ABSTRACT

Primary adrenal lymphoma is a very rare condition; it represents less than 0.5% of all adrenal tumors. Adrenal lymphoma is often bilateral and is mostly a B-cell type. T-cell lymphoma is exceptional. Diagnosis and treatment are challenging and prognosis is pejorative. We are reporting the case reports of two patients with primary and bilateral adrenal lymphoma. Both patients were operated without histological evidence, for the first, adrenalectomy with right nephrectomy and biopsy of the left mass were performed, histological result show the non-Hodgkin’s lymphoma cell-T, the evolution were marked with the death of patient by pulmonary embolism after the first cycles of chemotherapy, for the second patient left adrenalectomy and biopsy of the right mas were made, the histological result show the non-Hodgkin’s lymphoma cell-B, the deceased patient in the outcomes. A review of literature displays the pejorative prognosis of these tumors. The objective is to have a less aggressive etiological approach without exploratory surgery.

Keywords: Adrenal; Bilateral; Lymphoma; Primary.

INTRODUCTION

Primary adrenal lymphoma (PAL) is very rare, accounting for approximately 0.5% of all adrenal malignancies. About 200 cases have been reported to date. The bilateral presentation is frequent and large B-cell type is prominent in histology. Diagnosis and management are real issues and prognosis remains always pejorative. We report 2 cases of primary and bilateral non-Hodgkin’s lymphoma of adrenal.

CASE REPORTS

Case I:

A 43-year-old patient with no medical history, presented brutally an acute abdominal pain in the right flank with sweat, headache and fever. Clinical examination found sensitivity in the right flank. Abdominal CT scan revealed a rounded mass of the right adrenal gland pushing the right kidney and liver with contact with the inferior vena cava associated with another mass of the left adrenal gland, there was without hepatic metastases (Figure 1). Cortisolemia and methoxylated derivatives were normal. A bone marrow biopsy specimen was performed to rule out secondary lymphoma; the result was negative. Median laparotomy revealed a huge right adrenal mass with invasion of the right kidney and all the retroperitoneal space. One-piece right adrenalectomy with right nephrectomy were performed. Biopsy was taken from left mass biopsy (Figure 2). The histological study with immunohistochemistry confirmed the diagnosis of primary non-Hodgkin’s malignant lymphoma with
large T-cells phenotype (Figure 3). Chemotherapy was indicated. Patient died after pulmonary emboli following first chemotherapy.

**Figure 1:** Abdominal CT showing the two right adrenal masses pushing back the liver and right kidney and in contact with the inferior vena cava and left adrenal mass.

**Figure 2:** Adrenal and right kidney tumorectomy

**Figure 3:** Histological aspect of non-Hodgkin malignant large T-cell lymphoma (G x 20)

**Case II:** A 54 year-old patient with no medical history, was complaining from localized abdominal pain in both flanks during the last 04 months associated with vomiting, fever and important weight loss. Clinical examination found an altered patient with tachycardia at 110 and localized tenderness to both sides without lumbar contact and palpable mass on palpation. The abdominal ultrasound and CT scan showed two adrenal mass weakly enhanced after contrast injection (Figure 4): the right mass pushing back the IVC with invasion of the upper pole of the right kidney. There was no liver metastases or deep lymphadenopathy. Biologically the patient had hypokalemia with minimal elevation of normetanephrin. Median laparotomy showed the presence of two large adrenal masses. A left adrenalectomy with a biopsy of the right adrenal mass were performed. The histological study with immunohistochemistry revealed non-Hodgkin’s malignant lymphoma large phenotype B-cells (Figure 5). The patient died in the post-operating suites.

**Figure 4:** Abdominal CT showing the 2 hypodense adrenal lesions weakly enhanced after the PC measuring right 11x6cm and left 10x5cm. The right mass pushes the inferior vena cava and invades the upper pole of the kidney.

