EAU GUIDELINES ON RENAL CELL CARCINOMA

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B. Ljungberg (Chair), L. Albiges, K. Bensalah, A. Bex (Vice-chair), R.H. Giles (Patient Advocate), M. Hora, M.A. Kuczyk, T. Lam, L. Marconi, A.S. Merseburger, T. Powles, M. Staehler, A. Volpe Guidelines Associates: Y. Abu-Ghanem, S. Dabestani. S. Fernández-Pello Montes, F. Hofmann, T. Kuusk, R. Tahbaz

Epidemiology

The use of imaging techniques such as ultrasound (US) and computed tomography (CT) has increased the detection of asymptomatic renal cell carcinoma (RCC). The peak incidence of RCC occurs between 60 and 70 years of age, with a 3:2 ratio of men to women. Aetiological factors include lifestyle factors, such as smoking, obesity and hypertension. Having a first-degree relative with RCC is associated with a significantly increased risk of RCC.

Staging system

The current UICC 2017 TNM (Tumour Node Metastasis) classification is recommended for the staging of RCC (Table 1).

Table 1: The 2017 TNM staging classification system

| T-P | rimary Tumour | |
|--------------------------|---|--|
| TX | Primary tumour cannot be assessed | |
| T0 | No evidence of primary tumour | |
| T1 | Tumour ≤ 7 cm or less in greatest dimension, limited to the kidney | |
| | T1a Tumour ≤ 4 cm or less | |
| | T1b Tumour > 4 cm but ≤ 7 cm | |
| T2 | Tumour > 7 cm in greatest dimension, limited to the kidney | |
| | T2a Tumour > 7 cm but ≤ 10 cm | |
| | T2b Tumours > 10 cm, limited to the kidney | |
| Т3 | Tumour extends into major veins or perinephric tissues but not into the ipsilateral adrenal gland and not beyond Gerota fascia | |
| | T3a Tumour grossly extends into the renal vein or its segmental (muscle-containing) branches, or tumour invades perirenal and/or renal sinus fat (peripelvic fat), but not beyond Gerota fascia | |
| | T3b Tumour grossly extends into the vena cava below diaphragm | |
| | T3c Tumour grossly extends into vena cava above the diaphragm or invades the wall of the vena cava | |
| T4 | Tumour invades beyond Gerota fascia (including contiguous extension into the ipsilateral adrenal gland) | |
| N - Regional Lymph Nodes | | |
| NX | Regional lymph nodes cannot be assessed | |
| N0 | No regional lymph node metastasis | |
| N1 | Metastasis in regional lymph node(s) | |

| M - Distant metastasis | | | | |
|------------------------|--------------------|-------|----|--|
| M0 Nod | istant metastas | sis | | |
| M1 Dista | ant metastasis | | | |
| TNM stage | TNM stage grouping | | | |
| Stage I | T1 | N0 | M0 | |
| Stage II | T2 | N0 | M0 | |
| Stage III | T3 | N0 | M0 | |
| | T1, T2, T3 | N1 | M0 | |
| Stage IV | T4 | Any N | M0 | |
| | Any T | Any N | M1 | |

A help desk for specific auestions about TNM classification is available at http://www.uicc.org/tnm.

Clinical Diagnosis

Many renal masses remain asymptomatic until late disease stages. The classic triad of flank pain, visible haematuria, and palpable abdominal mass is rare and correlates with aggressive histology and advanced disease.

Paraneoplastic syndromes are found in approximately 30% of patients with symptomatic RCCs. A few symptomatic patients present with symptoms caused by metastatic disease, such as bone pain or persistent cough.

Imaging

Computed tomography imaging, before and after intravenous contrast, can verify the diagnosis and provide information on the function and morphology of the contralateral kidney and assess tumour extension, including extra-renal spread, venous involvement, and enlargement of lymph nodes (LNs) and adrenals.

