MiT Family Translocation Renal Cell Carcinomas

MiT Ailesi Translokasyon Renal Hücreli Karsinomlar

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Introduction

MiT family translocation renal cell carcinomas (RCCs) are particular neoplasms with their clinically aggressive behavior and histopathologically distinctive appearance (1). These tumors tend to occur in young age group and consist of nearly 40% of pediatric and 1.6-4% of adult RCCs (1,2).

These tumors are caused by two types of translocations involving TFE3 (transcription factor E3) and TFEB genes which are belong to the Microphthalmia-associated transcriptional factor family (MiT family) (2,3). TFE3 is located on Xp11.2, while TFEB is on chromosome 6. Tumors showing Xp11 translocation are much more common than those involving TFEB translocations. The most common subtypes of gene fusions are ASPSCR1-TFE3 and PRCC-TFE3 (1,3). t(6;11) RCC is the rare form of translocation RCCs which harbors gene fusion among TFEB and MALAT-1 (2). These gene rearrangements result in overexpression of several fusion proteins including TFE3 and TFEB which can be demonstrated with a nuclear labeling by immunohistochemistry as a sensitive and specific diagnostic method for each subtype.

The gross morphology is similar to other RCCs; they do not show a distinct appearance (1). Histopathologically, Xp11 translocation RCC is characterized by papillary structures lined by cells with large, usually clear, sometimes eosinophilic cytoplasm (Figure 1). Immunohistochemically, these tumors are usually negative for epithelial markers [pancytokeratins, epithelial membrane antigen (EMA)], pax-8, and positive for cathepsin K and melanocytic markers (HMB-45 and Melan-A) (2).

The differential diagnosis of MiT family translocation RCCs includes a variety of renal neoplasm demonstrating clear cell and papillary features. Conventional clear cell RCC demonstrates diffuse positive expression with epithelial markers and is negative with melanocytic markers. Papillary RCC is positive for cytokeratin 7 and alpha-methylacyl-CoA racemase. In addition, clear cell RCC is characterized by chromosome 3p25 deletion, papillary RCCs by trisomy 7/17. Some of the Xp11 translocation carcinomas include melanin pigment (5). Therefore, epithelioid angiomyolipomas should be considered in the differential diagnosis. Since TFE3 and TFEB gene rearrangements are not present in other neoplasms, fluorescent in situ hybridization can also be useful and helpful in diagnostically challenging cases.

Figure 1. A Xp11 translocation carcinoma having cells with prominent large clear cytoplasms lining papillary structures (x200, Hematoxylin&Eosin) Inset: TFE3 nuclear positivity (x200)
The prognosis of the Xp11 translocation RCCs is similar to clear cell RCCs but worse than PRCCs. t(6;11) RCCs are indolent than Xp11 translocation RCCs. To be aware of these neoplasms on microscope would provide correct diagnosis and predicting of prognosis.

Ethics
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