Bilateral primary non-Hodgkin's lymphoma of the adrenal glands with adrenal insufficiency: a case report

Authors: Utsunomiya, Masato; Takatera, Hiroshi; Itoh, Hiroshi; Itatani, Hiroaki; Tsujimura, Takahiro

Citation: 泌尿器科紀要 (1992), 38(3): 311-314

Issue Date: 1992-03

URL: http://hdl.handle.net/2433/117503

Type: Departmental Bulletin Paper

Publisher: Kyoto University
BILATERAL PRIMARY NON-HODGKIN'S LYMPHOMA OF THE ADRENAL GLANDS WITH ADRENAL INSUFFICIENCY: A CASE REPORT

Masato Utsunomiya, Hiroshi Takatera, Hiroshi Itoh, Takahiro Tsujimura* and Hiroaki Itatani

From the Departments of Urology and Pathology*, Sumitomo Hospital

A 72-year-old woman with bilateral non-Hodgkin's lymphoma of the adrenal glands causing adrenal insufficiency is reported. The left-side lymphoma (diffuse large B-cell lymphoma) was removed surgically but the right-side lymphoma could not be removed. Complete response was obtained with subsequent combined chemotherapy and was maintained for 6 months with repeated chemotherapies. However, the patient died of liver recurrence 1 year after admission. In cases of malignant lymphoma forming a bulky mass, surgery with subsequent chemotherapy appears to reduce the cancer volume and improve the therapeutic outcome.

Key words: Primary non-Hodgkin's lymphoma, Bilateral adrenal glands, Chemotherapy, Operation

INTRODUCTION

The primary lesions of malignant lymphoma in the urologic field are found in the kidneys, testes, bladder and related areas. Although lymphoma involving the adrenal glands is sometimes encountered in cases of widespread lymphoma, malignant lymphoma arising from the adrenal gland is extremely rare. We describe such a rare case of bilateral malignant lymphoma (diffuse large B-cell lymphoma) of the adrenal glands causing adrenal insufficiency and discuss the role of surgical procedure for this order.

CASE REPORT

A 72-year-old female was admitted to our hospital for diminished consciousness and for general fatigue on May 26, 1987. She had been febrile and confused during the week before admission. The serum sodium level was 129.0 mEq/l and the serum LDH elevated to 1,783 U. CT-scan showed bilateral, huge, homogeneous adrenal tumor but no other retroperitoneal adenopathies (Fig. 1). Although the serum cortisol level was at the lower limit of the normal range (11.7 mg/dl), the ACTH was elevated (170 pg/ml). We suspected that this condition was caused by adrenal insufficiency secondary to large adrenal masses. Therefore an adrenal steroid hormone with antibiotics was administered immediately. After two days of treatment, the patient had fully regained consciousness while her fever had subsided. Angiographic findings revealed bilateral tumors originating from the adrenal glands. Additional radiographic studies, including gastrointestinal series and chest film, revealed no other abnormal findings. Results of other hormonal studies of the serum and the urine were also normal. On June 2, 1987, we performed adrenalectomy based on the diagnosis of bilateral nonfunctioning adrenal tumors. The tumor on the left side could be resected, but that on the right side could not because of direct invasion to the liver, kidney and inferior vena cava. Histological examination showed the tumor to be a non-Hodgkin's lymphoma, diffuse large cell type (by Working Formulation) (Fig. 2), while immunohisto-
Fig. 1. CT-scan shows symmetrical enlarged adrenal masses. The right mass measures 9.3 x 5.7 cm and the left mass 8.9 x 6.0 cm. Additional caudal scans showed no retroperitoneal adenopathies.

Fig. 2. Histologic feature of the tumor shows non-Hodgkin’s lymphoma, diffuse large cell type. (H&E, x200)

Fig. 3. Immunostaining of the tumor cells using the ABC method with methyl green nuclear counterstaining. Many lymphoma cells are positive for L-26, B cell marker (DAKO, Ccurate Chemical, Westbury, NY). (x200)

Fig. 4. The unresectable right adrenal tumor had been completely eliminated after the second chemotherapy.

Fig. 5. $^{67}$Ga scintigram also shows complete disappearance of the right adrenal tumor. (A: Before B: After the chemotherapy)

chemical staining with the avidin-biotin-peroxidase complex (ABC) method of Hsu et al. proved that the tumor cells were of B cell origin (Fig. 3). After surgery, the patient was given 1 mg of vincristine, 750 mg of cyclophosphamide, 20 mg of doxorubicin and 50 mg of prednisone daily for 5 days. This regimen was repeated 3 times in 2 months. Serum LDH decreased rapidly to normal and the right adrenal tumor had completely vanished from the CT-scan and $^{67}$Ga scintigram after the second treatment (Fig. 4, 5). Complete response
was maintained for 6 months, but adrenal hormone replacement (hydrocortisone 20 mg/day) could not be ceased. Six additional chemotherapy courses were given (Fig. 6), but multiple recurrence in the liver as well as rapid elevation of serum LDH was noted on the 67Ga scintigram taken in January, 1988. Further chemotherapy modified with VP-16 and others was ineffective, and the patient died on May 8, 1988. Postmortem examination was not done.

DISCUSSION

Malignant lymphoma originating from the adrenal gland is rare, and only 4 cases of bilateral lymphoma originating in the adrenal glands have been reported in the literature6-8). Cases of bilateral malignancies in the adrenal gland are rarely accompanied by adrenal insufficiency9). However, 4 of the 5 cases including our case (80%) presented with adrenal insufficiency. Therefore, in patients with bilateral adrenal masses with adrenal insufficiency, the differential diagnosis should include malignant lymphoma. It is of interest that a case reported by Carey et al.8) was represented as an instance of reversal of Addison’s disease after antineoplastic therapy. Schnitzer et al.7) reported the first case of T-cell lymphoma of the adrenal glands based on careful immunochemical examination. Our case is the first case to show immunohistological evidence of B-cell origin lymphoma.

Considering the high mortality rate of non-complete responders with malignant lymphoma, it is undoubtedly important to obtain a complete response in the initial stage. In spite of better CR rates obtained with current chemotherapy, full dose chemotherapy is not often completed because of the patients' high age and poor condition. Also this fact means limiting CR rate of this order. Therefore, before the intensive chemotherapy, it is of value to reduce the cancer volume when it presents with a large mass. Although surgery is generally not considered curative for malignant lymphoma because of its systemic nature, surgical procedure is effective for reducing cancer volume as well as establishing the correct diagnosis. In our case, we could reduce the tumor volume by half and could gain complete response for 6 months with subsequent combined chemotherapy, offering palliation with minimum morbidity. In conclusion, we recommended surgical procedure with subsequent chemotherapy as an effective method for selected cases of malignant lymphoma, especially those with large masses.
REFERENCES


(Received on April 15, 1991)
(accepted on June 19, 1991)