Abdominal US and magnetic resonance imaging (MRI) are supplements to CT. Contrast-enhanced US can be helpful in specific cases (e.g., chronic renal failure with a relative contraindication for iodinated or gadolinium-based contrast media, complex cystic masses, and differential diagnosis of peripheral vascular disorders such as infarction and cortical necrosis).

Magnetic resonance imaging can be used in patients with possible venous involvement, or allergy to intravenous contrast. Chest CT is the most accurate for chest staging and is recommended in the primary work-up of patients with suspected RCC.

In younger patients who are worried about the radiation exposure of frequent CT scans, MRI may be offered as alternative for follow-up imaging.

Biopsy

Percutaneous renal tumour biopsies are used:

- to obtain histology of radiologically indeterminate renal masses:
- to select patients with small renal masses for active surveillance;
- to obtain histology before, or simultaneously with, ablative treatments;
- to select the most suitable form of medical and surgical strategy in the setting of metastatic disease.

In patients with any sign of impaired renal function, a renal scan and total renal function evaluation using estimated glomerular filtration rate should always be undertaken to optimise the treatment decision.

Renal biopsy is not indicated for comorbid and frail patients who can be considered only for conservative management (watchful waiting) regardless of biopsy results.

| Recommendations | Strength rating |
|--|-----------------|
| Use multi-phasic contrast-enhanced computed tomography (CT) of abdomen and chest for the diagnosis and staging of renal tumours. | Strong |
| Use magnetic resonance imaging to better evaluate venous involvement, reduce radiation or avoid intravenous CT contrast medium. | Weak |
| Use non-ionising modalities, mainly contrast-enhanced ultrasound, for further characterisation of small renal masses, tumour thrombus and differentiation of unclear renal masses. | Strong |
| Do not routinely use bone scan and/or positron-emission tomography CT for staging of renal cell carcinoma. | Weak |
| Perform a renal tumour biopsy before ablative therapy and systemic therapy without previous pathology. | Strong |
| Perform a percutaneous biopsy in select patients who are considering active surveillance. | Weak |
| Use a coaxial technique when performing a renal tumour biopsy. | Strong |
| Do not perform a renal tumour biopsy of cystic renal masses. | Strong |
| Use a core biopsy technique rather than fine needle aspiration for histological characterisation of solid renal tumours. | Strong |

Histological diagnosis

A variety of renal tumours exist, and about 15% are benign. All kidney lesions require examination for malignant behaviour.

Histopathological classification

The new WHO/ISUP classification will replace the Fuhrman nuclear grade system but will need validation.

The three most common RCC subtypes, with genetic and histological differences, are: clear-cell RCC (cc-RCC) (80-90%), papillary RCC (10-15%), and chromophobe RCC (4-5%). The various RCC types have different clinical courses and responses to therapy.

Prognostic factors

In all RCC types, prognosis worsens with stage and histopathological grade. Histological factors include tumour grade, RCC subtype, sarcomatoid features, microvascular invasion, tumour necrosis, and invasion of the perirenal fat and collecting system. Clinical factors include performance status, local symptoms, cachexia, anaemia, platelet count, neutrophil/lymphocyte ratio, C-reactive protein and albumin.

| Recommendations | Strength rating |
|--|-----------------|
| Use the current Tumour, Node, Metastasis classification system. | Strong |
| Use grading systems and classify renal cell carcinoma type. | Strong |
| Use prognostic systems in the metastatic setting. | Strong |
| In localised disease, use integrated prognostic systems or nomograms to assess risk of recurrence. | Strong |

Disease Management Treatment of localised RCC

Localised RCCs are best managed with partial nephrectomy (PN) rather than radical nephrectomy (RN), irrespective of the surgical approach. Partial nephrectomy is unsuitable in some patients with localised RCC due to:

- locally advanced tumour growth:
- unfavourable tumour location:
- · significant health deterioration.

If pre-operative imaging and intra-operative findings are normal, routine adrenalectomy is not indicated.

