Current management options in metastatic renal cell cancer

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Abstract

Renal cancer accounts for 2.4% of all cancers. Localised disease accounts for the majority of renal cancers (70-75%), however 20-25% of these ultimately develop distant metastasis. The median overall survival of untreated metastatic disease is 5 months with 1-year survival of only 29%. The management of metastatic renal cell cancers traditionally relied on systemic immunotherapy with attendant high morbidity but after the year 2005 the use of effective targeted therapy with tolerable side effect profile has improved the survival from 10.2 months in the cytokine era to 17.7 months. This article reviews the past, present and future options in the management of metastatic renal cancer.

Introduction

Renal cancer (RCC) currently ranks twelfth among other cancers in incidence, a position it holds along with pancreatic cancer. A lifetime risk of 1 in 63 (1.6%) has been reported for the general population with a median age of 65 years at diagnosis. GLOBOCAN 2012 data indicate renal cancer accounts for 2.4% of all cancers.1 The age-standardised rate worldwide is 4.4 per 100,000 population with males (6 per 100,000) more commonly afflicted than females (3 per 100,000).2 North America and Europe have a reported higher incidence compared to Africa and Asia with Czech Republic and Lithuania reporting an age standardised rate of 16.7 and 13.2 per 100,000.3 Time trends indicate an increasing incidence from 1975-2009, primarily due to detection of localised disease with incidence of distant disease remaining unchanged. The natural history of metastatic renal cancer

The 10 major histological types of renal cancer result in considerable variability in the natural history of the disease and this reflects in the five year survival rates which vary from 81% for stage I disease to only 8% for metastatic disease.5 Localised disease accounts for the majority of renal cancers (70-75%), however 20-25% of these ultimately develop distant metastasis. The median overall survival of untreated metastatic disease is 5 months with 1 yr survival of only 29%.5 RCC commonly metastasizes to lung (60%), bone (30%), liver, lymph nodes and the brain. The prognosis is known to vary with the involved target organ, liver and bone involvement resulting in adverse survival compared to lung and lymph nodal disease. This observation initially noted during the cytokine era has been reported to be unchanged with the targeted therapy regimes as well. Kroeger et al reported poor survival in patients with nodal involvement and recommended lymph nodal dissection during cytoreductive nephrectomy.7 The treatment of metastatic renal cell cancers traditionally relied on systemic immunotherapy with its attendant high morbidity but after 2006 the use of effective targeted therapy with tolerable side effect profile has improved the survival from 10.2 months in the cytokine era to 17.7 months with targeted therapy.

Molecular biology of renal cancer

Our knowledge of the abnormal molecular pathways involved in renal cancer genesis is attributed to study of hereditary renal cancers. Mutations in the Von Hippel Lindau (VHL) gene are the primary defect in 92% of sporadic and all hereditary clear cell cancers. The hereditary papillary, chromophobe and oncocytic tumors have defective MET and folliculin (FLCN) genes respectively but the sporadic variants of these types exhibit these abnormalities in only13% and 11% of tumours.8 The VHL gene product binds to hypoxia inducible factor (HIF-α) and targets it for ubiquitin mediated degradation. In clear cell cancers inactivating mutation of the VHL protein inhibit HIF-α degradation allowing accumulation of HIF-α function as a transcription factor and up regulates its target genes like VEGF, PDGF and GLUT-1 resulting in neovascularisation and tumor growth. The mTOR pathway functions in the downstream of PI3-K/Akt pathway and through its effector S6K regulates cell growth and proliferation in response to growth factors, nutrient, energy level and environmental stresses. Inactivating mutations of the FLCN gene negatively regulates the mTOR pathway. The MET proto-oncogene codes for hepatocyte growth factor (HGF) hence activating mutation as in papillary tumors promote cell growth, morphogenesis and differentiation.8
Targeting each of these pathways presents promising treatment strategies in advanced renal cancer. Targeting strategies include multistep inhibition of a single pathway (vertical) or single step inhibition of multiple pathways (horizontal).

