Isolated omental metastasis of renal cell carcinoma after extraperitoneal open partial nephrectomy: A case report

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A B S T R A C T

INTRODUCTION: Metachronous metastatic spread of clinically localized renal cell carcinoma (RCC) affects almost 1/3 of the patients. They occur most frequently in lung, liver, bone and brain. Isolated omental metastasis of RCC has not been reported so far.

CASE PRESENTATION: A 62-year-old patient previously diagnosed and treated due to pulmonary sarcoidosis has developed an omental metastatic lesion 13 years after having undergone open extraperitoneal partial nephrectomy for T1 clear-cell RCC. Constitutional symptoms and imaging findings that were attributed to the presence of a sarcomatoid paraneoplastic syndrome triggered by the development this metastatic focus complicated the diagnostic work-up. Biopsy of the [18F]-fluorodeoxyglucose (+) lesions confirmed the diagnosis of metastatic RCC and the patient was managed by the resection of the omental mass via near-total omentectomy followed by targeted therapy with a tyrosine kinase inhibitor.

DISCUSSION: Late recurrence of RCC has been reported to occur in 10–20% of the patients within 20 years. Therefore lifelong follow up of RCC has been advocated by some authors. Diffuse peritoneal metastases have been reported in certain RCC subtypes with adverse histopathological features. However, isolated omental metastasis without any sign of peritoneal involvement is an extremely rare condition.

CONCLUSION: To our knowledge, this is the first reported case of metachronously developed, isolated omental metastasis of an initially T1 clear-cell RCC. Constitutional symptoms, despite a long interval since nephrectomy, should raise the possibility of a paraneoplastic syndrome being associated with metastatic RCC. Morphological and molecular imaging studies together with histopathological documentation will be diagnostic.© 2016 The Authors. Published by Elsevier Ltd. on behalf of IJS Publishing Group Ltd. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).

1. Introduction

The incidence of renal cell carcinoma (RCC) has been rising, which includes both early stage and late stage disease [1]. Approximately 85% of all RCCs are of clear cell histology [2]. About 20–30% of patients have metastatic disease at the time of diagnosis, and about 20–40% of patients with clinically localized disease at diagnosis will eventually develop metastases [3]. RCC often metastasizes to the lung, bone, liver and brain. Herein, we report a case with isolated omental metastasis of RCC that has developed metachronously 13 years after the initially localized disease was managed by partial nephrectomy.

2. Description of the case

A currently 62-year-old male presented with decreased force of urinary stream, frequency, malaise, loss of apetite, difficulty in breathing and nonproductive cough. He was treated due to pulmonary sarcoidosis in the past and apart from hypertension he did not have any systemic comorbidity. He was using bronchodilators on demand due to airway-related problems and he received corticosteroids due to sarcoidosis in the past. He had undergone open extraperitoneal partial nephrectomy elsewhere due to T1 clear-cell RCC, 13 years ago. He also reported transurethral resection of the prostate (TURP) and consequent office-based urethral dilatation

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procedures due to post-TURP urethral stricture. He had complied with the postoperative surveillance protocol of RCC and until 2013 there was no sign of recurrence or metastasis.

Imaging workup began with an abdominal ultrasound, demonstrating a solid round mass superolateral to the urinary bladder with nonspecific heterogeneous echotexture. Magnetic resonance imaging (MRI) confirmed this mass as a well-circumscribed structure abutting the superolateral urinary bladder, with a thin but preserved intervening fat plane between it and the urinary bladder (Fig. 1). For further staging, [18F]-fluorodeoxyglucose (FDG) positron emission tomography–computed tomography (PET-CT) scan was undertaken, demonstrating the index supravesical lesion to be FDG-hypermetabolic, as well as innumerable FDG-hypermetabolic lesions with an osseous, renal, pleural, and lymphatic distribution (Fig. 2). Percutaneous biopsy was directed to both the index supravesical lesion as well as the most hypermetabolic extravesical site (right iliac bone lesion). Histopathology of the supravesical specimen revealed clear cell RCC (representing delayed isolated omental metastasis) while the osseous specimen was interpreted as sarcomatoid reaction (Fig. 3).

Then, we made a consultation with the Pulmonary Medicine department and the advice given by them was to remove the metastatic lesion since widespread FDG (+) lesions and accompanying constitutional symptoms may be a sort of paraneoplastic syndrome, which can be seen quite often in RCC. We opted for an open surgery to remove the mass and during exploration it was found to be located within the omentum as a clearly palpable, isolated nodularity (Fig. 4). Neither the intestinal segments nor the peritoneum seemed to be affected by the metastatic spread. We performed a near-total omentectomy and excised the mass with clear margins. In the same session, we performed direct visual endoscopic internal urethrotomy due to bulbomembranous urethral stricture which seemed explanatory for the emptying phase lower urinary tract symptoms (LUTS). Histopathological examination findings in the omental mass were concordant with the tru-cut biopsy result (Fig. 5). After having the opinions of Medical Oncology and Pulmonary Medicine specialists, the patient was scheduled for adjuvant targeted therapy with a tyrosine kinase inhibitor (sunitinib).