Lymphadenectomy should be restricted to staging because the survival benefit of extended LN dissection is unclear in patients with localised disease. In patients who have RCCs with tumour thrombus and no metastatic spread, prognosis is improved after nephrectomy and complete thrombectomy.

Nephron-sparing surgery versus radical nephrectomy

Based on current available oncological and quality of life outcomes, localised RCC is best managed by nephron-sparing surgery (NSS) rather than RN, irrespective of the surgical approach. Before routine nephrectomy, tumour embolisation has no benefit. In patients unfit for surgery with massive haematuria or flank pain, embolisation can be a beneficial palliative approach.

| Recommendations | Strength rating |
|--|-----------------|
| Offer surgery to achieve cure in localised | Strong |
| renal cell cancer. | |
| Offer partial nephrectomy to patients with | Strong |
| T1 tumours. | |
| Do not perform ipsilateral adrenalectomy if | Strong |
| there is no clinical evidence of invasion of | |
| the adrenal gland. | |
| Offer an extended lymph node dissection | Weak |
| to patients with adverse clinical features, | |
| including a large diameter of the primary | |
| tumour. | |
| Offer embolisation to patients unfit for | Weak |
| surgery presenting with massive | |
| haematuria or flank pain. | |

Radical- and partial nephrectomy techniques

| Summary of evidence | LE |
|---|----|
| Laparoscopic RN has lower morbidity than open | 1b |
| nephrectomy. | |
| Short-term oncological outcomes for T1-T2a tumours are equivalent for laparoscopic- and open RN. | 2a |
| | 01 |
| Partial nephrectomy can be performed, either by open-, pure laparoscopic- or robot-assisted approach, based | 2b |
| on surgeon's expertise and skills. | |
| Partial nephrectomy is associated with a higher percentage of positive surgical margins compared to RN. | 3 |

| Recommendations | Strength rating |
|--|-----------------|
| Offer laparoscopic radical nephrectomy | Strong |
| (RN) to patients with T2 tumours and | |
| localised masses not treatable by partial | |
| nephrectomy (PN). | |
| Do not perform minimally invasive RN in | Strong |
| patients with T1 tumours for whom a PN is | |
| feasible by any approach, including open. | |
| Do not perform minimally invasive | Strong |
| surgery if this approach may compromise | |
| oncological-, functional- and peri-operative | |
| outcomes. | |

Alternatives to surgery Surveillance

Elderly and comorbid patients with incidental small renal masses have a low RCC-specific mortality and significant competing-cause mortality. In selected patients with advanced age and/or comorbidities, active surveillance (AS) is appropriate to initially monitor small renal masses, followed, if required, by treatment for progression. The concept of AS differs from the concept of watchful waiting. Watchful waiting is reserved for patients whose comorbidities contraindicate any subsequent active treatment and who do not require follow-up imaging, unless clinically indicated.

Cryoablation and radiofrequency ablation

Currently there are no data showing oncological benefit of cryoablation or radiofrequency ablation (RFA) techniques over PN.

| Recommendation | Strength rating |
|---|-----------------|
| Offer active surveillance, radiofrequency | Weak |
| ablation or cryoablation to frail and/or | |
| comorbid patients with small renal masses. | |
| When radiofrequency ablation, cryoablation | Weak |
| and active surveillance are offered, inform | |
| patients about the higher risk of local | |
| recurrence and/or tumour progression. | |

Treatment of locally advanced RCC Management of clinically positive lymph nodes (cN+)

In the presence of clinically positive LNs (cN+), LND is always iustified but the extent of LND is still controversial.

Low level data suggest that tumour thrombus in the setting of non-metastatic disease should be excised. Adjunctive procedures such as tumour embolisation or inferior vena cava filter do not appear to offer any benefits in the treatment of tumour thrombus.

