Dual Pathology- Clear Cell Variant of Renal Cell Carcinoma and High Grade Papillary Urothelial Carcinoma of Urinary Bladder a Rare Entity: Case Report

Mukherjee Rina¹, Ganguly Tanmoy¹, Bandopadhyay Jayeeta², Das Dipkana³, Singh Neha⁴
¹Senior Consultant & HOD, ²,³ Senior Resident, ⁴DNB Post Graduate Trainee
Department Of Pathology, B. R. Singh Hospital AndCentre For Medical Education &Research, Eastern Railway, Sealdah, Kolkata-700014, India

Abstract: Simultaneous occurrence of clear cell variant of renal cell carcinoma and high grade papillary urothelial carcinoma of urinary bladder is a rare entity. The review of different literatures revealed incidence of carcinoma of urinary bladder and the prostate in patients with renal cell carcinoma, especially the papillary subtypes. Here we report a case of 61 years male patient presented with hematuria of one month duration. Further investigations revealed SOL in upper pole of left kidney measuring 4x4x3.5 cm and a separate friable growth in the urinary bladder. Urinary Bladder tumor specimen was diagnosed as high grade papillary urothelial carcinoma of urinary bladder and left radical nephrectomy specimen was diagnosed as Clear cell variant of renal cell carcinoma. The etiopathogenesis of this association remains debated. It seems to be based on genetic alterations that have not been formerly proved.

Keywords: Clear cell variant of renal cell carcinoma, high grade papillary urothelial carcinoma of urinary bladder, SOL in upper pole of left kidney, Radical nephrectomy.

I. Introduction

Several studies in the literature dealing with the occurrence of multiple primary malignant neoplasms report incidence rates ranging from 2.6 to 3.9% among cancer patients¹ ² and those studies that include tumors found at autopsy report even higher rates³. The most common sites for multiple tumors vary among the reports but as the occurrence of primary tumors in the genitourinary (GU) tract is higher compared to other organ systems, multiple tumors have a predilection to involve a GU tract organ. Multiple primary malignant tumors involving only the GU tract are rare and present either as multicentric malignant tumors of the urinary epithelium, tumors occurring in bilateral organs or as multiple tumors with dissimilar histology. We report a 61 years old male patient who first underwent transurethral resection of bladder tumor (TURBT) and reported as high grade papillary urothelial carcinoma of urinary bladder with squamoid differentiation. Later he underwent left radical nephrectomy and found to have clear cell variant of renal cell carcinoma.

II. Case Report

A 61 years old gentleman, known smoker, presented with complaints of burning sensation during micturition, abdominal pain and heaviness along with few episodes of hematuria and weakness of one month duration. On clinical examination, mass was felt on the left flank. Investigation includes radio-imaging studies like USG KUB, MDCT Urography and PET CT, showed a large solid-cystic lesion in upper pole of left kidney measuring 5.5x5.1x6.3cm and a separate neoplastic lesion in urinary bladder involving left lateral & posterior wall extending to the base measuring 7.1x5.1x5.5cm. Cystoscopy showed a vascular and friable tumor involving base, left lateral wall and anterior wall around bladder neck. Transurethral resection of bladder tumor was done and sent for histopathological examination. It was reported as high grade papillary urothelial carcinoma of urinary bladder with squamoid differentiation. Later he underwent left radical nephrectomy and found to have clear cell variant of renal cell carcinoma.

For kidney tumor, left radical nephrectomy was done and sent for histopathological examination. On cut section, a solid, well circumscribed variegated yellowish tumor in upper pole of the kidney measuring 4x4x3.5 cm, sharply demarcated from the surrounding tissues with foci of haemorrhage, necrosis and cystic changes was noted. Renal pelvis, ureter, renal vessels, and pericapsular area were grossly unremarkable. Histopathological examination revealed a pT1aNxMx, clear cell variant of renal cell carcinoma, Fuhrman grade II. No pericapsular extension was identified. Ureter and renal vessels were free of tumor invasion. No lymph node was found. Neo-adjuvant chemotherapy followed by radical cystectomy is awaited.
Images:

**Fig 1&2:** MDCT Urography and PET CT image showing left kidney and urinary bladder tumor.

**Fig 3 & 4:** Photomicrographs of TURBT specimen showing high grade papillary urothelial carcinoma of urinary bladder with squamoid differentiation.
III. Discussion

Clear cell variant is the most common subtype of renal cell carcinoma and papillaryurothelial carcinoma isthe most common urinary bladder carcinoma. Multiple primary malignant tumors can occur in all decades, but they are most common in the geriatric population. It is a well known fact that urinary tract is one of the most common sites for multiple primary tumors. Extensive search revealed that multiple primary tumors occur in 3.1% of total cancer patients and 13.9% of these have genitourinary tumors. Multiple malignant tumors involving more than one genitourinary site are rare. In one study, 5.4% of the patients with urinary bladder neoplasms had malignant tumors outside the urothelium and of these 4.3% were renal cell carcinomas. Several studies done by Begget al, Neuzillet et al, Jensen et al and others concluded that urinary bladder cancer patients have higher risk of development of renal cell carcinoma. Neuzillet et al and Rabbaniet al concluded that this dual association is usually observed in papillary subtypes of renal cell carcinoma. But in our study, this dual association was observed between the clear cell variant of renal cell carcinoma and high grade papillary urothelial carcinoma of urinary bladder. Extensive search does not reveal such coexistence in published literature. Smoking, drinking, alcohol oncogenic substances (like - radiation) and genetic mutations are the common risk factors for the development of primary cancers like renal and urothelial carcinomas.

Some studies reported that the chemo and/or radio therapy for one cancer may induce the secondary carcinoma. But in our study no such intervention was done. The carcinogenesis is an anomalous phenomenon determined by a clonal selection. Association of dual primary cancers affecting different organs necessitates the coexistence of genetic mutation or a multifactorial etiology.
IV. Conclusion

This case reported as a coexistence of clear cell variant of renal cell carcinoma with high grade papillary urothelial carcinoma of urinary bladder in an adult smoker. This dual association is least reported in different literature. Due to our limited resources we could not establish whether this association is due to some genetic alterations or it is just an incidental finding. But further researches are necessary to establish the actual etiopathogenesis of such togetherness.

References