**Figure 5:** Histological aspect showing non-Hodgkin’s large cell malignant lymphoma of B-cell phenotype

**DISCUSSION**

Anusual locations and special presentations of lymphoma has been reported in literature (biliary tract, conjonctiva, and others...) and interest in this entity is particular (1-3). Isolated Primary lymphomas of adrenal gland are exceptional, representing less than 0.5% of all adrenal neoplasm. Secondary forms due to lymphomatous infiltration...
are mostly reported (4, 5). Isolated Primary lymphomas of adrenal gland are prominent in men (sex ratio= 3/1) and mean of age is 39-89 year-old (7, 8). Bilateral involvement is most common and adrenal function often remains normal, the diagnosis is based on histology and immunohistochemistry findings (8, 8). Etiopathogeny is still unexplained but the role of Epstein Barr Virus has been suggested (9). The most common mode of presentation is either related to lymphoma itself with fever, altered general condition, abdominal pain and weight loss. Symptoms could be related to adrenal insufficiency (4, 10, 11), but usually the diagnosis is late and accidental (12); Maugendre reported 14 cases (13). Asymptomatic cases with post-operative diagnosis are exceptional (14, 15). Biology is not specific, it may show inflammatory biological syndrome, elevated LDH rates or hypercalcemia (16, 17). Diagnosis can be suspected on imaging findings. Abdominal ultrasound of adrenal masses can be echoic with liquid zones corresponding to hemorrhagic or necrotic areas (4,18-20). The adrenal CT scan usually shows a well-limited oval homogeneous mass that is slightly enhanced by the contrast agent (21). Arguments supporting the diagnosis of lymphoma are: the large size >3cm and the absence of fatty area or calcification or hypervascular character (22) but those are not specific criteria. The MRI allows the visualization of adrenal masses but does not make the difference between lymphoma and metastasis (17, 22). The definitive diagnosis is histological specimen (18-20); Biopsy specimen could be performed by ultrasound or CT scan guidance. Laparotomy must be suggested as a last option and after eliminating a pheochromocytoma. However, Luton proposes systematic surgery (23). In our cases, laparotomy was performed first with the intention of an adrenalectomy but due to the local invasion, only a simple biopsy was done. Primary adrenal lymphoma may be confused with undifferentiated carcinoma or metastatic lesions of the adrenal gland. Immunohistochemical study should be performed with lymphoid markers before definitive diagnosis is made (24). Large cell lymphoma (high grade according to Kiel) is clearly prominent (71% vs 11% small cell lymphoma). Phenotype B is the predominant according to the review of Lecaltier et al (18, 19). For our patients, T-cell and B-cell phenotypes were found. To retain the primitive character of lymphoma, it is necessary to confirm the absence of other localization. Clinical examination is important, checking the absence of nodes, hepatomegaly and splenomegaly. Bone marrow biopsy is indicated. Thoraco-abdominal CT-scan, upper gastrointestinal endoscopy, colonoscopy and bronchoscopy should be all performed with multiple biopsies and a (4). In our patients, all investigation showed negative results which confirmed the primary adrenal location of non-Hodgkin’s lymphoma. Regarding the severity of the disease, poly-chemotherapy based on CHOP or MACOP-B seems to be the treatment of choice (4, 18, 19). Surgery alone or in combination with chemotherapy or adjuvant radiotherapy presents limited benefit (4). The prognosis of primary adrenal lymphoma is still considered fatal (4). Al-Fiad et al reported survival rate of 10 days to 11 months after diagnosis (25) which is significantly shorter than thyroid lymphomas (10). The literature reported rare cases with excellent response to treatment with prolonged total or partial remission (18). Prognosis factors are: age over 60 years old, tumor size greater than 10 cm, histological aggression and phenotype, existence of more than one extra-lymph node localization, high levels of LDH and ß-2 macroglobulin up to 3 mg/l (18).

CONCLUSION

Bilateral primary adrenal malignant lymphoma is a rare entity with bad prognosis. It must be evocated before any bilateral mass of the 2 adrenals, the diagnosis is histological. Mini-invasive approach is recommended for biopsy.

CONFLICT OF INTEREST

There is no Conflict of Interest to declare.

REFERENCES