High grade primary adrenal intravascular large B-cell lymphoma manifesting as Addison disease

J. Venizelos, D. Tamiolakis¹, M. Lambropoulou¹, G. Alexiadis², G. Petrakis¹ and N. Papadopoulos¹

Department of Pathology, Ippocration General Hospital of Salonica. ¹Department of Histology-Embryology, Democritus University of Thrace. ²Diagnostic Radiology Center of Alexandroupolis, Thrace, Greece

INTRODUCTION

It is known that, development of adrenal failure requires destruction of more than 90% of the adrenal gland tissue (1). Most cases of adrenal insufficiency caused by malignant neoplasms are due to metastases from a lung, breast, or stomach primary, and malignant melanoma (2). Malignant lymphoma could occur in any body organ. However, non-Hodgkin lymphoma (NHL) arising in the endocrine system represents only 3% of extranodal malignant lymphomas, and is usually confined to the thyroid gland (3). Adrenal infiltration at the time of presentation is rare and difficult to quantify. Surprisingly secondary involvement by lymphoma of the adrenal glands in autopsy findings rises 35% (4). Primary adrenal lymphoma tends to be predominant in males compared to females (2.5-7 times), occurs with an average age of 61 years (38-81), morphology of large cell lymphoma and a B cell phenotype with a small number of T cell phenotype (5). Of the 70 cases reported so far, bilateral cases constitute 70% of the total adrenal lymphoma (5). Symptoms may occur due to the tumor itself or adrenal failure.

CASE STUDY

A 68-year-old male was diagnosed 7 months ago with anaemia due to duodenal ulcer and subsequently treated with anti-ulcer drugs and ferrum, but he failed to respond. Four months prior to admission, he complained of nausea and developed fever and diarrhoea. He presented with generalized pigmentation, and organomegaly without peripheral lymphadenopathy. Because of anaemia, lymphocytosis, and hepatosplenomegaly, the patient underwent an abdominal CT investigation that revealed bilateral adrenal masses, hepatic and splenic enlargement, but no other pathology. Laboratory tests showed normal complete blood count and no abnormal results on urinalysis. Biochemical tests demonstrated a
high LDH level (427IU/l). Serum electrolyte analysis revealed no hyponatremia, but elevated values of serum potassium and serum CRP. Endocrinological data showed plasma ACTH 820 pg/ml, plasma cortisol 4.4 μg/day, and urinary-cortisol 2.3 μg/day. Furthermore, urinary 17 KS, urinary 17-OHCS, serum DHEA-S, serum estradiol, and plasma aldosterone were low, confirming adrenal failure.

Abdominal CT detected two masses measuring 5 cm and 2 cm in the right and left adrenal gland respectively, with comparable and inhomogeneous density after contrast administration. The right adrenal revealed hypodense center due to necrosis (Fig. 1). No celiac lymph node enlargement was shown. Magnetic resonance imaging revealed low signal intensity on T1 weighted images and high signal intensity with heterogeneous appearance in both of the adrenal glands on T2-weighted images. We suspected metastatic tumors and searched the entire body for the primary lesion with negative results. 67Ga scintigraphy did not reveal any abnormal uptake in the adrenals or other organs.

The patient was administrated with hydrocortisone replacement therapy resulting in immediate improvement of symptoms. A surgical excision of the left adrenal gland followed.

**PATHOLOGICAL FINDINGS**

Gross examination showed a 9.5 x 7 x 1.5 oval, encapsulated, rubbery, brown-white coloured tumor. No adrenal tissue was recognised. Histopathologically (Fig. 2), there was complete and diffuse infiltration by large pleomorphic cells with round hypochromatic nuclei, 2 to 3 prominent nucleoli at the periphery of the nuclear envelope and a slightly eosinophilic cytoplasm. There were also medium sized cells some of them with nuclear grooves, a few immunoblasts, and plasma-like cells. Mitotic figures were frequent.

**IMMUNOHISTOPATHOLOGY**

Immunohistochemical staining demonstrated that most tumor cells were positive for LCA (leukocyte common antigen) (Fig. 3), B-cell markers (CD20, CD79a), but although stained for bcl-2, they did not stain for CD5. Malignant cells were surrounded by endothelial (CD34 positive) cells (Fig. 4).

