapy disease process was stable on follow up after three months. Steroids are first line therapy if there is no evidence of any mass effect such as compression of surrounding structure, vascular extension into the surrounding veins and arteries, tubular compression like ureters or organ function compromise. Steroid resistant disease can be treated by other immunosuppressive agents including cyclophosphamide, methotrexate, mycophenolate mofetil [21] colchicine [22] and tamoxifen [23].

Follow up of disease activity can be done by serial imaging including contrast enhanced CT, MRI and PET. Measuring Serum Ig4 level [24] ESR/CRP [25] and renal function periodically may also be helpful, though the validity of these studies for following up of disease activity remains to be validated in larger studies. If the disease doesn’t respond to these measures, diagnosis should be reconsidered and other causes must be ruled out. Complication like ureteral entrapment and consequent renal dysfunction is one of the most recognized complications of RPF [3]. Studies have shown that incidence of DVT and PE is also increased in patients with RPF and prophylactic anticoagulation should be considered on individual case by case basis in such patients with evidence of inferior vena cava and/or iliac vessel involvement causing venous stasis and lower extremity edema [3]. To our knowledge there have been no randomized trials evaluating the duration of treatment and prophylactic use of anticoagulation in this patient population. A strong argument can be made regarding more frequent surveillance i.e every 3 to 6 months, by different imaging modalities to access the progression of the disease instead of subjecting the patient to risks of anticoagulation including but not limited to hemorrhagic complications.

Conclusion

Our patient presented with lower extremity edema and imaging revealed extension of RPF to involve common iliac vessels. With beta-blockers as a possible inciting event, RPF causing venous stasis, iliac vein compression and thus DVT/PE is the most plausible explanation. A-fib is a very important symptom of PE especially in patients who had their beta blockers discontinued recently.

References


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