Case report of a patient with a rare location of renal cell carcinoma metastasis

Prikaz slučaja pacijentice s rijetkom lokacijom metastaze karcinoma bubrega

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Abstract. Aim: Renal cell carcinoma is one of the deadliest cancers which takes the third place among malignant carcinomas of the genitourinary tract. This case report describes the rare case of a large fast-growing metastasis of renal cell cancer in an unusual location with the short time of its appearance, but with an unexpected clinical course after its treatment.

Case report: A 60-year-old female patient presented with a fist-sized formation on the scalp. During physical examination, the mass was determined to be firm, infiltrative, fixed, and painless on palpation. The formation had appeared a month prior to arrival, characterized by fast growth. The patient underwent a radical nephrectomy six months earlier due to RCC. A cranial radiogram showed osteolysis of the neurocranium, while an MRI of the brain confirmed the mass belonged to the scalp, no brain infiltration, measuring 82 × 60 × 74 mm. Fine needle aspiration isolated malignant cells of an adenocarcinoma, origin undetermined. The patient underwent surgery to remove the mass along with the effected neurocranium, and the resulting bone defect was reconstructed using Palacos. Postoperative recovery went well, and the pathohistological analysis confirmed renal cancer metastasis. PET-CT didn’t show suspicious lesions for distal metastasis for a period of 2 years. Conclusion: Renal cell carcinoma commonly metastasizes to the lung. This case report presents the rare case of a large fast-growing renal cancer metastasis to an unusual location, with a short time for metastasis appearance. In spite of that we have shown two years follow up of remission.

Key words: neoplasm metastasis; osteolysis; renal cancer

Sažetak. Cilj: Karcinom bubrega zauzima treće mjesto među malignim tumorima genitouri
narnog trakta. Cilj je ovog prikaza slučaja je opisati rijedak slučaj gigantske i brzorastuće meta-
staze ovog karcinoma te rijetko mjesto njezинog sijela s kratkim vremenom njezīnog pojavljava-
nja i neocljekinih kliničkim tijekom nakon njezīnog lečenja. Prikaz slučaja: 60-godišnje žena do-
lazi zbog tvorbe u predjelu mekog oglavka glave veličine šake. Prilikom pregleda tvorba se očitovala kao tvrda infiltrativna, fiksirana za podlogu i palpatorno bezbolna. Tvorba se pojavila mjesec dana pred dolazak te se očitovala vrlo brzinom rastom i napredovanjem. Kod pacijentice je šest mjeseci unatrag bila učinjena radikalna nefrectomija zbog karcinoma bubrega. Kranigram je pokazao osteolizu skeleta neurokranijuma, a MR mozga potvrdio navedeno. Tvorba je pripadala mekom oglašku glave bez infiltracije parenhima mozga i mjerila je 82 × 60 × 74 mm. Citološkom punkcijom nađene su maligne stanice adenokarcinoma, ali nije razjašnjeno njihovo podrijetlo. Pacijentica je bila podvrgnuta neurokromatološkoj operaciji kojom je tvorba izrezbijena i rekonstruirana uključivši palakos. Postoperativni tijek protokolo je uredno, a PH nalaz pokazao je da se radilo o metastazi karcinoma bubrega. PET-CT nije pokazao udaljene metastaze u periodu praćenja od dvije godine. Zakočen je u podvrgnutoj operaciji karcinoma bubrega. U praćenju ne došlo do obnavljanja metastaze. Unatoč tome kliničkim praćenjem prikazan je period dvije godine remisije.

Ključne riječi: karcinom bubrega; metastaza neoplazme; osteoliza

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INTRODUCTION

About 2-3% of malignant tumors in patients are renal cell carcinomas (RCC), representing the third most common genitourinary cancer following bladder and prostate cancers, and the fifth most common cancer in men1,2. RCC is among one of the most lethal urological cancers, along with bladder cancer. Incidence of RCC increases 2.5% annually, due to the frequent use of non-invasive procedures, as well as the influence of genetic and environmental factors3,4.

This case report has shown the patient with rare, large, fast-growing metastasis of renal cancer in an unusual location, demonstrating its aggressiveness and high grade. MRI of the brain confirmed the mass belonged to the scalp with bone osteolysis, but no brain infiltration.