### Risk stratification in renal cancer

The varied spectrum of biological behaviour of renal cancer coupled with a myriad of approved targeted therapy makes selection of appropriate therapy for an individual patient difficult. The use of prognostic or predictive factors can help circumvent this issue but each approach has its inherent limitations. Several prognostic models like the MSKCC risk model, the Cleveland clinic foundation (CCF) model, Groupe Francais d’Immunotherapie (French) model, the IKCWG model and the International Metastatic IMDC model have been reported. The Motzer (MSKCC) model uses low Karnofsky performance status (PS<80%), high serum lactate dehydrogenase (LDH>1.5 times upper limit of normal-ULN), low hemoglobin (Hb=lower limit of normal-LLN), high corrected serum calcium (>10 mg/dL), and time since diagnosis and treatment initiation <1 year to stratify patients with metastatic into good risk (0 risk factors), intermediate (1-2 risk factors) and poor risk groups. Mekhail et al. added prior radiotherapy and number of metastatic sites to motzers criteria and subsequently validated the same (CCF model). The French model uses presence of hepatic metastases, short interval from renal tumor to metastases (<1 year), more than one metastatic site and elevated neutrophil counts (ANC) as prognostic determinants and patients who combined at least three of these factors have >80% probability of rapid progression despite treatment. The Heng model (IMDC) excludes serum LDH from the motzer model but includes high neutrophil and platelet counts to stratify risk similar to the motzer model. The IKCWG model uses performance status, number of metastatic sites, time from diagnosis to treatment, and pre treatment hemoglobin, white blood count, lactate dehydrogenase, alkaline phosphatase, and serum calcium into a risk score that is subsequently used to stratify patients into good, intermediate and poor risk groups. It is important to note while the motzer model was developed for use in the cytokine era, the IKCWG and CCF models have been tested on patients treated with targeted therapy and have been found to be valid. The motzer and the Heng model have been used in the inclusion of study participants in the management defining clinical trials and consequently a working knowledge of risk stratification is essential for selection of appropriate therapy in practice. A tabulated summary of each model is presented in Table 1 for quick review.

| Table 1. Prognostic models in advanced RCC. |
|--------------------------------------------|
| **Prognostic factor** | **MSKCC Motzer** | **CCF** | **HENG** | **IKCWG** |
| Time from RCC diagnosis to start of therapy <1 yr | + | + | + | + |
| PS <80% Karnofsky | + | + | + | + |
| Sr LDH >1.5x ULN | + | - | - | + |
| Corrected Serum ca >ULN | + | + | + | + |
| Hb<LLN | + | - | + | + |
| Prior RT | - | + | - | - |
| >2 metastatic sites | - | + | - | + |
| ANC >ULN | - | - | + | + |
| Platelet >ULN | - | - | + | - |
| Prior immunotherapy | - | - | - | + |
| Elevated alkaline phosphatase | - | - | - | + |
| Risk category | **Good = 0** | **Favourable = 0 or 1** | **Favourable = 0** | **Favourable = 0** |
| **Intermediate = 1 or 2** | **Intermediate = 1 or 2** | **Intermediate = 1 or 2** | **Intermediate = 1 or 2** |
| **Poor = >2** | **Poor = >2** | **Poor = >2** | **Poor = >2** |

### Surgical management

Surgical options include cytoreductive nephrectomy, metastectomy and local therapies including radiotherapy. The benefit of cytoreductive nephrectomy in combination with cytokine therapy has been proved in several phase III trials. The southwest oncology group reported a 31% reduction in the risk of death in the nephrectomy plus interferon α 2b arm compared to interferon only arm (13.6 vs 7.8 months median survival). The European genito urinary group in a similar study design demonstrated a17 vs 7 months survival benefit to the interferon α 2b and nephrectomy arm. An identical benefit has been noted with interleukin-2 and cytoreductive nephrectomy. The ideal candidate for this combined therapy has been a source of considerable debate. It is generally accepted that a resectable tumor, performance status 0 or 1, lung only metastasis, good risk status as ascertained by the motzer criteria and absence of brain, bone and liver metastasis identify the cohort that derives the best results from this therapy. It is important to note despite a survival benefit, cytoreductive nephrectomy is associated with higher morbidity and mortality. Abdollah et al. after a database review of 1063 patients reported higher mortality (2.4% vs 0.9%), longer hospital stay (8.4 vs 5.7 days), higher second procedure (28% vs 10%), and post op complications (26% vs 19%) in the nephrectomy patients. Advanced age ≥ 75 years and comorbidity score of ≥3 was associated with poor outcomes, hence despite recommendations a prudent clinical decision after thorough discussion with the patient is required. It is noteworthy that the above benefit has been noted in clear cell histological type only. No phase III prospective trial data exists for combining nephrectomy with targeted agent, however several retrospective data indicate a benefit of 19.8 vs 9.4 months while combining cytoreductive nephrectomy with VEGF targeted therapy.

