Radiological aspects of retroperitoneal fibrosis

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Abstract

Objective: Evaluate radiological findings on retroperitoneal fibrosis.

Patients and methods: A retrospective analysis conducted in the internal Medicine Department in Sahloul teaching Hospital between 1999 and 2016.

Results: Thirty-two patients were included. There were 24 men and 8 women with a mean age of 58 years. Lower back pain is the most common symptom. Ultrasound showed a dilatation of the urinary cavities (n=16, 50%) and the CT scan confirmed the bilateral ureteral catheterization (n=11). A fibrous mass that surrounds abdominal aorta, the inferior vena cava and the iliac vessels was noted in CT scan (23 cases), in MRI (9 cases). Intravenous urography (IVU) noted ureteropelvic dilatation in 4 cases. Five patients suffered from RFP associated to pulmonary fibrosis (n=2), sarcoidosis (n=1), immunoglobulin G4-related disease (n=1), Riedel thyroiditis and orbital pseudo tumor (n=1).

Conclusion: The diagnosis of RPF is based on imaging, which seeks an etiology, an associated disease or a complication. CT-scan is the preferred imaging method. MRI may facilitate assessment of disease extent.

Introduction

Retropertineal fibrosis (RPF) is a rare fibroblastic inflammatory process. It results on the progressive transformation of retroperitoneal adipose tissue into a retractile fibrous tissue. It compresses ureters and abdominal vasculature [1]. The incidence is more pronounced in men and the peak of frequency is between 40 and 60 years old [2]. Clinical and biological polymorphism causes diagnostic and therapeutic difficulties. However, imaging had a valuable aid in the diagnosis especially by CT scan, MRI and renal PET-Scann.

The objective of our study was to determine the interest of radiological tools in the management of retroperitoneal fibrosis.

Patients et méthodes

It was a retrospective study which included all patients having RPF. It was conducted in the Internal Medicine Department between 1999 and 2016 in Sousse, Tunisia. Data was collected from medical records. Radiological examinations were interpreted and reviewed by a radiologist at the Sahloul University Hospital.

Data collection

All patients’ demographic criteria were studied including age, gender, comorbidities, presenting symptoms and laboratory findings. Imaging findings were reviewed. Due to the age of some files, modern explorations were practiced only in a limited number. We note the progression of the imaging as they are introduced into the technical platform of our hospital.

For the statistics, we used SPSS version 20 software.

Results

Demographics

Between 1999 and 2016, we enrolled 32 patients. It included 24 males (75%) and eight females (25%) with a mean age 58 years old. Low back pain was the most important symptom (100%).

Several radiological tools were used to establish the diagnosis of RPF in our analysis.

Intravenous Urography (IVU)

UVI was practiced in four patients (12.5%). It had shown hydronephrosis in all cases.

Abdominal ultrasound

Abdominal ultrasound was done in 20 cases. It was normal in four cases (20%) and it had shown ureters sheath in all cases. Unilateral ureteral dilation was noted in 12 cases (37.5%). however, it was bilateral in four cases (12.5%). Ultrasound imaging showed hydronephrosis in four cases (12.5%). Anechoic fibrous lesion was noted in three cases (9.4%). Abdominal aortic aneurysm was detected in two cases (6.2%).

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Abdominal CT-scan

It was practiced in 26 cases (81.2%). It showed retroperitoneal sleeve with homogenous density and regular contours associated with abdominal vascular sheath in 23 cases (71.8%) (Figure 1). Peri ureteral lesions were noted in 11 cases (34.3%). Thrombosis of the renal vein extended to the inferior vena cava was identified in one patient (3.1%). An anerysm of the abdominal aorta was noted in two cases (6.2%).

Renal MRI

Renal MRI was done in eleven cases (34.3%). It showed vascular sheath in nine cases (81.8%). Renal venous thrombosis extended to the inferior vena cava was noted in one case (9%) (Figure 2).

