ADRENAL METASTASES OF RENAL CELL CARCINOMA WITH INTRAVENOUS TUMOR THROMBI OF THE INFERIOR VENA CAVA
(two case reports with literature review)*

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In clinical studies the incidence of ipsilateral adrenal metastases in patients with renal cell carcinoma (RCC) ranges from 1.1% to 10%, and the rate of contralateral metastases usually does not exceed 1% [1–4]. Information about bilateral RCC metastases is limited to about two dozens of reports [5–9].

Intravenous extension is one of the attributes characterizing the neoplastic process. Microscopically this phenomenon is characteristic for many tumors, however massive macroscopic growth into the vena cava can be detected only with certain types of tumors though, specifically, renal cell carcinoma and various types of adrenal tumors [10–13]. So far in the worldwide literature only a few reports describe intravenous extension of distant RCC metastases, in particular, adrenal metastases [14–18].

Case study #1. A 60-year-old patient was hospitalized in October 2016 with complaints of right lower back pain and general weakness. Ultrasound and MDCT revealed a lesion in the lower pole of the right kidney of 60×68 mm. Right renal vein had no signs of tumor thrombosis. Retroperitoneal lymph nodes were not enlarged. In both lungs multiple soft tissue masses ranging from 5 to 12 mm were detected. In both right and left adrenal glands masses of 48 mm and 56 mm, respectively, were found. Assessment of the inferior vena cava
demonstrated the spread of the right adrenal mass into its lumen within 1.5 cm.

The patient’s general condition was satisfactory (ECOG performance status 0). Blood pressure monitoring revealed moderate arterial hypertension (130/90 mm Hg). The level of cortisol, metanephrine, normetanephrine, blood aldosterone, 17-OH corticosteroids and 17 oxysteroids of 24-hour urine had normal values. In this regard, the adrenal neoplasms were recognized as hormonally inactive. Hb = 122 g/l, erythrocytes = 3.8 × 10¹², leukocytes = 8.0 × 10⁹ g/l. Blood urea nitrogen = 6.8 mmol/l, creatinine = 98 µmol/l, glucose = 5.6 mmol/l. Taking into account the data obtained from the assessment, the diagnosis was the right kidney tumor T3aN0M1 with synchronous metastases into both adrenals and the right adrenal gland tumor extension into the inferior vena cava. After discussion with the patient the treatment plan was agreed: cytoreductive nephrectomy with resection of the left adrenal gland and right adrenalectomy or bilateral adrenalectomy.

The patient underwent Chevron incision laparotomy. The duodenum was mobilized using Kocher maneuver. Isolation of the inferior vena cava and both renal veins revealed the right adrenal gland tumor penetrating the subhepatic segment of the IVC with the left adrenal gland tumor expanding through the supra-renal vein up to its fusion with the left renal vein. The right renal artery was ligated and transected in the interaortocaval segment. After dissection and ligation of the right renal vein and the ureter, the kidney was removed.

Both adrenals were isolated from the surrounding tissue. In addition, it was discovered that the adrenal tissue was completely replaced by the tumors. Due to infeasibility of resection of neither one of the adrenal glands it was decided to perform bilateral adrenalectomy. Vascular clamps were placed on the IVC and on the left adrenal vein distal to the tumor masses. Lateral resection of IVC was performed in the area of the right adrenal vein mouth. The left adrenal vein was dissected away from the left renal vein. Both adrenals were removed along with the tumor intravenous thrombi. The vein defects were closed with Prolene suture 5-0.

The postoperative period was uneventful. Hormone replacement therapy with glucocorticoids and mineralocorticoids was used at the same time. The patient was discharged on the 9th day after surgery. Hormone replacement therapy is ongoing under the care of an endocrinologist. 1 month after the surgery the first line of targeted therapy was initiated (pazopanib 800 mg once daily). The 2- and 6-month MDCT demonstrated no signs of tumor progression. Most lung tumor lesions decreased in size or remained without change. The patient had no complaints. Blood pressure was
110/70 mmHg. Addison disease symptoms were absent.