Approximately 1.5-3% of patients have solitary metastasis and
can be considered for metastatectomy. A survival advantage has been demonstrated in retrospective studies when such surgery has been combined with immunotherapy. Metastatectomy is done with a curative intent and is generally preceded or combined with radical nephrectomy. Metastatectomy has been reported for several organs including lung, bone and brain metastasis, however lung metastasis has been observed to have better overall survival compared to other sites. No clear selection criteria are recommended but good performance status, a completely resectable lesion preferably lung metastasis and absence of major comorbidities indicate an ideal surgical candidate.¹⁹

**Systemic therapy**

**Chemotherapy**

Systemic therapy for metastatic renal cancer was long considered ineffective as early trials with medroxy progesterone and chemotherapy with 5FU failed to demonstrate significant antitumor effects. A review of 51 phase II trials that included 33 chemotherapy drugs showed an overall response rate of only 5.5%.¹⁸ Recently combination chemotherapy with 5FU and gemcitabine has been reported to have a response rates of 10-15% in phase II trials.²¹ The addition of 5FU to immunotherapy has failed to improve survival in phase III trials.²²

**Cytokine therapy**

Interleukin-2 (IL-2) and interferon α-2b (IFN) represented the standard of care until the advent of targeted therapy in 2005. IL-2 has been in use to treat renal cancer since 1985 with response rate varying from 7-27%. High dose IL-2 in a dose of 600,000 IU/Kg for 14 doses or 720,000 IU/Kg for 12 doses every 8 hours per week produced complete response in 7% that was maintained for a median of 80 months. Substantial toxicity was the major limitation with 3-4% succumbing to therapy related mortality. In current practice high dose IL-2 though listed as first line therapy in patients with good performance status, lung metastasis and clear cell histology is not preferentially used in lieu of targeted therapy.¹⁹ IFN monotherapy produces responses similar to hormonal agents with response rate of 6-15%. A Cochrane meta analysis confirmed a moderate benefit compared to placebo but the response was not reproducible in the intermediate risk category.¹⁹ Combination of IFN 9MU three times a week with bevacizumab 10 mgs/kg biweekly is an approved first line therapy in metastatic clear cell renal cancer in patients with favourable risk assessment.

**Targeted therapy**

Pazopanib is an oral anti angiogenic tyrosine kinase inhibitor approved for first line therapy and in patients progressive after IFN therapy. It is also effective against PDGFR α, β and cKit. The VEG 105192 trial tested pazopanib 800 mgs/day against placebo in both treatment naïve and patients progressing on IFN, with 89% of study participants nephrectomised in both arms. A significant PFS survival advantage to pazopanib was demonstrated (9.2 vs 4.2 months, HR 0.46), the benefit was larger in treatment naïve (11.1 vs 2.8, HR 0.40) compared to those receiving prior cytokines (7.4 vs 4.2, HR 0.54). The trial design permitted crossover which may have confounded the non significant overall survival results (22.9 vs 20 months). Treatment discontinuation due to adverse effects was only 14% with diarrhoea (52%), nausea (26%) and haematological toxicity (32%) commonly reported. The commonest grade 3/4 toxicity was hypertension (40%). Toxicities were manageable by dose reduction to 400 mgs and subsequent increments done in 200 mgs steps. The COMPARZ, non-inferiority trial tested pazopanib 800 mgs/day against sunitinib 50 mgs/day in a 4 week on 2 week off cycle in treatment naïve patients. Pazopanib was found to be non-inferior as assessed with the primary end point of PFS (8.4 vs 9.5 months) and with overall survival (28.3 vs 29.1 months). Pazopanib was found to be less toxic with lower fatigue, hand foot syndrome, and thrombocytopenia, and had better patient acceptability with only 24% discontinuing due to adverse effects. Treatment related mortality was identical at 1% in both groups. The side effects were managed by dose decrements of 200 mg for pazopanib and 12.5 mgs for sunitinib.²⁴ The PISCES crossover study tested patient preference between pazopanib and sunitinib and confirmed better patient preference for pazopanib over sunitinib.²⁵

