GUIDELINES ON RENAL CELL CANCER

J. Carballido, S. Hellsten, H. Mensink
H. Schulze, G. Mickisch
1. BACKGROUND

Renal cell carcinoma (RCC) accounts for about 2% of all cancers, with an worldwide annual increase of 1.5-5.9% (1,2). The mean age at the time of diagnosis is about 70 years and there is a predominance of men over women in the range of 1.5-3.1. The mortality from RCC is increasing parallel to trends in incidence (2). Worldwide mortality is expected to increase from 54,000 deaths in 1985 to 102,000 deaths in 2000. It may reach or even exceed that of bladder cancer in certain areas.

The increased incidence of RCC is primarily due to enhanced detection of tumours by expanded use of imaging techniques, such as ultrasound and computed tomography (CT) (2). At present, 25-40% of clinically diagnosed RCC are found incidentally. A total of 25-30% of patients with RCC have overt metastases at initial presentation and, in addition, a substantial fraction of patients have subclinical metastases at that time explaining the hitherto unsatisfactory outcome of treatment (3,4).

A slight to moderate improvement in survival has been observed in most countries. Survival is closely related to initial stage; 5-year survival is 50-90% for localised disease, decreasing to 0-13% for metastatic disease (3).

2. CLASSIFICATION

RCC represents the greater part of malignant tumours of the kidney (80-90%). The remainder include transitional cell carcinomas, non-epithelial kidney tumours and Wilms’ tumours (5,6). The TNM 97 classification is recommended and differs from TNM 92 in stage T1 (tumour size < 7 cm) and stage T2 (> 7 cm). It also differs in N1 (one node) and N2 (more than one node) involvement, while the N3 subcategory has been removed.

Robson’s classification (1969) is commonly used and the relationship with TNM 97 is as follows:
- Robson’s Stage I = T1-2
- Robson’s Stage II = T3a
- Robson’s Stage IIIa = T3b-c
- Robson’s Stage IVa = T4
- Robson’s Stage IIIb = N1-2
- Robson’s Stage IVb = M1 (6-8).

Traditionally RCC have been classified according to the nuclear (7,9) or cellular morphology (10). New morphologic, cytogenetic and molecular studies make it possible to distinguish five types of carcinomas:
- Clear - cell: 60-85%
- Chromophilic: 7-14%
- Chromophobic 4-10%
- Oncocytic: 2-5%
- Collecting duct: 1-2% (10).

Recent attempts have been made to generate a molecular classification (11).

There are no generally accepted risk factors for RCC. There are some epidemiologic data indicating that a smoking habit, obesity or exposure to certain heavy metals such as cadmium may favour the development of RCCs.

3. DIAGNOSIS

Clinical symptoms of RCC, such as haematuria, palpable tumour and flank pain, are becoming less frequent. Asymptomatic tumours are more commonly diagnosed (12). Clinical examination has a limited role in diagnosing RCC, but it may be valuable in assessing co-morbidity (12). In case of haematuria, additional tumours of the genitourinary tract should be excluded (13). The most commonly assessed laboratory parameters are:
- Haemoglobin and erythrocyte sedimentation rate: prognosis
- Creatinine: overall kidney function
- Alkaline phosphatase: liver metastasis, bone metastasis
Serum calcium is frequently included in the preoperative assessment because of its association with paraneoplastic manifestation, which may have clinical implications (14).

The majority of tumours are diagnosed by abdominal ultrasound performed for various reasons. Standard radiological procedure is an abdominal CT scan with and without contrast medium. It serves to document the diagnosis of RCC and provides information on the function and morphology of the contralateral kidney (15). Additional diagnostic procedures, such as magnetic resonance imaging, angiography or fine needle biopsy, have a very limited role, but may be considered in selected cases (16).

EXTENSION ASSESSMENT
Abdominal CT scan assesses primary tumour extension and provides information on venous involvement and on metastatic spread to loco-regional lymph nodes, adrenals, contralateral kidney, liver, etc (15). Chest X-ray is performed to assess pulmonary spread. If indicated by signs and symptoms, other diagnostic procedures may be applied, such as bone scan, brain CT, chest CT (12).

REFERENCES
4. TREATMENT

Only radical surgery offers a reasonable change of curing the disease (1). The chances of cure by surgery most strongly depend on stage (primarily) and grade (secondarily) of the disease (e.g. following TNM classification) (2). Standard operative procedure is a radical nephrectomy including Gerota’s fascia (3). There is no evidence to favour a specific surgical approach. In selected cases of small (< 4 cm) peripheral lesions, an organ sparing approach may be considered. Final evaluation of oncologic efficacy is pending (4, 5).

