Wilms’ Tumor in Horseshoe Kidney

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Keywords
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Abstract
Wilms’ tumor is the most common malignant kidney tumor found in children. The Horseshoe kidney is the most common renal fusion malformation. However, Wilms’ tumor is rarely identified in horseshoe kidney patients. Multimodal treatments in Wilms’ tumor can play important roles in increasing the survival rate. In this study, we report the case of a 6-year-old boy in whom a Wilms’ tumor was identified in a horseshoe kidney. The tumor was successfully treated with preoperative chemotherapy, followed by surgical resection.

Introduction

Nephroblastoma, also known as Wilms’ tumor, is the most commonly identified pediatric renal mass, accounting for 87% of all renal masses and representing 7% of all malignant tumors identified in children [1]. The median age at which this tumor is identified is 3 years [2]. Wilms’ tumor has been associated with a number of syndromes, including WAGR syndrome (Wilms’ tumor, aniridia, genitourinary anomalies, and range of developmental delays), Beckwith-Wiedemann syndrome, Denys-Drash syndrome, and Edwards or Perlman syndrome [3]. Horseshoe kidney occurs in approximately 1 in 500 cases, with a male
preponderance [4]. However, the occurrence of Wilms’ tumor in horseshoe kidney is outstandingly rare, with an estimated incidence of approximately 0.4–0.9% of all Wilms’ tumors [5]. In this report, we describe a rare case of Wilms’ tumor identified in a horseshoe kidney to highlight the effectiveness of using preoperative chemotherapy during the treatment of this tumor.

**Case Report**

A 6-year-old boy presented to the hospital due to left upper abdominal pain after a falling accident. The patient’s blood pressure and pulse were within the normal ranges. A physical examination revealed a large mass on the left upper abdomen. Laboratory studies demonstrated that complete blood counts, liver function tests, renal function tests, and urinalysis results were normal. An abdominal computed tomography (CT) scan was performed, which showed the existence of an isthmus connecting the right and left kidneys, anterior to the aorta and inferior vena cava. A large mass was observed, measuring 7 × 8 cm, located in the isthmus of the horseshoe kidney, which primarily developed toward the left side of the abdomen (Fig. 1a). This mass showed heterogeneous enhancement with less enhancement relative to the normal kidney parenchyma (Fig. 1a, arrowhead). Abdominal CT image after 12 weeks of chemotherapy treatment revealed a significant reduction in tumor size (Fig. 1b, arrowhead). CT, computed tomography.

**Fig. 1.** Abdominal CT images revealed the presence of a large mass located in the isthmus of the horseshoe kidney, which primarily developed toward the left side of the abdomen (a, arrow). This mass showed heterogeneous enhancement, with less enhancement relative to the normal kidney parenchyma (a, arrowhead). Abdominal CT image after 12 weeks of chemotherapy treatment revealed a significant reduction in tumor size (b, arrowhead). CT, computed tomography.
Discussion

During the fetal period, the kidneys develop and ascend, developing first in the pelvis and then gradually ascending into position below the thorax, on either side of the lumbar spine [6]. During this ascending process, the kidneys also rotate, which typically occurs by the gestational ninth week [6]. Renal fusion anomalies may occur during this process [7]. The isthmus of the horseshoe kidney may contain functioning renal parenchyma or a fibrous band [8]. In up to 80% of cases of horseshoe kidney, the isthmus contains functional renal parenchymatissue, and in >90% of cases, fusion occurs at the lower pole [6]. Patients with horseshoe kidney are often asymptomatic and are typically discovered incidentally, often due to symptoms secondary to pelvic ureteric junction obstruction and infection [9]. These patients are thought to be at increased risk of developing malignancies, such as renal cell carcinoma, Wilms’ tumor, and carcinoids, among which renal cell carcinoma is the most common [7]. However, Wilms’ tumor is the most common malignant kidney tumor identified in children [10]. The risk of Wilms’ tumor in children with horseshoe kidney is 2–6 times that of children in the general population [11]. Approximately 50% of Wilms’ tumors in horseshoe kidney develop from the isthmus, likely due to the abnormal proliferation of metanephric blastema in the isthmus [7]. The same anomaly that causes the development of horseshoe kidney may also lead to the development of Wilms’ tumor [12]. Patients with Wilms’ tumors are often asymptomatic; approximately 10% are discovered incidentally after trauma, whereas 25% present with microscopic hematuria or hypertension secondary to renin production [13]. Ultrasound is used to diagnose horseshoe kidney, whereas CT and magnetic resonance imaging are often used for staging purposes [14]. On ultrasound, the mass presents as a large renal mass, which can be either solid or cystic, with large hypoechoic areas due to central necrosis and cyst formation [13]. Areas characterized by fat deposits, calcification, or hemorrhage may appear [13]. On CT, the tumors are lower density and enhance less than the normal renal parenchyma [15]. Tumors are often characterized by heterogeneous contrast enhancement and may feature punctuated calcifications [15]. On magnetic resonance imaging, the tumors have low signal intensity on T1-weighted images, with either low or high signal intensity on T2-weighted images and restricted diffusivity on diffusion-weighted images [13]. CT is also used for the detection of lung metastasis or local recurrence [13]. Wilms’ tumors contain variable quantities of embryonic renal elements, such as blastema, epithelium, and stroma [16]. Wilms’ tumor can be divided into 2 types, based on prognosis: favorable (over 90%) and unfavorable (6–10%) [13]. Histopathological analysis is the current gold standard.
standard for diagnosing Wilms’ tumor. Surgery, chemotherapy, and radiotherapy are also used to treat Wilms’ tumor [10]. The National Wilms Tumor Study Group (NWTS)/Children’s Oncology Group (COG) and the International Society of Paediatric Oncology (SIOP) have established the major guidelines regarding the management of Wilms’ tumor [17]. SIOP recommends using preoperative chemotherapy to reduce the tumor size and prevent intraoperative spillage due to tumor rupture [10]. In contrast, the NWTS/COG recommends the application of primary surgery before any adjuvant treatments [17]. The overall survival of children with Wilms’ tumor in horseshoe kidney appears to be similar to that among children with Wilms’ tumor in normal kidneys [5].

Although multiple guidelines may be used for the management of Wilms’ tumor, this patient was treated with chemotherapy to reduce the tumor size before surgery. Because the tumor arose from the isthmus and primarily developed to the left, the left kidney, isthmus, and tumor were all removed completely. After 2 years of follow-up, no evidence of tumor recurrence or metastasis was observed.

**Conclusion**

Horseshoe kidney is a common congenital kidney anomaly. Patients with horseshoe kidney have an increased risk of Wilms’ tumor than patients with normal kidneys. Imaging plays a pivotal role in the diagnosis and follow-up posttreatment. Neoadjuvant chemotherapy can reduce the size of the tumor prior to surgery while maintaining normal kidney parenchyma.

**Statement of Ethics**

All treatments and examinations followed the guidance of the Declaration of Helsinki. Informed consent for treatment was obtained from the legal guardian of the patient. The legal guardian of the patient has given written informed consent to publish the case, including the publication of images.

**Conflict of Interest Statement**

The authors have no conflicts of interest to declare.

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**Author Contributions**

Doan Tien Luu and Nguyen Minh Duc contributed equally to this article as co-first authors. Doan Tien Luu, Nguyen Minh Duc, and Thieu-Thi Tra My contributed to the acquisition of data and writing of the manuscript. Nguyen Minh Duc and Thieu-Thi Tra My provided supervision, assessment, and interpretation of data, along with mentorship. All authors approved the final manuscript.
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