Radiological aspects of pathologies associated with RPF

RPF was isolated without any other pathology in 27 cases (84.4 %). Auto-immune pathology associated with RPF was noted in five cases (15.6 %). It was pulmonary fibrosis in two cases (40%). Sarcoidosis and immunoglobulin G4-related were noted in one case each one (10%). The last patient had Riedel thyroiditis and orbital pseudo-tumor (10%).

Pulmonary fibrosis

Pulmonary and mediastinal fibrosis was noted in two patients who were 48 and 54 years old respectively. Diagnosis was made by the CT-scan aspects.

Sarcoidosis

Observation: It was a male patient who had 55 years old. He was treated for cervical adenopathy in the last two years. He had asthenia, loss of weight and bilateral low back pain. Laboratory investigations showed inflammatory biological syndrom. CT-scan had shown retroperitoneal sleeve with vascular sheath (Figure 3). Histology examination showed multiple epithelioid and giganto-cellular granulomas. We started him with corticoistertoids with favourable outcomes.

Riedel thyroiditis and orbital

Observation: It was a female patient who had 45 years old. She was treated with Tamoxifene. Then, medical examination found goiter associated with diplopia. CT-scan showed thyroid and mediastinal fibrosis. (Figure 4).

Brain CT-scan showed orbital fibrostic sleeve with orbital pseudotumor (Figure 5).

Hyper Ig4 syndrome

Observation: It was a male patient who was 71 years old with a medical history of diabetes, hypetension and anemia. He consulted for fatigue and asthenia. Abdominal CT-Scan showed peri aortic and peri iliac artery fibrosis. Laboratory investigations showed increased IgG4 (= 1,940 g/l). we did not complete investigations because the patient refused any other tests.

Discussion

RPF is a rare inflammatory pathology. It is still mysterious and is requiring further explanation. There are limited available data about RPF in the litterature. Retroperitoneal fibrosis can involve any organ proximate to the retroperitoneum. It is characterized by the fibous involvement of the renal vessels and the ureters causing the urinary excretion impairment [3]. Intestinal or biliary-pancreatic involvement and other pelvic organ and peripheral nerve involvement have all been reported. The clinical presentation includes low back pain, renal failure, anemia and accelerated sedimentation rate. [2]. CT-scan and MRI are the most comonly used for diagnosis and the therapeutic follow up [4-6]. Although intravenous urography was the key radiological exploration in previous bibliographic references, it is currently abandoned in favour of more efficient and less invasive explorations [7].

Diagnosis of RPF

Intravenous Urography

It includes the usual triad of pyelocaliceal and proximal ureter dilation (20% -70%), extrinsic compression and medial deviation of the ureters (Figure 6) [8, 9].
IVU is abandoned in favour of CT-scan and MRI which offer more detailed and specific information about RPF [10].

It was practiced in four cases and it showed indirect signs of obstruction in all cases.

**Abdominal ultrasound imaging**

In some cases, ultrasound reveals a well-defined mass with variable echostructure in close contact with the abdominal vessels [2,11].

It frequently shows a repercussion on the upper urinary tract with hydronephrosis [2,9].

This imaging is no more used in the positive diagnosis of RPF. It makes it possible to suspect the disease without confirming it. Renal doppler ultrasound shows indirect signs of RPF [10,12,13]. In our analysis, ultrasound showed ureter sheath associated with dilation in 16 cases (50%).

The abdomino-pelvic ultrasound was justified in several patients by the initial symptomatology which was nonspecific and dominated by lumbar pain. It will serve as a descrambling examination and will be supplemented by more sensitive examinations for the RPF.

**Abdominal CT-scan**

In our series, the CT performed in 26 patients made the diagnosis in 23 cases in 88% of the cases. Our results are similar to those of the literature, since CT led to the diagnosis of RPF in 72% of cases in the Lugosi study [10]. In the Sérézal study [15], the CT confirmed the diagnosis of RPF in 100% of cases.

Thus, abdominal CT remains the standard imaging method in diagnosis [14,16]. It shows a well-delimited fibrous tissue with homogeneous density which is engaging the large vessels and the ureters and enhancing to the injection of contrast in the early stages (Figure 7) [3,17].