In patients unfit for surgery, or with non-resectable disease, embolisation can control symptoms, including visible haematuria or flank pain. At present there is no evidence for the use of adjuvant therapy following surgery.

Treatment of advanced/metastatic RCC Management of RCC with venous tumour thrombus

| Recommendations | Strength rating |
|--|-----------------|
| In patients with clinically enlarged lymph | Weak |
| nodes (LNs), perform LN dissection for | |
| staging purposes or local control. | |
| Remove the renal tumour and thrombus | Strong |
| in case of venous involvement in non- | |
| metastatic disease. | |

Cytoreductive nephrectomy

Tumour nephrectomy is curative only if all tumour deposits are excised. This includes patients with the primary tumour in place and single- or oligometastatic resectable disease. For most patients with metastatic disease, cytoreductive nephrectomy (CN) is palliative and systemic treatments are necessary.

| Summary of evidence | LE |
|---|----|
| Deferred CN with pre-surgical sunitinib in intermediaterisk patients with cc-metastatic RCC (mRCC) shows a survival benefit in secondary endpoint analyses and selects out patients with inherent resistance to systemic therapy. | 2b |
| Sunitinib alone is non-inferior compared to immediate CN followed by sunitinib in patients with MSKCC intermediate and poor risk who require systemic therapy with vascular endothelial growth factor receptor (VEGFR)-tyrosine kinase inhibitor (TKI). | 1a |
| Cytoreductive nephrectomy for patients with simultaneous complete resection of a single metastasis or oligometastases may improve survival and delay systemic therapy. | 3 |
| Patients with MSKCC or IMDC poor risk (≥ 4 risk factors) do not benefit from local therapy. | 1a |

| Recommendations | Strength rating |
|---|-----------------|
| Do not perform cytoreductive nephrectomy | Strong |
| (CN) in MSKCC poor-risk patients. | |
| Do not perform immediate CN in MSKCC | Weak |
| intermediate-risk patients who have an | |
| asymptomatic synchronous primary | |
| tumour and require systemic therapy with | |
| vascular endothelial growth factor receptor | |
| (VEGFR)-tyrosine kinase inhibitor (TKI). | |
| Start systemic therapy without CN in | Weak |
| MSKCC intermediate-risk patients who | |
| have an asymptomatic synchronous | |
| primary tumour and require systemic | |
| therapy with VEGFR-TKI. | |
| Discuss delayed CN in MSKCC intermediate- | Weak |
| risk patients under VEGFR-TKI therapy who | |
| derive long-term sustained benefit and/or | |
| minimal residual metastatic burden. | |
| Perform immediate CN in patients with | Weak |
| good performance who do not require | |
| systemic therapy. | |
| Perform immediate CN in patients with | Weak |
| oligometastases when complete local | |
| treatment of the metastases can be | |
| achieved. | |

IMDC = International Metastatic RCC Database Consortium; MSKCC = Memorial Sloan-Kettering Cancer Center.

Local therapy of metastases in metastatic RCC

A systematic review of the local treatment of metastases from RCC in any organ was undertaken. The heterogeneity of the data will only allow for cautious recommendations.

| Summary of evidence | LE |
|--|----|
| All studies included in a Panel systematic review were | 3 |
| retrospective, non-randomised comparative studies, | |
| resulting in a high risk of bias associated with non- | |
| randomisation, attrition, and selective reporting. | |
| With the exception of brain and possibly bone | 3 |
| metastases, metastasectomy remains by default the | |
| only local treatment for most sites. | |
| Retrospective comparative studies consistently point | 3 |
| towards a benefit of complete metastasectomy in | |
| mRCC patients in terms of OS, CSS and delay of | |
| systemic therapy. | |
| Radiotherapy to bone and brain metastases from RCC | 3 |
| can induce significant relief from local symptoms | |
| (e.g. pain). | |

| Recommendations | Strength rating |
|--|-----------------|
| To control local symptoms, offer ablative | Weak |
| therapy, including metastasectomy, to | |
| patients with metastatic disease and | |
| favourable disease factors and in whom | |
| complete resection is achievable. | |
| Offer stereotactic radiotherapy for clinically | Weak |
| relevant bone- or brain metastases for local | |
| control and symptom relief. | |