**Case study #2.** A 53-year-old patient was admitted to hospital in February 2013, with complaints of hematuria and weight loss (4 kg within 6 months). MDCT examination revealed the right kidney middle segment tumor of 10×12 cm without signs of local invasion, extension to the regional lymph nodes and distant metastasis.
metastases. Considering these findings, radical nephrectomy with paracaval lymphadenectomy was performed. Right adrenalectomy was not performed, since computed tomography had revealed no signs of the right adrenal gland lesions. The final histopathological examination revealed poorly differentiated (Fuhrman Grade 3) clear cell renal cell carcinoma with signs of perinephric fat invasion (T3aN0M0). The postoperative period was uneventful. The patient was discharged from hospital 9 days later in satisfactory condition. However, after 6 months the follow-up MDCT revealed a tumor of the right adrenal gland of 2.5×3.0 cm with extension to the lumen of the inferior vena cava via the adrenal vein. The intravenous part of the tumor measured 2.2×1.2 cm. No distant metastases to the lungs or other organs were found.

The patient’s ECOG performance status was 0. Blood pressure monitoring revealed arterial hypertension (140/90 mm Hg). The level of cortisol, metanephrine, normetanephrine, blood aldosterone, 17-OH corticosteroids and 17-OH corticosteroids of 24-hour urine had normal values. Hb = 116 g/l, erythrocytes = 3.6×10¹², leukocytes = 7.0×10⁹ g/l. Blood urea nitrogen = 8.8 mmol/l, creatinine = 112 µmol/l, glucose = 4.6 mmol/l. Following the assessment, the diagnosis was established: metachronous renal cell carcinoma metastasis into the ipsilateral adrenal gland with intravenous extension to the subhepatic segment of the inferior vena cava. In October 2013 the patient underwent right adrenalectomy with thrombectomy and lateral resection of the inferior vena cava with the use of thoracolaparotomic approach to the 10th intercostal space. For vascular isolation of the tumor thrombus vascular clamps were placed on the IVC above and below the thrombus. After opening the vena cava it was found that intraluminal tumor masses were attached to its wall in the site of the confluence of the right adrenal vein. Furthermore, 10 mm below the tumor thrombus on the surface of the caval endothelium another tumor was found - metastasis to the IVC wall measuring about 5 mm. In this regard, lateral resection of the IVC was performed 3.0 cm. The resection margins were located within 5 mm of intraluminal tumor lesions.

The postoperative period was uneventful. Patient was discharged from hospital on the 10th day after surgery. After 6 months multiple lung metastases were found. The therapy with pazopanib (800 mg once daily) was initiated. 3 months later tumor progression (lung metastases increase in number and size) was identified. The patient died in 5 months (the total of 14 months after surgery) from tumor intoxication.

DISCUSSION

Intravenous extension of renal cell carcinoma adrenal metastases can be viewed in several ways. It is possible, that this phenomenon represents a specific feature of this neoplasm. Though, this assumption must be supported by similar incidence of RCC metastases invasion into the lumen of the major venous vessels. On the other hand, primary adrenal gland tumors by themselves have a distinct tendency to intravenous extension. Specifically, the adrenal pheochromocytomas are accompanied by this phenomenon in 2% of cases, and adrenocortical carcinomas in 9%–30% [19–21]. A large retrospective study in an Australian hospital presented 30 years’ experience of treatment of 464 patients with metastases into the adrenal glands. The authors have demonstrated that the rate of detection of adrenal metastases on autopsy is 3.1%, while at adrenalectomy is 7.5% [22].

In another work, which included 52 patient with various tumor metastases into the adrenal glands, localization of primary tumors was distributed as follows: the kidneys (28.8%), the lungs (21.2%), the intestine (13.4%), unknown localization (9.6%), the stomach (5.8%), melanoma (5.8%), others (15.4%) [23]. High proportion of adrenal metastases, originating from RCC, is also confirmed by clinical trials data, which studied the rate of occurrence of adrenal metastases of this tumor type. The studies have demonstrated that adrenal metastases are detected in 1%–10% of patients with renal cell carcinoma [1–4].

A distinct tendency of RCC towards macroscopic intravenous extension is widely known and well documented by many authors (20%–26%) [24, 25]. Though, can we declare the affinity to the venous route of
tumor extension for this type of cancer metastases? From our point of view, this is not a general tendency, since there are only a few reports on this phenomenon in the literature. In particular, a group of British authors presented a case study of metastatic renal cancer extending from the thoracic spine via v. azygos into the superior vena cava. In addition, there were significant clinical signs of the superior vena cava obstruction [26].