A radiological classification recently proposed by Scheel [18] is available in some published studies [15]. This classification involves distributing lesions of fibrosis as follows:

- **Classe I**: aortic and iliac artery location;
- **Classe II**: inferior vena cava involvement;
- **Classe III**: lateral extension to the ureter;
- **Classe IV**: Extension to the renal hilum with involvement of the renal artery or the renal vein.

**Renal MRI**

MRI offers more advantages than CT-scan. It analyses of the characteristics of the peri-aortic fibrous sleeve and its anatomical ratios (Figure 8).

It also allows the acquisition in many planes. The injection of contrast agent based on gadolinium is significantly less nephrotoxic than the iodine product for used for CT [2]. Finally, MRI helps to differentiate between malignant RPF and non-tumoral forms [16]. This technique allows the diagnosis, the assessment of obstructive complications and the follow-up of RPF [10].

In our study, it showed RPF in 82% of cases.

**PET- scann (Figure 9) [19]**

This imaging seems to be useful in the initial review of the RPF [10].
Although fixation does not systematically show morphological data visualized by CT scan and MRI, PET scan assesses the extension of RPF, the vascular and the perivascular lesions [20,21].

It could also help in the monitoring of the RPF [21]. In the study of Serezal [15], this technique was performed in 11 patients, at different stages of the evolution of the RPF. Retroperitoneal lesions were then fixed in 8 cases out of 11 (72.7%).

Moreover, in the Desbois study [22], nine DFG-TEP-CT were performed, showing discrete hypermetabolic foci in six cases (66.6%). The PET scanner can be used for the detection of multifocal fibrosis lesions when an inflammatory disease is associated with RPF [23]. It also has an interest in the follow up of the disease during corticosteroid therapy [24].

Renal scintigraphy with 2,3-dimercaptosuccinic acid (DMSA)

It can be proposed to evaluate the function of each [24]. Renal scintigraphy provides a complementary functional element to the etiological analysis of renal insufficiency during the RPF [25]. No renal scintigraphy was requested in our study (Table 1).

### Diagnosis of illness associated with RPF

FRP may be isolated or associated with systemic diseases (Table 2) [2,26,27-29]. Many publications report associations with multiple diseases whose substratum can be immunological or inflammatory.

In the Lugosi study [10], RPF was associated with systemic vasculitis in three cases: granulomatosis with polyangiitis (n = 1) and

### Table 1. Medical Interest of Imaging [24]

| Imaging          | Signs                              |
|------------------|------------------------------------|
| Ultrasound Imaging | Caliceal dilation                   |
| CT-scan          | Obstruction, vascular and peri vascular lesions |
| IVU              | Obstruction                         |
| MRI              | Vascular and peri vascular lesions  |
| Renal scintigraphy | Functional study of each kidney    |
| TEP-Scan         | Multifocal fibrosis                 |

Figure 7. CT-scan showing fibrous lesion with vascular involvement

Figure 8. Abdominal MRI perivascular and retroperitoneal lesion in hyposignal T2(A) ureteral involvement (B).
Takayasu’s disease (n = 2). Riedel’s thyroiditis was associated with RPF in one case. In the study of Ouertani [13], FRP was associated with systemic vasculitis in 2 cases: granulomatosis with polyangiitis and Takayasu’s disease in one case each. In our study, FRP was associated with pulmonary fibrosis (2 cases), sarcoidosis (1 case), Riedel’s thyroiditis and an orbital pseudotumor (1 case) and hyper-IgG4 syndrome in the latter case.

**Conclusion**

RPFremains a rare inflammatory disease. Its pathogenesis is still poorly understood. Clinical signs and symptoms are not very specific. The biological assessment is essential but often not very useful for the diagnosis. The radiological assessment helps to demonstrate the fibrous lesion, to evaluate its extension and its renal repercussion and finally to orient the biopsies for the positive diagnosis.

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