Sunitinib is a multi tyrosine kinase receptor inhibitor effective against VEGF,PDGF and cKit. It was one of the first targeted therapy (2006) to be approved for metastatic renal cell carcinoma in cytokine refractory patients and in those with no prior therapy. A multicenter international study compared standard dose sunitinib with IFN in treatment naïve metastatic RCC. Nephrectomy rates approached 90% and 77% had more than 2 metastatic sites. An interim analysis with PFS endpoint favoured the sunitinib arm (11 vs 5 months, HR 0.41) subsequently the trial permitted crossover not surprisingly an overall survival benefit remained insignificant (114 vs 95 weeks, HR 0.82). Adverse event were more in the sunitinib arm and grade 3/4 toxicities were identical in both arms. Side effects requiring dose reductions were more frequent in the sunitinib arm (32 vs 21%). Despite more general adverse effects with sunitinib quality of life parameters favoured sunitinib (P<0.001).²⁶ The EFFECT trial tested sunitinib in standard dosing schedule with a modified dose of 37.5 mgs/day continuous regime. The time to progression was longer with the standard dosing schedule with identical tolerability with either scheduling.²⁷ Sunitinib in a 2/1 dosing schedule has been recently proven to be less toxic than the standard 4/2 dosing schedule with preserved oncological benefits (RESTORE TRIAL).²⁸

Sorafenib was the first TKI to be licensed for metastatic renal carcinoma. Escudier et al. in a multi center international phase III trial (TARGET) compared sorafenib with placebo in intermediate and low risk groups (MSKCC risk) with metastatic clear cell renal cancer in a dose of 400 mgs twice daily in 6 weekly cycles for first 24 weeks followed by 8 weekly cycles. 93% had prior nephrectomy and 81% had received cytokine therapy with either IL-2 or IFN. A PFS advantage to sorafenib (5.5 vs 2.8 months, P<0.001) was noted at interim analysis and the benefit also translated into a non-significant overall survival trend (19.3 vs 15.9 months, P=0.02). Ten and 8% discontinued treatment due to adverse effects in each arm. Dose reductions (13% vs 3%) and dose interruptions (21% vs 6%) were more with sorafenib. The dose reduction protocol was 400 mgs/day as initial step, further reducing to alternate day treatment. Two treatment related deaths was reported in the sorafenib arm.²⁹,³⁰ The PREDICT multicenter non interventional study was done to evaluate safety and efficacy of sorafenib in the practice setting. Advanced renal cancer and absence of contra indication to sorafenib were the inclusion criteria and the study did not have specific disease related exclusion criteria. This study confirmed tolerability and efficacy of sorafenib, the median PFS was 7.3 months but no overall survival data was reported. The study consequent to its heterogeneous inclusion criteria was able to demonstrate a benefit in poor risk, brain metastasis and non clear cell histology all excluded in the previously reported phase III TARGET trial.³¹ It is important to note the efficacy was assessed by the investigators using subjective criteria as opposed to contem-
porary practice of using the RECIST criteria. A phase II trial evaluated sorafenib against IFN in treatment naive metastatic renal cancer and failed to show a benefit to sorafenib; however, this trials showed a dose dependent response to sorafenib as patients progressing on a dose of 400 mgs showed a good response after increasing the dose to 600 mgs twice daily.32 Sorafenib continues to be the standard arm in several phase III trials testing third or second line agents and has been tested against tivozanib and dovitinib both found to be not superior to sorafenib.33,34