3. Discussion

Late recurrence is one of the biological behaviors of RCC. It has been reported that the recurrence rate 5 years after surgery is 8.8% and the rate after 10 years is 11% [4,5]. A previous study of patients with RCC who did not develop recurrence within a 10-year follow-up observation period reported that late recurrence rates were 10.5 and 21.6% at 15 and 20 years, respectively [6]. Therefore, some authors have highlighted the importance of postoperative surveillance and recommended life-long follow-up of RCC after the initial treatment [6]. Supporting this phenomenon; our case, who had pT1 clear cell RCC at the time of diagnosis, has developed an isolated omental metastasis 13 years following partial nephrectomy.

Some aggressive RCC subtypes with adverse histopathological features (sarcomatoid differentiation etc.) can present with diffuse peritoneal metastases involving the omentum [7,8]. However, development of metachronous omental metastasis(es), particularly in the absence of diffuse peritoneal involvement, is an extremely rare condition in the natural history of RCC. Win et al. have recently reported a patient with an initially T3N0M0 chromophobe RCC, who developed an interaortocaval lymph node enlargement 1 year after laparoscopic radical nephrectomy [9]. They have incidentally discovered an additional omental metastatic deposit while removing this interaortocaval mass. Despite surgical excision and adjuvant therapy with tyrosine kinase inhibitors, the omental nod-
ules recurred and they were able to detect this recurrence via FDG avidity on surveillance PET/CT studies [9].

RCC is unique for its association with various paraneoplastic syndromes. These can range from those manifesting as constitutional symptoms (ie., fever, cachexia, and weight loss) to those that result in specific metabolic and biochemical abnormalities (ie., hypercalcemia, nonmetastatic hepatic dysfunction, amyloidosis, etc.). The constitutional symptoms of our patient was initially attributed to the progression of sarcoidosis. However, disseminated osseous involvement is quite rare in sarcoidosis and in the absence histological proof of non-caseating granulomata on biopsy specimens it would not be possible to conclude upon the diagnosis of sarcoidosis. Moreover, the probability of metastatic dissemination must be evaluated in a patient who has been operated due to localized RCC. On the other hand, Willis et al. have reported that sarcoidosis and RCC can coexist in the same individual. In their case, the undiagnosed condition of sarcoidosis complicated the staging of bilateral RCC. Prior to the initiation of immunotherapy due to the initial diagnosis of mRCC, they also performed an FDG PET/CT scan and the difference in metabolic activity between the primary site of disease and the presumed metastatic disease suggested the possibility of two separate disease processes. The patient underwent bilateral partial nephrectomies and was ultimately diagnosed with stage T1 bilateral RCC. Demonstration of non-caseating granulomas on pulmonary nodules confirmed the diagnosis of coexistent sarcoidosis in this case [10].

This patient underwent extensive imaging work-up in an effort to exclude RCC metastasis, however the summary of noninvasive studies provided equivocal results. For example, initial ultrasound and MRI findings revealed the solitary index lesion (supravesical mass). However, the subsequent FDG PET/CT scan demonstrated the FDG-hypermetabolic index lesion but also innumerable disseminated FGD-hypermetabolic foci. Moreover, FDG PET-CT is known to carry some limitation in the evaluation of RCC (unless the RCC is high grade). Therefore, combining the known pulmonary and surgical history with the patient’s delayed presentation and dramatic PET-CT findings in the context of benign-appearing renal fossae, tissue sampling was deemed necessary to guide the next steps in care. Eventually, we thought that most of the presenting symptoms, except emptying phase LUTS, and the FDG-hypermetabolic lesions should be considered as a component of a sarcomatoid paraneoplastic syndrome triggered by metastatic progression of an initially localized RCC.

Surgical resection of the primary tumor and metastatic deposits continues to play an important role in managing patients with mRCC when aiming for complete remissions, the likelihood of which is further supplemented with the addition of systemic targeted therapy. Therefore, after the removal of omental metastasis, our patient was scheduled to receive targeted therapy with sunitinib.

4. Conclusions

To our knowledge, this is the first reported case of metachronous isolated omental metastasis of an initially T1 clear-cell RCC. Metastatic spread can develop many years after the surgical
Fig. 3. (A) Histopathology of the suprapubic mass demonstrate a neoplasm composed of cells with clear vacuolated cytoplasm and hyperchromatic small nuclei (H&E, ×90). (B) Immunohistochemically, tumor cells show nuclear positivity with PAX8 (IHC stain ×150). (C) Immunohistochemically, tumor cells show strong, diffuse positivity with CD10 (IHC stain ×150). (D) H&E stained sections of the needle biopsy from the iliac bone showing granulomatous inflammation (×150).
treatment of localized RCC. Constitutional symptoms may be a sign of paraneoplastic syndrome that was triggered by RCC progression. Coexistent medical conditions, such as sarcoidosis as in our case, may complicate the diagnostic process. Morphological and functional imaging studies together with histopathological assessment of biopsy materials are necessary to confirm the diagnosis of mRCC and rule out other systemic disease processes.

**Conflicts of interest**

None.

**Funding**

None.

**Consent**

Written informed consent was obtained from the patient for publication of this case report and accompanying images.

**Author contribution**

Ömer Acar has written the manuscript. The operation was carried out by Tuna Mut, Ömer Acar and Tarik Esen. Pathology was reported by Yeşim Sağlıcan. Fatih Selçukbiricik, Levent Tabak have contributed to the clinical management of the patient. The radiological imaging studies were reported and managed by Alan Alper Sag, Okan Falay.

**Guarantor**

Tarik Esen.

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