Previously we described a patient with multiple RCC metastases into the liver, one of which penetrated into the lumen of a portal vein branche [27]. This feature was discovered by ultrasonography and was histologically verified during autopsy. It is noteworthy, that the patient had been operated on 4 months before for renal cancer with tumor thrombus at the level of the subhepatic IVC segment. Along with liver metastasis, invading the portal vein, the patient also had a recurrent neoplastic thrombus, reaching retrohepatic segment of the inferior vena cava. A very interesting case was presented by Yachida S, et al., who discovered invasion of pancreatic RCC metastasis into Wirsung's duct in the form of tumor «thrombus» in three patients [28]. There are two more similar cases in the literature. At the same time it is believed that extension into the main pancreatic duct is a distinguishing feature of renal cancer metastases to the pancreas, since it is not typical for primary pancreatic tumors. There is an undoubtful similarity between intravenous and intraductal invasion, although the small number of observations does not allow us to claim its accuracy.

No less unique are the reports on extension of hepatic renal cell carcinoma metastases into either the hepatic duct lumen or the common bile duct lumen [29]. In addition, the tumor inside the duct also resembles a thrombus. In the world literature only three cases of this phenomenon are described. However, the invasion into the lumen of the biliary tract is not unique to renal cancer metastases, since similar cases of colon cancer and malignant melanoma metastases are described [30].

How common is intravenous extension of adrenal metastases? In the world literature we found only 7 cases, including two clinical cases of our own. Table 1 presents a summary on the subject.

Among all 7 patients ipsilateral metastases were found in 2 cases, contralateral in 3 cases, and bilateral in 2. Synchronous adrenal metastases were revealed only in 3 patients, in the rest of cases the metastases occurred at various intervals after surgery. One of our patients had extremely rare synchronous RCC metastases into both adrenal glands with synchronous extension into the venous system. Interesting, that renal tumor had no signs of intravenous invasion. Previously no such reports were presented in the literature.

How often do other cancer adrenal metastases propagate into the venous system? The literature analysis demonstrated 27 cases of this phenomenon with various types of tumors (Table 2). More than half of these clinical cases (55.6%, 15 patients) referred to he-

Table 1

<table>
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<tr>
<th>№</th>
<th>Authors</th>
<th>Lesion side</th>
<th>Synchronicity</th>
</tr>
</thead>
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<tr>
<td>1</td>
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<td>bilateral metastases with one thrombus</td>
<td>synchronous</td>
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<tr>
<td>3</td>
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<td>ipsilateral</td>
<td>synchronous</td>
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<tr>
<td>4</td>
<td>Ploumidis A, 2015 [18]</td>
<td>contralateral</td>
<td>metachronous</td>
</tr>
<tr>
<td>5</td>
<td>Piotrowicz, 2015 [15]</td>
<td>contralateral</td>
<td>metachronous</td>
</tr>
<tr>
<td>6</td>
<td>Authors’ case report # 1</td>
<td>bilateral metastases and bilateral thrombi</td>
<td>synchronous</td>
</tr>
<tr>
<td>7</td>
<td>Authors’ case report # 2</td>
<td>ipsilateral</td>
<td>metachronous</td>
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</table>
patocellular carcinoma, which were described mostly by Japanese authors.

The data presented indicates that extension of adrenal RCC metastases into the lumen of the large venous vessels is not rare and is the second most common cancer after hepatocellular carcinoma. However, due to the small total number of observations we cannot consider this phenomenon as manifestation of affinity of this cancer type to intravenous extension.

Synchronous bilateral RCC metastases occur quite rarely. Thus, von Knobloch R. et al. found adrenal metastases during surgical treatment of 617 patients with RCC in 23 (3.7%) cases. At the same time, bilateral metastases were found in 6 (1%) cases [44]. In the study by Antonelli A. et al. analyzing the results of treatment of 1179 patients the incidence of bilateral adrenal lesions did not exceed 0.2% [2]. Nevertheless, the literature does not present a lot of studies on the rate of occurrence and clinical features of bilateral adrenal RCC metastases. Basically, these are reports on a few clinical cases [5–9].

Surgical management of renal cell carcinoma with metastases into both adrenal glands is a complex task, since in most cases it requires bilateral adrenalectomy and subsequent hormone replacement therapy with glucocorticoids and mineralocorticoids. In one of our patients with bilateral adrenal metastases this problem was considerably more difficult, since the right-sided tumor synchronously extended into the inferior vena cava, and the left-sided one invaded the left adrenal vein. We decided on bilateral adrenalectomy with one stage thrombectomies, since the general condition of the patient was stable and nephrectomy passed without complications. It should be noted that resection of tumor thrombi with adrenal tumors has some peculiarities.