The monoclonal antibody against VEGF, bevacizumab administered 10 mgs/kg every 2 weeks with IFN 9MU thrice a week is an established first line therapy in the favourable and intermediate risk groups. The AVOREN trial tested this combination with IFN alone and demonstrated a PFS benefit of 10.2 vs 5.4 months (HR 0.63, P<0.001), but the long term results with regards to overall survival was not significant (23.3 vs 21.3 months), the authors hypothesizing lack of OS benefit to confounding by post progression second line therapy.35 The CALGB 90206 had an identical design except not requiring a placebo in the IFN arm and in not requiring a prior nephrectomy. The results were similar to the AVOREN with a significant PFS benefit (8.5 vs 5.2 months) but no OS benefit.36 In both trials the combined arms had more toxicity related issues with fatigue, anorexia, hypertension and proteinuria dominating. The exact contribution of IFN to the efficacy of the combination together with the mechanism of action remains unknown. Bevacizumab monotherapy has been tested only in phase II trials and a median PFS of 8.5 months noted.37

Axitinib is a second generation TKI against VEGFR offering greater potency and specificity in inhibition. It is currently approved as a first, second and third line agent in metastatic renal cancer. The phase III AXIS evaluated axitinib 5 mgs twice daily against sorafenib 400 mgs twice daily on patients with prior cytokine and VEGFR antagonist exposure. The PFS was longer in axitinib (8.3 vs 5.7 months, HR 0.65) but no OS advantage was demonstrable (20.1 vs 19.2 months, HR 0.96). Patient reported outcomes and toxicity related variables were identical in both groups with only 4% treatment discontinuation rate. The quantum of benefit was longer in the cytokine subgroup compared to sunitinib subgroup perhaps indicating lack of resistance to VEGFR inhibitors.38 A similar clinical activity was observed in third line setting also.39 In treatment naive patients axitinib compared to sorafenib showed a non significant trend towards better median PFS (10 vs 6.4 months) establishing clinical activity with acceptable toxicity profile.40

Two mTOR inhibitors, temsirolimus and everolimus are approved for use in renal cancer, particularly for VEGF refractory disease. The Global advanced renal cell carcinoma phase III tested temsirolimus 25 mgs weekly with IFN 3 MU thrice weekly escalated to 18MU or combination of temsirolimus 15 mgs weekly plus IFN 3MU thrice weekly escalated to 6MU. Inclusion criteria required any histology, no prior therapy and at least 3 of 6 risk factors.18% had non-clear cell histological types. The Median OS was 10.9, 7.3 and 8.4 months for temsirolimus, IFN and combination arms, however despite a significant PFS advantage and OS benefit could not be demonstrated. Toxicity profile (67% vs 78% vs 87%) favoured the temsirolimus arm, it is important to note the dose of IFN used was higher than in other trials. The best results were observed in age >65 years and in the clear cell type.41

Table 2. Practice points on clinical pharmacology of selected targeted agents.

