A Case Report of Large Renal Cell Carcinoma

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Abstract

We report a case of 70 yr old woman with history of mass per abdomen since 5 months, which was insidious in onset and gradually progressed to present size. It was not associated with fever, haematuria, but gives history of loss of weight and loss of appetite. She was emaciated, poorly nourished with no significant past history. On examination she had mass per abdomen occupying Right lumbar, hypochondrium, umbilical, Right iliac fossa and hypogastrum abutting the anterior abdominal wall. No free fluid, no organomegaly. No evidence of swellings in other part of body, and no supraclavicular lymphadenopathy.

Keywords: Mass abdomen; Renal cell carcinoma; CT abdomen

Introduction

Renal cell carcinoma (RCC) was first reported by Paul Grawitz in 1883. It was named after him as Grawitz tumor, or hypernephroma, according to his belief that the tumor originated in adrenal rests at the upper pole of the kidney. Later, the origin of this tumor in renal tubular cells was documented [1]. Accounting for 2% of all adult malignancies [2], RCC has tendency to spread into the renal vein and the IVC (inferior vena cava). Late presentation is the common initial diagnosis in our part of the world [3], making operative treatment more challenging. Kidney cancer is among the 10 most common cancers in both men and women.

Case Report

We report a case of 70 yr old woman with history of mass per abdomen since 5 month’s, which was insidious in onset and gradually progressed to present size. It was not associated with fever, haematuria, but gives history of loss of weight and loss of appetite. She was emaciated, poorly nourished with no significant past history. On examination she had mass per abdomen occupying Right lumbar, hypochondrium, umbilical, Right iliac fossa and hypogastrum abutting the anterior abdominal wall. No free fluid, no organomegaly. No evidence of swellings in other part of body, and no supraclavicular lymphadenopathy.

Blood investigations: Haemoglobin- 9.9 gms%; Hematocrit-28.2%; Total Count- 7700 cells/mm3, Neutrophils- 85, Lymphocytes-10, Eosinophils-02, Monocytes-03. Bleeding Time- 2’ 15”, Clotting Time – 4’ 30”. Peripheral smear – normocytic normochromic anemia. Blood urea-21 mg%, serum creatinine - 0.85 mg%, Random blood sugar (RBS) - 84 mg%; Liver function tests (LFT), Alkaline phosphatase (ALP)- 471 IU/L, Gamma Glutamyl transaminase (GGT)- 232 IU/L. X Ray chest – no secondaries, no effusion. USG (Ultrasonography) abdomen revealed a large heterogeneous mass occupying Right lumbar, Right hypochondrium, Umbilical and crossing the midline measuring 15 × 12 × 10.5 cm. Right kidney not visualized, Left kidney normal. Liver shows multiple hyperechoiec lesions in both lobes largest measuring 3.2 × 2.6 × 3.5 cm. Bilateral pleural effusion seen (Figures 3,4 and 5). FNAC (Fine needle aspiration cytology) shows round to polygonal cells having coarse chromatin. 2-4 nucleoli arranged in discreates. The nuclear (inferior vena cava) compression by the mass. No thrombosis of the IVC (inferior vena cava) and renal vein. After 24 hrs scan small amount of contrast excreted at periphery. No free fluid. Other solid organs were normal.

Liver multiple hypodense areas were seen, largest measuring 3.2 × 2.6 × 3.5 cm. Bilateral pleural effusion seen (Figures 3,4 and 5). FNAC (Fine needle aspiration cytology) shows round to polygonal cells having coarse chromatin. 2-4 nucleoli arranged in discretes. The nuclear

Computerized Axial Tomography

Abdomen (Plain and Contrast) a well defined large lobulated mass seen in Right renal region extending from sub hepatic region to pelvis, medially crossing midline extending to the medial surface of Left kidney. Encasing retroperitoneal great vessels and renal vessels, measuring 17 × 14 × 13.5 cm. multiple hypodense areas and calcified areas noted. On contrast mass shows heterogenous enhancement, IVC

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margin prominent, the cytoplasm is scant these cells are arranged in solid cells clumps with a flowery arrangement featured suggestive of adeno carcinoma / Renal cell carcinoma [4].

Discussion

Malignant neoplasms involving the kidney may be primary or secondary tumors. Although metastatic lesions out number primary tumors, secondary renal neoplasms are usually clinically insignificant and are principally discovered at postmortem examination.

Patients with Renal cell carcinoma (RCC) present with a range of symptoms, but many are asymptomatic until the disease is advanced [5]. At presentation, approximately 25 percent of individuals either have distant metastases or significant local-regional disease. Other patients, even some with only localized disease, present with a wide array of symptoms and/or laboratory abnormalities. Because of this unusual characteristic, RCC has been labeled the “internist’s tumor” [1]. Today, most tumors are diagnosed incidentally [6].

The classic triad of flank pain, hematuria and flank mass is uncommon (10% cases) and is indicative of advanced disease. Renal cell carcinoma represents a heterogenous group of tumors, the most common of which is clear cell adenocarcinoma [7]. RCC accounts for 3% of adult tumors. The incidence has increased more than 30% over the past two decades. It is generally accepted that the increased incidence rates reflect earlier diagnosis at an earlier stage, largely due to more liberal use of radiological imaging techniques. However advanced disease has also been diagnosed more frequently and mortality has increased as well [6].

Symptomatic presentation correlates with aggressive histology and advanced disease. Incidental tumors may be frequently detected in female and elderly patients, as these groups traditionally seek general medical care more regularly. The mode of presentation can independently predict an adverse patient outcome. Indicators of symptomatic presentations include flank pain, flank mass, varicocele, constitutional symptoms, paraneoplastic syndromes and bone pain related to metastatic disease [8].

Ultrasound scan was found to be useful screening test, but CT (Computerized tomography) is the imaging study of choice to identify malignant features. MRI (Magnetic resonance imaging) can be used in equivocal cases [6].

Pre-operative clinical variables may be used instead of the pathologic stage to determine the risk of recurrence [3].

References