First of all, with the right-sided metastases the tumor easily extends via the adrenal vein to the retrohepatic segment and «higher» segments of the inferior vena cava. In most cases either mobilization of the liver with control of supradiaphragmatic part of the inferior vena cava, or the use of cardiopulmonary bypass is required. Secondly, with the left-sided metastases the tumor thrombus can invade not only the left adrenal vein, but the left renal vein also. Therefore, in order to preserve the left kidney,

Table 2
Case reports of intravenous extension of adrenal metastases of various cancer types

<table>
<thead>
<tr>
<th>№</th>
<th>Authors</th>
<th>Number of patients</th>
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</tr>
</thead>
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<tr>
<td>1</td>
<td>Takada K, 1989 [31]</td>
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</tr>
<tr>
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<td>Sakamoto Y, 1999 [32]</td>
<td>3</td>
<td>Hepatocellular carcinoma</td>
</tr>
<tr>
<td>3</td>
<td>Morimoto T, 1999 [33]</td>
<td>1</td>
<td>Hepatocellular carcinoma</td>
</tr>
<tr>
<td>4</td>
<td>Shimahara Y, 1999 [34]</td>
<td>1</td>
<td>Hepatocellular carcinoma</td>
</tr>
<tr>
<td>5</td>
<td>Momoi H, 2002 [35]</td>
<td>7</td>
<td>Hepatocellular carcinoma</td>
</tr>
<tr>
<td>7</td>
<td>Paparel P, 2004 [36]</td>
<td>1</td>
<td>Thyroid cancer</td>
</tr>
<tr>
<td>8</td>
<td>Ben Abid S, 2005 [37]</td>
<td>1</td>
<td>Hepatocellular carcinoma</td>
</tr>
<tr>
<td>9</td>
<td>Fernández Sarabia MT, 2008 [38]</td>
<td>1</td>
<td>Breast cancer</td>
</tr>
<tr>
<td>10</td>
<td>Wakayama K, 2013 [39]</td>
<td>1</td>
<td>Hepatocellular carcinoma</td>
</tr>
<tr>
<td>11</td>
<td>Tokue H, 2014 [40]</td>
<td>1</td>
<td>Ovarian cancer</td>
</tr>
<tr>
<td>12</td>
<td>Yang ST, 2014 [41]</td>
<td>1</td>
<td>Uterine leiomyosarcoma</td>
</tr>
<tr>
<td>13</td>
<td>Filippi L, 2014 [42]</td>
<td>1</td>
<td>Lung cancer</td>
</tr>
<tr>
<td>14</td>
<td>Vishnevskii AV, 2015 [43]</td>
<td>1</td>
<td>Colorectal cancer</td>
</tr>
</tbody>
</table>
resection and difficult vascular reconstruction of the renal vein may be necessary.

It is believed that in order to maintain adequate adrenal function 10% of adrenal tissue is required [7, 45]. Therefore, in recent years for adrenal tumors the organ sparing surgeries have been actively performed [46]. It should be noted, that these surgeries may not be always feasible though.

One should consider the size of the lesion, its localization, the volume of remnant adrenal tissue, as well as the degree of integrity of the adrenal vessels after resection. In our clinical case it was impossible to perform neither right-sided, nor left-sided organ sparing surgery, since the tumors had replaced the larger portion of the adrenal glands and extended into the main adrenal veins.

Although modern replacement therapy with glucocorticoids and mineralocorticoids is effective and safe, the patients with Addison disease due to bilateral adrenalectomy can have signs of acute adrenal insufficiency in various stress situations. Besides, in most cases, these patients suffer from side effects of steroid therapy, such as gastritis, hypertension and hypokalemia [5, 46].

Another important issue is feasibility of targeted therapy in presence of iatrogenic Addison disease. In our patient treatment with pazopanib was prescribed due to presence of pulmonary metastases after nephrectomy with bilateral adrenalectomy. At the same time, we found no significant side effects, and hormone replacement therapy did not require any adjustment.

CONCLUSION

We have presented two case reports on intravenous extension of renal cell carcinoma adrenal metastases. One of them for the first time in worldwide literature presents intravenous extension of synchronous bilateral adrenal metastases from RCC. Although these cases are extremely rare, it is necessary to take into account possibility of presence of venous tumor thrombi with adrenal metastases. The incidence of this phenomenon of RCC is in the second place after hepatocellular carcinoma. Organ-sparing surgical technique in these patients is not feasible. Targeted therapy along with hormone replacement therapy was not accompanied by increased side effects or exacerbation of adrenal insufficiency after removal of both adrenal glands.