Systemic therapy for advanced/metastatic RCC Chemotherapy

| Recommendation | Strength rating |
|--|-----------------|
| Do not offer chemotherapy to patients with | Strong |
| metastatic renal cell carcinoma. | |

Immunotherapy

Interferon- α monotherapy and combined with bevacizumab, has been superceded as standard treatment by targeted therapy of advanced cc-mRCC.

Immune checkpoint inhibition of programmed death receptor (PD-1) and ligand (PD-L1) inhibition have been investigated in mRCC. Randomised data support the use of nivolumab (a PD-1 inhibitor) in VEGF-refractory disease. The combination of two immune checkpoint inhibitors: ipilimumab and nivolumab showed superior survival in intermediate- and poor-risk patients while the combination of pembrolizumab and axitinib showed survival advantage for patients in all risk groups.

| Summary of evidence | LE |
|--|----|
| Interferon- α monotherapy is inferior to VEGF-targeted therapy or mammalian target of rapamycin (mTOR) inhibition in mRCC. | 1b |
| Nivolumab leads to superior OS compared to everolimus in patients failing one or two lines of VEGF-targeted therapy. | 1b |
| The combination of nivolumab and ipilimumab in treatment-naïve patients with cc-mRCC of IMDC intermediate and poor risk demonstrated OS and ORR benefits compared to sunitinib. | 1b |
| The combination of pembrolizumab and axitinib in treatment-naïve patients with cc-mRCC across all IMDC risk groups demonstrated OS and ORR benefits compared to sunitinib. | 1b |
| Currently, PD-L1 expression is not used for patient selection. | 2b |
| Axitinib can be continued if immune-related adverse events results in cessation of axitinib and pembrolizumab. Re-challeange with immunotherapy requires expert support. | 4 |
| Patients who do not receive the full 4 doses of ipilimumab due to toxicity should continue on single-agent nivolumab, where safe and feasible. Re-challange with combination therapy requires expert support. | 4 |
| Treatment past progression can be justified but requires close scrutiny and the support of an expert multidisciplinary team. | 1b |

| Nivolumab plus ipilimumab and pembrolizumab plus axitinib should be administered in centres with experience of immune combination therapy and appropriate supportive care within the context of a multidisciplinary team. | 4 |
|--|----|
| The combination of nivolumab and ipilimumab in the intention-to-treat population of treatment-naive unselected patients with cc-mRCC leads to superior survival compared to sunitinib. | 2b |
| Due to the exploratory nature of PD-L1 tumour expression, the small sample size, the lack of OS data and the premature results in this subpopulation, definitive conclusions cannot be drawn relative to the usefulness of PD-L1 expression. | 2b |
| Nivolumab plus ipilimumab was associated with 15% grade 3-5 toxicity and 1.5% treatment-related deaths. | 1b |

| Recommendations | Strength rating |
|---|-----------------|
| Offer pembrolizumab plus axitinib to | Strong |
| treatment-naïve patients with any | |
| IMDC-risk clear-cell metastatic renal cell | |
| carcinoma (cc-mRCC). | |
| Offer ipilimumab plus nivolumab to | Strong |
| treatment-naïve patients with IMDC | |
| intermediate- and poor-risk cc-mRCC. | |
| Administer nivolumab plus ipilimumab and | Weak |
| pembrolizumab plus axitinib in centres with | |
| experience of immune combination therapy | |
| and appropriate supportive care within the | |
| context of a multidisciplinary team. | |

