A rare cause of renal atrophy: Subcapsular collection presented as a huge perirenal complex cyst

Renal atrofinin nadir bir sebebi: Dev perirenal kompleks kist şeklinde prezente olan subkapsüler koleksiyon

Subcapsular collection presented as perirenal cyst

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Abstract
If left untreated, renal subcapsular collections may result in renal atrophy. The diagnosis could be challenging at chronic stage due to recurrent episodes of hemorrhage which is the reason for the complex appearance of the collection. Herein we report a rare cause of renal atrophy in a 44-year-old man.

Keywords
Atrophic Kidney; Complex Cyst; Subcapsular Collection; MRI; CT

Öz
Renal subkapsüler koleksiyonlar tedavi edilmezse renal atrofi ile sonuçlanabilir; Tekrarlayan hemorajı epizodları koleksiyonun karmaşık görünümü olması neden olduğu için kronik dönemde tani koymak oldukça zor olabilir. Biz bu yazımızda 44 yaşındaki bir erkek olguda böbrek atrofisinin nadir bir sebebinin sunuyoruz.

Anahtar Kelimeler
Atrofik Böbrek; Kompleks Kist; Subkapsüler Koleksiyon; MRG; BT

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Introduction
The contents of the renal subcapsular collections may vary depending on the etiology [1,2]. Subcapsular collections may induce significant pressure on renal parenchyma due to boundary effect of renal capsule. In the setting of renal vessel compression, one may identify the new onset of increased blood pressure due to activation of the renin-angiotensin system, which is also called as Page kidney [2,3]. However, in the case of long standing compression, the kidney may undergo atrophy and imaging based evaluation could be challenging. Herein we report a rare case of chronic renal subcapsular collection presented as a huge complex cystic mass resulting in significant right kidney atrophy.

Case Report
A previously healthy 44-year-old man was admitted to our hospital with a two days history of right back pain radiating through to right inguinal region. He was afebrile, and his blood pressure was within the normal range. There was no history of trauma. The urinary test revealed lack of hematuria. Serum creatinine level was normal (0.76 mg/dL). Abdominal radiography demonstrated a calcified ovoid mass in right flank (Figure 1). Ultrasonography revealed the cystic nature of the lesion with fluid-fluid level (Figure 2). Abdominal CT examination demonstrated a peripherally calcified, huge (sized 10 x 13 cm) cystic mass in right renal fossa (Figure 3). The right kidney was absent however there was a non-specific soft tissue density with a few punctate calcifications between the cystic mass and inferior vena cava (Figure 3). The left kidney was unremarkable except for hypertrophy. The right adrenal gland and urinary bladder were normal. Contrast-enhanced MRI examination was suggestive of right renal atrophy, however, neither renal artery nor renal vein were apparent on postcontrast images. The extension of the cyst was indicative of subcapsular origin (Figure 4). The patient underwent surgery, and histopathological examination revealed collagenized and calcified cyst wall associated with atrophic renal parenchyma and pelvicalyceal system. There was no sign of granulomatous reaction in histopathological examination, and also acid-fast bacillus staining was negative for tuberculosis. Urine culture test and chest radiograph were also unremarkable.
Discussion

Herein reported case indicates that renal subcapsular collections may result in significant parenchymal atrophy at late stage. Compensatory hypertrophy of the contralateral kidney also suggests chronic state of the insult. The main limitation for further conclusion is that we were not able to evaluate previous radiological examinations of the patient. The patient’s presenting symptom was newly onset of back pain. Although back pain is a nonspecific complaint that could be seen in various conditions, one may suspect complication of an existing cyst. However pathological examination revealed lack of such association as well. The presenting symptom could be related to another cause (i.e., lumbar disc hernia or myalgia), and the renal atrophy could be incidentally diagnosed.

A hematoma is the most frequent type of renal subcapsular collection and may occur following trauma, anticoagulant therapy, and interventional procedures [1, 4-6]. Renal subcapsular pseudocyst may also be encountered as a complication of acute pancreatitis [3]. A urinoma can also be the underlying cause of subcapsular collection particularly in the setting of obstructive uropathy (i.e., ureteral stone, retroperitoneal fibrosis, bladder cancer) complicated by fornical rupture [7,8]. On the other hand, trauma may also result in subcapsular urinoma as well [2]. In case of bilateral renal subcapsular collection presence of renal parenchymal diseases (i.e., membranoproliferative and focal-segmental glomerulonephritis) could be questioned [9,10]. In our case, neither imaging findings nor clinicopathological signs were able to reveal the underlying cause of renal subcapsular collection.

Lin et al. reported a case of tuberculosis autonephrectomy presented with a huge cystic mass with lack of normal appearing right kidney [11]. The cyst had calcified wall, and imaging findings were similar to our case. However, the urine culture test was positive for Mycobacterium tuberculosis in their case which is contrary to our findings. Moreover, the presence of compressed and atrophic kidney seen in our case is unusual for tuberculosis autonephrectomy.

In conclusion, renal subcapsular collections may result in renal atrophy, and the underlying cause may remain unknown particularly in patients presented at late stage. Presence of atrophic kidney adjacent to cystic mass indicates capsular-subcapsular origin of the cyst.

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Author contributions


Informed Consent

Written informed consent was obtained from the patient who participated in this study.

Conflict of Interest

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References


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