AMSER Case of the Month
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2 year old male with abdominal mass

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Patient Presentation

- **HPI:** 2 year old male with no PMH presented to outside ED with abdominal mass found on routine checkup by pediatrician. A mass was noted in the left upper quadrant on exam. The outside ED completed an ultrasound which showed a 13.5 cm x 12 cm x 9 cm mass, and transferred the patient for further workup.
- **Medical and Surgical History:** None
- **Family History:** Mesothelioma in paternal grandmother. Melanoma on both sides of family. History of HTN and TIA on father’s side of family.

**Physical Exam**
- Vitals: T: 36.9 BP: 112/68 HR: Unable to Record RR: 24 SpO2: 100%
- General: Alert, NAD
- Skin: Warm, intact
- Cardiac: Regular rate and rhythm, no murmurs
- Respiratory: Clear to auscultation bilaterally
- GI: Soft, nondistended, non tender, *mass in LUQ that does not cross midline*
- MSK: Normal ROM, no swelling

Labs were remarkable for a BUN of 22 and an LDH of 444.
What Imaging Should We Order?
# ACR Appropriateness Criteria


<table>
<thead>
<tr>
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<th>Relative Radiation Level</th>
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<tr>
<td>CT abdomen with IV contrast</td>
<td>Usually Appropriate</td>
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This was ordered by the outside hospital emergency room physician - dopplers were repeated by the ED at our facility.
Findings (Unlabeled US)
Findings (Labeled US)

Solid left renal mass
Patent left renal vein
IVC without tumor thrombus

IVC without tumor thrombus
### ACR Appropriateness Criteria

**Variant 1:**

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This was ordered by the pediatric heme/onc team upon admission.
Findings - Unlabeled CT (Coronal)
Findings - Labeled CT (Coronal)

Left renal heterogeneously enhancing soft tissue mass

Claw sign indicating mass originates from left kidney

Patent IVC without thrombus
Findings - Unlabeled CT (Axial)
Findings - Labeled CT (Axial)

- Normal right kidney
- Mass effect on stomach
- Patent left renal vein
- Heterogeneously enhancing soft tissue mass
- Left kidney
- Aorta displaced to right
Post-Op Axial CT (unlabeled)
Post Left Radical Nephrectomy Axial CT (Labeled)

Surgical clips
Abdominal Mass Differential Diagnosis

- Wilms tumor
- Neuroblastoma
- Mesoblastic nephroma
- Renal cell carcinoma
- Renal Rhabdoid Tumor
- Clear cell sarcoma
- Renal medullary carcinoma
Final Dx:

Wilms Tumor
Case Timeline

- Abdominal US was performed at outside hospital with concern for neuroblastoma versus Wilms tumor. Patient was transferred for further workup
- Abdominal US with dopplers was performed at VCU ED, which showed no tumor extension into the renal vein or IVC. Chest, abdomen and pelvis CT with contrast was ordered. Chest CT was unremarkable. Abdominal CT showed mass arising from left kidney
- Left radical nephrectomy was performed
- Follow up CT showed post-surgical changes
Case Discussion

**Epidemiology and Presentation**
- Accounts for 87% of pediatric renal masses
- Peak incidence of 3-4 years of age with 80% of patients presenting before 5 years of age
- Can be bilateral in 4-13% of children
- May be associated with other congenital anomalies such as cryptorchidism, hemihypertrophy, hypospadias, and sporadic aniridia
- Discovery of tumor most commonly follows detection of palpable mass
- Hematuria and pain are infrequent clinical findings, and hypertension can be noted in 25% of cases

**Features**
- Tumor arises from mesodermal precursors of renal parenchyma
- Solid intrarenal mass with pseudocapsule and distortion of renal parenchyma and collecting system
- Tumor typically spreads by extension and displaces adjacent structures but does not elevate or encase the aorta (which would be a distinguishing feature of a neuroblastoma)
- May demonstrate vascular invasion of renal vein and inferior vena cava with occasional extension into right atrium
- Metastases are most commonly found in the lungs, liver and regional lymph nodes
Case Discussion

Radiographic Features
- Often very large on presentation and displaces adjacent structures without insinuating between them
- Initial evaluation on ultrasound the tumor has heterogeneous echogenicity representing hemorrhage, fat, necrosis or calcification
- Examination of IVC using dopplers on ultrasound is crucial to detect tumor extension
- CT demonstrates heterogeneous soft tissue mass with infrequent areas of calcification and fat density regions
- Claw sign on CT can help differentiate Wilms tumor from neuroblastoma
- Important to assess contralateral kidney for presence of disease
- 10-20% of cases have lung metastases found on CT at the time of diagnosis, so it is important to complete a chest, abdomen and pelvis CT with contrast in order to evaluate for metastases
- While MRI is the most sensitive modality for evaluating caval patency, it may require sedation in many children and so surgical planning can be done using US and CT
- If MRI is completed, T1 imaging shows heterogeneous hypointensity and T2 imaging shows heterogeneous hyperintensity
Case Discussion

**Treatment**
- Unilateral Wilms tumors are treated by a combination of nephrectomy and chemotherapy
- Occasionally, especially in cases where tumors are bilateral, chemotherapy can precede surgery
- Radiotherapy has a limited role in treatment
- Cure is possible in 90% of cases
- Recurrence can be seen in the tumor bed or with metastases in the lungs or liver
References