REFERENCES


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Intravenous extension is one of the features of neoplastic process and is very characteristic for tumors of the kidneys and adrenal glands. However, reports about the penetration of adrenal metastases into the venous system are extremely rare. We have presented two case reports on intravenous extension of renal cell carcinoma adrenal metastases. One of them for the first time in worldwide literature presents intravenous extension of synchronous bilateral adrenal metastases RCC into inferior vena cava and left renal vein. The patient underwent bilateral adrenalectomy with the use of hormone replacement therapy and targeted therapy with pazopanib. MDCT after 2 and 6 months did not show tumor progression. Lungs metastases are reduced in size or remained unchanged. The patient does not complain. Blood pressure 110/70 mm Hg. Symptoms of Addison's disease are absent.

In the second case, there was a metachronic contralateral metastasis of RCC, which penetrated through the right adrenal vein into the subhepatic section of the inferior vena cava. The patient underwent a right-sided adrenalectomy with thrombectomy. After 6 months multiple pulmonary metastases were identified. Pazopanib therapy has initiated. Three months later, tumor progression was revealed. The patient died after 5 months (14 months after surgery) from tumor intoxication.

Although these cases are extremely rare (7 cases in the world literature), it is necessary to take into account possibility of presence of venous tumor thrombi with adrenal metastases. The incidence of this phenomenon of RCC is in the second place after hepatocellular carcinoma (15 cases). Organ-sparing surgical technique in these patients is not feasible. Targeted therapy along with hormone replacement therapy was not accompanied by increased side effects or exacerbation of adrenal insufficiency after removal of both adrenal glands.

Key words: renal cell carcinoma, adrenal metastases, intravenous invasion, tumor thrombus.
Хотя данные наблюдения являются экстремально редкими (7 случаев в мировой литературе), необходимо учитывать возможность существования опухолевых венозных тромбов при метастазах почечно-клеточного рака в надпочечники. По частоте встречаемости данного феномена ПКР занимает второе место после гепатоцеллюлярного рака (15 случаев). Органосохраняющая хирургическая тактика у данных пациентов не является возможной. Таргетная терапия на фоне заместительной гормональной терапии не сопровождалась увеличением числа побочных эффектов или усилением адrenalной недостаточности у пациента после удаления обоих надпочечников.

Ключевые слова: почечно-клеточный рак, адренальные метастазы, инравенозное распространение, опухолевый тромб.

Внутривенозное поширнення є однією з особливостей неопластического процесу і дуже характерно для пухлин нирок і надниркових залоз. Однак повідомлення про проростання у венозну систему адренальних метастазів є дуже рідкісними. Ми представили два клінічних спостереження внутрішньовенозного поширення адренальних метастазів нирково-клітинного раку. Одне з них вперше в світовій літературі описує синхронну внутрішньовенозну інвазію білатеральних адренальних метастазів нирково-клітинного раку в нижню порожністю вену ліворуч і в основну ліву ниркову вену праворуч. Пациєнту була виконана білатеральна адреналектомія з подальшою замісною гормональною і таргетною терапією пазопанібом. МДКТ через 2 і 6 місяців не продемонструвала прогресії пухлини. Більшість об’ємних утворень зменшилася в розмірах або залишалося без змін. Пациєнт не пред’являє скарг. Артеріальний тиск 110/70 мм рт. ст. Симптоми хвороби Аддісона відсутні.

У другому випадку мав місце метахронний контралатеральний метастаз НКР, який через праву адренальну вену проник до підпечінкового відділу нижньої порожністої вени. Пациєнції була виконана правооблична адреналектомія з тромбектомією. Через 6 місяців у неї виявлені множинні легеневі метастази. Розпочато терапію пазопанібом. Три місяці по тому була ідентифікована прогресія пухлини. Пацієнка померла через 5 місяців (через 14 місяців після операції) від пухлинної інтоксикації.

Хоча ці клінічні випадки є екстремально рідкісними (7 випадків), необхідно враховувати можливість існування пухлинних венозних тромбів при метастазах нирково-клітинного раку в наднирники. За частотою зустрічаємості даного феномена нирково-клітинний рак посідає друге місце після гепатоцеллюлярного раку (15 випадків). Органосберігаюча хірургічна тактика у даних пацієнтів не можлива. Таргетна терапія на тлі замісної гормональної терапії не супроводжувалася збільшенням кількості побічних ефектів або посиленням адrenalної недостатності у пацієнта після видалення обох наднирників.

Ключові слова: нирково-клітинний рак, адренальні метастази, інравенозне поширення, пухлинний тромб.