Based on pathological assessment, the final diagnosis was intravascular large B-cell lymphoma, a subtype of extranodal diffuse large B-cell lymphoma characterized
by the presence of lymphoma cells in the lumina of small vessels.

Following the resection of the left adrenal gland, serum electrolytes improved under the replacement therapy with hydrocortisone (30 mg/day). The pigmentation gradually disappeared, and plasma ACTH decreased to 18.7 pg/ml. However, after the operation, persistence of symptoms of general fatigue, appetite loss, low-grade fever, steal sweat and elevation of CPR and LDH was observed. The activity of NHL remained high. Subsequent abdominal enhanced CT revealed multiple low intensity areas in the liver and celiac lymph node enlargement, which had not been detected by then. The disease was staged as IVB according to the Ann Arbor system and the patient underwent chemotherapy with CHOP (Cy clophosphamide, doxorubicin, vincristine, and prednisolone), two cycles twice a week, resulting in remission. Seven months later the patient passed away due to infection in the lungs.

**DISCUSSION**

We treated a 68 year old man who was diagnosed with bilateral primary adrenal malignant lymphoma. The most common manifestations of malignant lymphoma are the involvement of the amygdale, stomach and pharynx, in addition to the lymph nodes, although the involvement of various other organs has been described. Involvement of the adrenal gland in malignant lymphoma cases is reported to be 25% at autopsy (6), but malignant lymphoma arising in the endocrine glands represents only 3% of extranodal malignant lymphomas, and is usually confined to the thyroid gland. Therefore, primary adrenal malignant lymphoma is rare.

The present patient showed typical symptoms and endocrinological findings of adrenal failure. Primary adrenal malignant lymphoma was described in 24 patients of 67 reported cases complicated with adrenal failure including subclinical or partial adrenal failure (7,8). These reports showed certain characteristic features for primary adrenal malignant lymphoma, such as male predilection, involvement of both adrenal glands and being of the diffuse large B cell type. CT and MRI revealed heterogeneous tumors in our case. The heterogeneous tumors on imaging studies are usually tumors with bleeding and necrosis, making it very difficult to differentiate malignant lymphoma from adrenal metastatic tumors or adrenal cortical carcinomas (9,10). Necrosis is associated with rapid tumor development. Surgery was performed to allow rapid histopathological examination under operation for the final diagnosis. The final diagnosis was intravascular lymphoma (IVL) based on histopathological examination. IVL was classified as diffuse large B-cell lymphoma subtype, according to the new WHO classification (11). Several cases of IVL have been reported since the first patient was reported by Pfleger and Tappeiner (12). Generally, antemortem diagnosis of IVL is very difficult because of the exceedingly rapid clinical course and protean presentation of the disease. Additionally, the poor prognosis of IVL results partially from delays in diagnosis. IVL is characterized by massive proliferation of atypical mononuclear cells within arterioles, capillaries, and venules. This disease is therefore rarely diagnosed clinically but could present with various clinical features, such as fever, skin eruptions, neurological symptoms, and weight loss (13,14). Moreover, the most common manifestations of IVL are infiltration of the skin and central nervous system, although infiltration of atypical cells has been described in various organs (15). Numao et al. (16) indicated that primary and secondary adrenal gland infiltration of IVL was detected in 67% of patients diagnosed with IVL.

The sensitivity of IVL to systemic chemotherapy is very high and particularly when early diagnosis is made followed by treatment at an early stage. Therefore, early diagnosis and treatment of IVL is crucial for prognosis. Recent studies showed that de novo CD5+ diffuse large B-cell lymphoma (CD5+ DLBCL) is phenotypically and genotypically different from CD5− DLBCL (17). The survival of patients with CD5+ DLBCL is significantly worse than that of patients with CD5− DLBCL. Moreover, several patients with
CD5+ IVL have been described (18,19). However, in the present case the tumor cells did not stain for CD5.

IVL should be suspected in patients with bilateral adrenal tumors who present with rapidly progressive adrenal failure.

REFERENCES