| Patients who do not receive the full 4 doses of ipilimumab due to toxicity should continue on single-agent nivolumab, where safe and feasible. | Weak |
|--|---------|
| Offer axitinib as subsequent treatment to patients who experience treatment-limiting immune-related adverse events after treatment with the combination of axitinib and pembrolizumab. | Weak |
| Treatment past progression can be justified but requires close scrutiny and the support of an expert multidisciplinary team. | Weak |
| Do not re-challange patients who stopped immune checkpoint inhibitors because of toxicity without expert guidance and support from a multidisciplinary team. | Strong |
| Offer nivolumab after one or two lines of vascular endothelial growth factor-targeted therapy in mRCC. | Strong |
| Offer sunitinib or pazopanib to treatment- naïve patients with IMDC favourable-, intermediate-, and poor-risk cc-mRCC who cannot receive or tolerate immune checkpoint inhibition. | Strong |
| Offer cabozantinib to treatment-naïve patients with IMDC intermediate- and poor-risk cc-mRCC who cannot receive or tolerate immune checkpoint inhibition. | Strong* |

^{*} While this is based on a randomised phase II trial, cabozantinib (weak) looks at least as good as sunitinib in this population. This justified the same recommendation under exceptional circumstances.

IMDC = International Metastatic RCC Database Consortium.

Targeted therapies

At present, several targeting drugs have been approved for the treatment of mRCC.

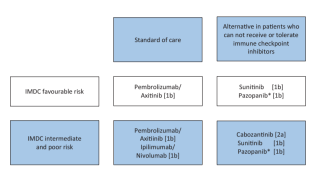
| Summary of evidence | LE |
|--|----|
| Single agent VEGF-targeted therapy has been superseded by immune checkpoint-based combination therapy. | |
| Pazopanib is non-inferior to sunitinib in front-line mRCC. | 1b |
| Cabozantinib in intermediate- and poor-risk treatment- naïve cc-RCC leads to better response rates and PFS but not OS when compared to sunitinib. | 2b |
| Tivozanib has been EMA approved, but the evidence is still considered inferior over existing choices in the front-line setting. | 3 |
| Single-agent VEGF-targeted therapies are preferentially recommended after front-line PD-L1-based combinations. Re-challenge with treatments already used should be avoided. | 3 |
| Single-agent cabozantinib or nivolumab are superior to everolimus after one or more lines of VEGF-targeted therapy. | 1b |
| Everolimus prolongs PFS after VEGF-targeted therapy when compared to placebo. This is no longer widely recommended before third-line therapy. | 1b |
| Both mTOR inhibitors and VEGF-targeted therapies have limited activity in non-cc-mRCC. There is a non-significant trend for improved oncological outcomes for sunitinib over everolimus. | 2a |

| Lenvatinib in combination with everolimus improved | 2a |
|---|----|
| PFS over everolimus alone in VEGF-refractory disease. | |
| Its role after immune checkpoint inhibitors is | |
| uncertain. There is a lack of robust data on this | |
| combination making its recommendation challenging. | |

| Recommendations | Strength rating |
|--|-----------------|
| Offer nivolumab or cabozantinib for | Strong |
| immune checkpoint inhibitor-naive | |
| vascular endothelial growth factor receptor | |
| (VEGFR)-refractory clear-cell metastatic | |
| renal cell carcinoma (cc-mRCC). | |
| Sequencing the agent not used as second- | Weak |
| line therapy (nivolumab or cabozantinib) for | |
| third-line therapy is recommended. | |
| Offer VEGF-tyrosine kinase inhibitors as | Weak |
| second-line therapy to patients refractory | |
| to nivolumab plus ipilimumab or axitinib | |
| plus pembrolizumab. | |
| Offer cabozantinib after VEGF-targeted | Strong |
| therapy in cc-mRCC. | |
| Sequence systemic therapy in treating | Strong |
| mRCC. | |

IMDC = International Metastatic RCC Database Consortium.