Adrenalectomy is generally recommended. The sparing of the ipsilateral adrenal gland in the case of a smaller tumour of the lower half of the kidney is currently being evaluated in ongoing clinical research (6). A formal lymph node dissection is a valuable diagnostic tool (staging); however, therapeutic efficacy is unproven (7).

If surgery cannot eradicate all tumour deposits, tumour nephrectomy remains palliative therapy and should be considered in the context of multimodality treatment (e.g. in conjunction with immunotherapy or experimental therapies) (8,9). Certain cases, such as bilateral tumours, a solitary tumour-bearing kidney, multifocal lesions, renal insufficiency, or an occasional palliative situation, will require individual decisions not amenable to general guidelines.

5. FOLLOW-UP

Rational for follow-up
Follow-up of patients with RCC after surgical treatment is recommended to detect local recurrence and distant metastases as early as possible to permit additional treatment when indicated and if possible. Such therapy may include resection of pulmonary metastasis or local recurrences; certain cases may also be candidates for immunomodulating therapy. With this background in mind, a regular postoperative follow-up of patients with RCC is proposed (10-12).

Principles
Prognostic factors and the type of surgical intervention (radical vs. partial or nephron sparing surgery) are relevant in determining the most efficient follow-up regimen. The only established prognostic factor is tumour stage according to the TNM system (12). After nephron sparing tumour resection (elective or mandatory indication), the local recurrence rate may vary between 0 and 10% (4,11). In a small proportion of patients with a genetic predisposition, a different follow-up procedure may be required (13,14).

Follow-up procedures
The first assessment is at 4-6 weeks and includes:
• Physical examination to exclude surgical complications
• Serum creatinine to assess the remaining kidney function
• Haemoglobin to assess recovery of perioperative blood loss.

If these values are normal, repeat investigation is usually unnecessary. Urine analysis is not needed for routine follow-up.

If alkaline phosphatase is abnormal preoperatively, repeat measurement is recommended because recurrent or persistent alkaline phosphatase elevation after surgery suggests distant metastasis, or residual tumour (15,16). Alkaline phosphatase elevation together with bone pain is suspicious for bone metastasis. Elevation may also occur in case of liver metastasis or paraneoplastic manifestations.

A chest X-ray is recommended to detect pulmonary metastases, which occur most commonly within 3 years after surgery. Imaging of the contralateral kidney is advocated in case of enhanced risk of developing metachronous occurrence (as in familial papillary RCC or VHL (von Hippel-Lindau disease). Imaging of the retroperitoneum by abdominal CT or ultrasound is recommended only after nephron sparing surgery or after radical surgery in locally advanced disease, e.g. T3, T4.

A recommended follow-up scheme is shown in Table 1.
REFERENCES


## Table 1: Recommended follow-up scheme for renal cell carcinoma

<table>
<thead>
<tr>
<th>Stage</th>
<th>Visit</th>
<th>Examination</th>
<th>Optional</th>
<th>Purpose</th>
</tr>
</thead>
<tbody>
<tr>
<td>All T</td>
<td>4-6 weeks after surgery</td>
<td>Physical exam, Creatinine, Hb</td>
<td>AP&lt;sup&gt;1&lt;/sup&gt;</td>
<td>Exclude complications of surgery, Establish remaining kidney function&lt;sup&gt;2&lt;/sup&gt;, To check recovery of perioperative blood loss</td>
</tr>
<tr>
<td>T1,2</td>
<td>Every 6 months for 3 years, Every year from 3-5 years&lt;sup&gt;3&lt;/sup&gt;</td>
<td>Physical exam, Chest X-ray</td>
<td>AP&lt;sup&gt;2&lt;/sup&gt;, Kidney imaging</td>
<td>Exclude complications of surgery and LR and LN metastases, Exclude pulmonary metastases and LR after partial nephrectomy</td>
</tr>
<tr>
<td>T3,4</td>
<td>Every 6 months for 3 years, Every year from 3-10 years</td>
<td>Physical exam, Chest X-ray, Retroperitoneal imaging</td>
<td></td>
<td>Exclude complications of surgery and LR and LN metastases, Exclude pulmonary metastases and LR after partial nephrectomy, To detect LR, contralateral metastases or neo-occurrence</td>
</tr>
</tbody>
</table>

AP = alkaline phosphatase; LR = local recurrence; LN = lymph node.

1. If elevated preoperatively (recurrent or persistent elevation suggests distant metastases or residual tumour) when bone pain is present, suspicious of bone or liver metastasis.
2. If the postoperative level is abnormal, it should be repeated at the regular visits.
3. There is a small, but continuous, risk of recurrence or metastasis from 5-15 years.