Recent case reports claim 60-70% of patients with RCC are asymptomatic when diagnosed5. The majority of tumors discovered are in an early stage of the disease, and can be treated surgically. Nephrectomy is the gold standard in the treatment of disease that is clinically limited to the kidney, however, due to the aggressive nature and malignancy of RCC, recurrence rates can be as high as 40% after nephrectomy for localized disease6. Approximately 20-30% of patients have metastatic disease at the time of diagnosis, while 30% of patients after radical surgery will develop a metastatic disease. Less than 4% of patients have bilateral tumors, usually within hereditary syndromes7,8. Survival rate is well correlated with the TNM stage of disease, serving as the basis for RCC prognosis. Congruently, the 5-year survival rate from stages I to IV decreases respectively from 91%, 74%, 67%, to only 32% for stage IV, in other words, a major decrease in the 5-year survival rate is seen between stage III and IV9. RCC is prone to metastasize to almost all soft tissues, with the highest incidence developing in the lung, making up about 16% of all metastatic localization, followed by bone in about 2-8% of cases, liver in a maximum of 7%, and the brain in only 4% of all secondary sites of RCC10. Resection is preferred for solitary lung metastasis, resulting in 5-year survival rates reaching 54%11, resection is also preferred for loco-
fat, as well as the retroperitoneal lymph node among the tissue received. The tumor was localized at the apical and mesorenal parts of the kidney, and infiltrated the renal sinus and perirenal fat. Based on a cross section, the tumor was described as soft in consistency, yellowish, partially cystic and solid, measuring about $13 \times 10 \times 9$ cm, with signs of central hemorrhage and necrosis. Histologically, based on the World Health Organization (WHO) Classification of Tumours on Pathology and Genetics of Tumours of the Urinary System and Male Genital Organs, 2004, the tumor cells were determined to originate from renal cells, more precisely identified as a clear cell type renal carcinoma. The tumor was defined as grade II based on the nuclear grading system by Fuhrman. A pseudocapsule was present, which was infiltrated, along with the surrounding fat tissue reaching Gerota’s fascia. Microvascular invasion of the tumor was present. It was noted that the tumor did not infiltrate the large muscle veins, nor the renal vein. The surgical margin of the tumor was clear, as well as the renal artery and vein, and the ureter. The adrenal gland presented normal structure. Two lymph nodes found in the perirenal fat tissue and the extirpated retroperitoneal lymph node were completely infiltrated by tumor cells. The patient returned six months later when a suspicious tumor was noticed on the head in the area above the coronal suture of the skull on the left. A cranial radiogram was immediately taken, the report described an area of osteolysis or destruction of the parietal bone, measuring about $5 \times 3$ cm, with irregular edges and intense periosteal reaction. Magnetic resonance imaging (MRI) of the brain confirmed the presence of a subcutaneous expansive process measuring $82 \times 60 \times 74$ mm above the coronal suture of the skull on the left with osteolysis and epidural propagation, without infiltration of the brain. All other findings were normal (Figures 1, 2). These findings were highly suspect for metastasis of the earlier operated RCC and fine needle aspiration (FNA) confirmed RCC metastasis. After imaging, the patient was hospitalized for the second time at the Department of Neurosurgery for the planned surgical treatment of the neurocranial tumor, or solitary RCC metastasis. Neurological status of the patient was normal, and there were no contraindications for surgical treatment. The patient was operated in the supine position with the head in anteflexion at about 20 degrees and fixed in a Mayfield head clamp (Figure 3). The skin covering the tumor was excised in a “tear-drop” shape revealing a limited soft tissue mass which was removed and sent for pathohistological analysis (Figure 4). After removing the tumor, partially destroyed frontal and parietal bones to the left of the median line were noticed, and a frontoparietal craniectomy was performed, removing the effected bone, which was sent for pathohistological analysis as well (Figure 5). Dura mater was...
After surgery the final pathohistological examination confirmed renal cancer metastasis. After imaging no suspicious lesions have been shown for a follow-up period of 2 years. It should be noted that every lesion should be suspected as metastasis in renal cancer patients due to its aggressiveness and unpredictable behavior.

The tissue received earlier identifying RCC, clear cell type, grade III.

Postoperative recovery passed without complications. The surgical wound healed per primam without complications. The patient was discharged on the seventh postoperative day in good overall health. The surgical drain was removed on the tenth postoperative day due to the prolonged drainage from the wound, along with the stitches. A thoracic radiogram and positron emission tomography combined with computed tomography (PET-CT) were ordered to assess for distal metastases, neither of which showed any suspicious lesions in the lungs or other organs for a follow-up period of 2 years.

DISCUSSION

RCC is a complex cancer that requires a complex approach, namely due to its aggressive and unpredictable behavior, and a propensity to metastasize\textsuperscript{14}. Rare or unusual locations of RCC metastases are defined as the sites anatomically distant from the kidney or outside of the typical path of metastatic spread of renal tumors. When breaking down the occurrences of RCC metastatic locations to percentages, the numbers vary considerably, as a result, there are several case reports presenting rare metastatic sites\textsuperscript{15}. The mean time for appearance of metastasis of RCC is 1-2 years after operation\textsuperscript{16} although in the literature there is a case report with record lung metastasis from RCC 37 years after nephrectomy\textsuperscript{17}. When compared to this
our patient had a very early appearance of metastatic disease even received sunitinib treatment that should decrease the relapse of disease\textsuperscript{18}. On the other side there were very interesting two years of remission after radical resection of solitary metastatic lesion with no signs for distal metastatic disease in PET-CT examination\textsuperscript{19}. Our patient underwent radical nephrectomy, after which intact surgical margins were confirmed, as well as adrenal gland and ureter. The possibility of local recurrence of disease was considerably greater in this case than the occurrence of distant metastases due to pathohistologically confirmed capsule infiltration and lymphovascular invasion with positive lymph nodes\textsuperscript{8}. In spite of this, in the six months following nephrectomy, the patient developed a large solitary RCC metastasis in the subcutaneous tissue of the scalp with osteolysis and periostatic reaction of the underlying cranium, which is the first such case seen in our clinic, which correlates with the low incidence of RCC metastasis in this location as described earlier. Stage IV RCC has a 5-year survival rate of only 32\%, our patient after successful removal of the scalp metastasis, had a two-year follow up without lung or other metastases\textsuperscript{9}. Follow up after radical nephrectomy treatment of RCC, which includes thorough anamnesis, clinical examination, and diagnostic procedures like PET-CT and MRI scans to detect possible metastases, is the key for prompt and adequate treatment, as shown in the case of our patient\textsuperscript{19}.

**CONCLUSION**

In conclusion, RCC commonly metastasizes to the lung. This case report presents the rare case of a large fast-growing RCC metastasis to an unusual location, demonstrating its aggressiveness and high grade. In spite of that we showed two years follow up of remission. It should be noted that every lesion should be suspected as metastasis in RCC patients, confirmed via pathohistological analysis.

**Conflicts of interest statement:** the authors report no conflicts of interest.

**REFERENCES**