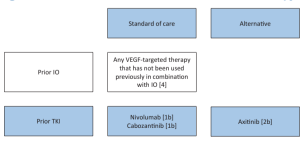
Figure 1: Updated EAU Guidelines recommendations for the treatment of first-line and following lines in clear-cell metastatic renal cancer.



IMDC = International Metastatic RCC Database Consortium.
*pazopanib for intermediate-risk disease only.
[1b] = based on one randomised controlled phase III trial.

[2a] = based on one randomised controlled phase II trial.

Figure 2: Guidelines Recommendations for later-line therapy



IO = immunotherapy; TKI = tyrosine kinase inhibitors; VEGF = vascular endothelial growth factor. [1b] = based on one randomised controlled phase III trial.

[2b] = subgroup analysis of a randomised controlled phase III trial

[4] = expert opinion.

Recurrent RCC

Locally recurrent disease can occur either after nephrectomy, PN, or after ablative therapy. After nephron-sparing treatment approaches the recurrence may be intra-renal or regional, e.g. venous tumour thrombi or retroperitoneal LN metastases. Isolated local recurrence in the true renal fossa after RN is rare.

Patients can benefit from a complete surgical resection of local recurrent disease. In cases where complete surgical removal is not feasible due to advanced tumour growth and pain, palliative treatments including radiation treatment can be considered

Surveillance following surgery for RCC

The aim of surveillance is to detect either local recurrence or metastatic disease while the patient is still surgically curable. Surveillance after treatment for RCC allows the urologist to assess:

- postoperative complications:
- renal function:
- local recurrence:
- recurrence in the contralateral kidney;
- · development of metastases.

Depending on the availability of new effective treatments, more intensive follow-up schedules may be required, particularly as there is a higher local recurrence rate after cryotherapy and RFA. At present there is no evidence-based standard for the follow-up of patients with RCC, or for the optimal duration of follow-up. An example of a surveillance

algorithm monitoring patients after treatment for RCC that recognises not only the patient's risk profile but also treatment efficacy is provided in Table 2. For patients with metastatic disease, individualised follow-up is indicated.

Table 2: Proposed surveillance schedule following treatment for RCC, taking into account patient risk profile andtreatment efficacy (expert opinion [LE: 4])

| Risk profile | Surveillance | | | | |
|------------------------|--------------|----|-----|-----|--|
| | 6 mo | 1y | 2 y | 3 y | >3y |
| Low | US | СТ | US | СТ | CT once every 2 years; counsel about recurrence risk of ~10% |
| Intermediate / High | СТ | СТ | СТ | СТ | CT once every 2 years |

CT = computed tomography of chest and abdomen, alternatively use magnetic resonance imaging for the abdomen; US = ultrasound of abdomen, kidneys and renal bed.

Summary of evidence and recommendations for surveillance following RN or PN orablative therapies in RCC

| Summary of evidence | LE |
|---|----|
| Surveillance can detect local recurrence or metastatic disease while the patient is still surgically curable. | 4 |
| After NSS, there is an increased risk of recurrence for larger (> 7 cm) tumours, or when there is a positive surgical margin. | 3 |
| Patients undergoing surveillance have a better OS than patients not undergoing surveillance. | 3 |
| Repeated CT scans do not reduce renal function in chronic kidney disease patients. | 3 |

| Recommendations | Strength rating |
|--|-----------------|
| Base follow-up after RCC on the risk of | Strong |
| recurrence. | |
| Intensify follow-up in patients after nephron- | Weak |
| sparing surgery for tumours > 7 cm or in | |
| patients with a positive surgical margin. | |
| Base risk stratification on pre-existing | Strong |
| classification systems such as the | |
| University of California Los Angeles | |
| integrated staging system or the SSIGN | |
| score. | |

SSIGN = (Mayo Clinic) stage, size, grade, and necrosis score.

This short booklet text is based on the more comprehensive EAU Guidelines (ISBN 978-94-92671-07-3), available to all members of the European Association of Urology at their website: http://www.uroweb.org/quidelines/.