| Table 2. Practice points on clinical pharmacology of selected targeted agents. |
|---|---|---|---|---|---|
| **Pazopanib** | **Sunitinib** | **Sorafenib** | **Axitinib** | **Everolimus** | **Temsirolimus** |
| **Dose** | 800 mgs/day | 50 mgs/day | 400 mgs BD | 5 mgs BD | 10 mgs OD |
| **Dose modification protocol** | 400 mgs/day then in 200 mgs steps | 12.5 mgs decrements | 400 mgs OD then to alternate day | 3 mgs or 2 mgsBD | 2.5 mgs OD then to 5 mgs OD |
| **Interactions** | | | | | |
| CYP3A4 inhibitors | ↓ dose to 400 mgs | 37.5 mgs | ↓ dose | ↓ dose | ↓ dose |
| CYP3A4 inducers | 87.5 mgs | | 5 mgs BD | 12.5 mgs/wk |
| **Renal dysfunction** | No dose reduction | None even in severe impairment | None required | None required | None required |
| **Hepatic dysfunction** | | | | | |
| Child A | No change | No change | No change | No change | ↓ dose |
| Child B | 200 mgs/day | No change | No change | No change | ↓ dose |
| **Cardiac dysfunction** | | | | | |
| Stop if LVEF ↓ >15% cQT interval >50 msec | Stop if LVEF >15% cQT interval >50 msec | Cardiac ischemia/ CCF | None required | None required | None required |
| **Monitoring therapy** | | | | | |
| BP- weekly | BP- weekly | BP- weekly | BP-weekly | CBC weekly |
| LFT Bi weekly | LFT each cycle | LFT Biweekly | LFT 2weekly | RFT, LFT |
| LVEF | LVEF | LVEF | Urine protein | Glucose |
| TFT | TFT | TFT | Urine protein | Bi weekly |
| Urine protein | | | | |
| **Comments** | PPI ↓ absorption by 40% | Caution in bleeding or perforation | Caution in bleeding or perforation | Avoid in GI bleed or perforation | Watch for pneumonitis and infections |
| | | | | | Watch for interstitial lung disease, bowel perforations, and infections |
The INTORSECT trial evaluated temsirolimus against sorafenib in patients failing on sunitinib, no PFS difference was demonstrable but an OS advantage to sorafenib (12 vs 16 months, P=0.01) was observed, consequently temsirolimus is not recommended as a second line therapy. The RECORD-1 trial compared everolimus vs placebo in patients progressed or intolerant to VEGF-TKI, a median PFS of 4.9 vs 1.9 months (HR 0.33) was observed but no OS benefit. The phase II RECORD-3 trial compared everolimus followed by the sunitinib, with sunitinib initial therapy followed by everolimus at progression. The median OS (22.4 vs 29.5 months, HR 1.09) supported the combination of sunitinib followed by everolimus.

Current inclusions to standard therapy

Nivolumab is a human monoclonal antibody against programmed death receptor-1 (PD-1). It blocks the interaction of PD-1 with its ligands resulting in enhanced T cell mediated tumor surveillance. The checkmate 025 trial (phase III) evaluated Nivolumab at a dose of 3 mks/kg every 2 weeks against everolimus 10 mgs/day in patients with Karnofsky performance status of >70 and pre-treated with at least 2 prior anti angiogenic agents. CNS metastasis were specifically excluded from the study. This trial showed a significant objective response difference favouring nivolumab (21.5% vs 3.9%) that translated into an overall survival benefit of 25 vs 19.6 months (HR 0.73, P=0.0018). Nivolumab had a favourable toxicity and quality of life profile than everolimus. Fatigue, nausea and pruritus were the most commonly observed side effects of nivolumab with treatment discontinuation in only 8% (13% in everolimus) of patients. Interestingly predictive marker inclusion criteria-based was not applied and the observed benefit noted irrespective of PD-L1 expression status. The above trial formed the basis for approval of nivolumab in 2015 as second line therapy for metastatic renal cancer. Cabozantinib is an inhibitor of MET, VEGFR and AXL tyrosine kinases. MET and AXL are unregulated as a consequence of VHL gene inactivation in renal cancer and the two have been postulated as possible resistance inducing factors to VEGFR therapy. A phase III RCT (METEOR) compared cabozantinib 50 mgs/day with everolimus 10 mgs/day in advanced renal cancer including those with stable brain metastasis and who had prior therapy. A benefit to cabozantinib over everolimus in the primary end point of PFS (7.4 vs 3.8 months, P 0.001) was noted with 42% reduction of disease progression. The results were reproducible in all risk categories and sub groups analysed. Though discontinuation due to adverse drug effects was nearly identical (10% vs 9%), dose reductions were 60% in cabozantinib group compared to 25% in everolimus arm. Grade 3/4 toxicities were more in the cabazantinib (68% vs 58%) with hypertension, fatigue and diarrhoea predominating. All toxicities were managed with dose reductions at 50% decrements in both arms and only 1 treatment related death reported in each study arm. Lenvatinib with everolimus is the only combination therapy approved for advanced renal cancer. In a phase II trial on pre-treated patients lenvatinib 18 mgs plus everolimus 5 mgs was tested to either drug as monotherapy. A median PFS of 14, 7.4 and 5.5 months was observed for the combination, levantinib and everolimus respectively, an OS advantage was elusive. The combination arm has more toxicities but all manageable with dose reductions.

Selection of therapy

The list of approved therapies for advanced renal cancer has rapidly increased in the past decade, but the selection of ideal therapeutic agent for an individual patient is still unclear. Treatment decisions depend upon histological type, risk category and coexistent illnesses. It is important to realise most of the trials that formed the basis for approval of these targeted therapy have excluded non clear cell histology, poor risk category and brain metastasis. It also worthwhile to note that most of the trials that evaluated second line therapy have included non clear cell histology. First line therapy in the non-clear cell group may be initiated with one of the following agents everolimus, temsirolimus or sunitinib. Temsirolimus appears to be the choice in poor risk category while sunitinib, pazopanib and bevacizumab/IFN is recommended as initial therapy in the favourable and intermediate risk category, the ideal among these may be decided by coexistent illness in the individual patient. A tabulated summary of clinical pharmacology of these agents is presented in Table 2. NCCN also recommends high dose IL-2 and sorafenib as first line therapy in carefully selected patients. IFN-α monotherapy has been found to be inferior to sunitinib as first line therapy and hence is no longer recommended. Best results are seen in the nephrectomised patients

Table 3. Selected ongoing trails in advanced RCC.

| Type          | Design                          | Clinical setting       | Status      |
|---------------|---------------------------------|------------------------|-------------|
| Phase III     | IM motion 151                   | Atezolizumab + bevacizumab vs sunitinib | First line | Active |
|               |                                 |                        | Not recruiting |         |
| Phase III     | Checkmate 214                   | Nivolumab + ipilimumab vs sunitinib | First line | Active |
|               |                                 |                        | Not recruiting |         |
| Phase III     | Javelin renal 101               | Avelumab + axitinib vs sunitinib | First line | Active |
|               |                                 |                        | Recruiting   |         |
| Phase III     | Javelin renal 101               | Avelumab + axitinib vs sunitinib | First line | Active |
|               |                                 |                        | Recruiting   |         |
| Phase II      | Pazopanib for 1 yr vs placebo after metastatectomy | First line | Recruiting |
| Phase II      | Pembrolizumab + axitinib vs sunitinib | First line | Recruiting |
| Phase II      | Pembrolizumab in clear cell and non clear cell types | First line | Recruiting |
| Phase II      | Metastatectomy in advanced RCC   | Curative               | Recruiting |
| Phase II      | Sunitinib + gemcitabine vs sunitinib in sarcomatus histology | First line | Recruiting |
| Phase II      | Pazopanib in non clear cell carcinoma | First line | Recruiting |
and the sequencing of nephrectomy and targeted therapy in patients presenting with synchronous metastasis is unclear.

Progression after first line therapy requires treatment with one of the following agents cabozantinib, nivolumab, axitinib and lenvatinib/everolimus combination. Progression after cytokine therapy may be treated with any VEGFR inhibitors or mTOR inhibitors or nivolumab, while failure after initial VEGFR therapy requires everolimus, axitinib, sorafenib or other approved agents as above. The ideal sequencing schedule of these agents is still unexplored. A list on ongoing clinical trials on advanced renal cancer is presented in Table 3.

Spurred by the benefits in metastatic disease several trials evaluating the adjuvant role of targeted agents in RCC are underway. The ASSURE and S-TRAC have published their interim results with ASSURE showing no disease free survival benefit but the S-TRAC showing a benefit particularly in the node positive and high risk cohort. The adjuvant setting continues to be an emerging indication for expanding the role of TKIs in renal carcinoma.

Conclusions

Survival outcomes for metastatic renal cancer have improved in the past decade particularly with the approval of targeted therapeutic agents. Sequential monotherapy after cytoreductive nephrectomy is the preferred approach but the ideal sequencing strategy of targeted agents remains unknown. Immunotherapy relegated to a selective role is re emerging as an important treatment approach with the licensing of nivolumab. Treatment of progressive disease beyond first and second line therapy is rapidly evolving but despite these optimism it is important to note these therapies have significant adverse effects and require diligent clinical application.

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