ABSTRACT

Historically, advanced renal cell cancer stood as a devastating diagnosis due to the paucity of systemic therapy options. Cytokine therapy including high dose IL2 and interferon alpha comprised the mainstay of advanced renal cell cancer treatment with modest benefit. Better understanding of the biology of the von Hippel-Lindau axis in the past 2 decades gave rise to antiangiogenic therapeutics starting late 2005 with currently 8 agents now approved by the FDA for the treatment of advanced renal cell cancer. Further, most recently approved is a checkpoint inhibitor – a form of “targeted immunotherapy” – which has shown significant clinical efficacy in the management of this disease, marking a resurgence of immunotherapy in a more sophisticated fashion. In this review, the historical perspective, current treatment options and what is on the horizon in the treatment of renal cell cancer will be discussed. With advances in the understanding of this disease, our treatment armamentarium has remarkably expanded, thereby improving overall prognosis of patients affected with this disease.

© 2016 The Authors. Published by ACT Publishing Group Ltd.

Key words: Renal cell cancer; Targeted therapy; Immunotherapy; VEGF inhibition; Check point inhibition

Basu A, Kim JJ, Changing Landscapes in the Treatment of Advanced Renal Cell Carcinoma. Journal of Nephrology Research 2016; 2(1): 99-108 Available from: URL: http://www.ghrnet.org/index.php/jnr/article/view/1356
targeted therapies as well as newer developments in the Renal Cell Carcinoma therapeutic space.

**VEGF TARGETING IN RENAL CELL CARCINOMA**

**Biology of Kidney Cancer: Underlying mechanisms of VEGF activation**

The activation of hypoxia pathways forms a critical underpinning for the pathogenesis of renal cancer. The inactivation of the VHL (von-Hippel-Lindau) gene through mutation or methylation is the most commonly described genetic anomaly associated with sporadic Renal Clear Cell Carcinoma reported present in almost 75-90% of the cases of this subtype. The VHL gene is responsible for degradation of the transcription factor HIF-1α (Hypoxia-Inducible Factor 1-alpha) and HIF-2α, which is a key driver of angiogenesis. The loss of VHL contributes to tumorigenesis through an unregulated buildup of the factor complex, in turn resulting in increased coding for VEGF receptors as described in the figure. This downstream activation has made VEGF inhibition an attractive target for the development of targeted agents for Renal Cell Carcinoma. Activation of other receptors such as c-met (Figure 1) are also clinically significant and intricately involved in the VEGF activation pathway.

**First generation anti-VEGF therapy**

Until the year 2002, the space for management of metastatic renal cell carcinoma was dominated by Interferon-Alfa (IFN-α) and Interleukin-2 (IL-2); systemic therapies that relied on generating a non-specific immune response. Although these options provide a small number of durable responses in the order of 5-10% based on various reports, the vast majority of patients did not respond and progressed. In 2002, Bevacizumab, a monoclonal antibody to VEGF ligand, was one of the first targeted molecules to enter clinical trials for RCC. In a landmark phase II study, high dose Bevacizumab almost doubled time to progression in patients with metastatic RCC at 4.8 months vs. 2.5 months (HR 2.55, p <0.001). This was a big proof of concept for the use of antiangiogenic therapy in patients with advanced renal cell cancer, and was followed by FDA approval for bevacizumab in combination with Interferon in the front line setting.

**Oral Small molecule tyrosine kinase inhibitors of the VEGF receptor**

With the relevance of VEGF pathway in RCC established, small molecule inhibitors, namely Tyrosine Kinase Inhibitors or TKIs were developed for more potent blockade of pro-angiogenic cellular signaling through the VEGF and PDGF pathways. The first TKI to be approved for RCC was Sorafenib in 2005. Sorafenib is a multikinase inhibitor of VEGFR, Flt-3, PDGFR, and c-KIT. Following early demonstration of tolerability and activity in phase I studies, it was evaluated in a phase II trial in patients with advanced renal cell carcinoma. At 24 weeks, almost 50% of the Sorafenib-treated patients were progression free versus 18% of placebo-treated patients. Sorafenib also demonstrated a similar progression free survival when compared with traditional Interferon therapy (5.7 months with Sorafenib vs. 5.6 months with Interferon) but with a much milder side effect profile. Also, as part of this trial, switching to Sorafenib after progression on Interferon therapy resulted in an improvement in PFS (5.3 months versus 3.6 months). Sorafenib was then finally evaluated in second line setting in a phase III trial involving 903 patients. This study showed a significant improvement in the PFS (5.8 vs. 2.8 months) in favor of Sorafenib without any statistically significant overall survival (OS) benefit (17.8 vs. 15.2 months) on an intent to treat basis, likely due to a significant months cross-over effect in the final data analysis. Sorafenib was approved by the FDA in December of 2005.

After the approval of Sorafenib, the next TKI to be approved was Sunitinib, a similar VEGFR, PDGFR, FLT-3 and c-KIT inhibitor. Sunitinib has since become the cornerstone of VEGF targeted therapy in the first line setting in renal cell carcinoma. Initial studies began in 2003, where Sunitinib showed an objective response to treatment in 3 out of 4 RCC patients treated in its first phase I trial. In the follow-up phase II study, among 68 patients with cytokine refractory metastatic RCC, almost 40% of the patients achieved partial response and an additional 27% had stable disease for 3 months and greater. This was a large step forward in the treatment of mRCC. Larger confirmatory trials were conducted such as a phase II trial of 106 patients, where 34% of the patients had a partial response with a median time to progression of 8.3 months. Fatigue and diarrhea were the most commonly reported side effects. The magnitude of responses prompted the evaluation of Sunitinib in a first line setting, where it was compared with Interferon-alpha as the standard of care treatment at the time. Sunitinib achieved an objective response rate (ORR) of 31% compared to 6% for Interferon-alpha, and PFS doubled to 11 months compared with 5 months. With these unprecedented results, Sunitinib was approved in the front line setting in Jan of 2006. Sunitinib is now largely the preferred VEGF targeted first line treatment option for mRCC. There continue to be ongoing trials focusing on improving the efficacy and tolerability of Sunitinib by modifying dosing schedule or using it in combination therapy.

Pazopanib was the next TKI to appear on the horizon around 2009. In a phase I trial, among 63 patients, of whom 12 patients had renal cell carcinoma, 2 patients had a partial response while 4 had stable disease. Further studies in phase II showed that Pazopanib had a similar efficacy profile to Sunitinib, with overall response rates around 30%-35%, and a median PFS of approximately 11.1 months to 16 months depending on previous treatment. While Pazopanib demonstrated a milder clinical side effect profile, it was associated with a higher incidence of liver injury. This led to a head to head comparison between the de-facto standard Sunitinib and Pazopanib in a phase III setting. The COMPARZ trial randomly assigned 1110 patients with metastatic clear cell carcinoma to receive Sunitinib or Pazopanib and was powered to show the non-inferiority of Pazopanib versus Sunitinib. In this study Pazopanib was shown to be non-
inferior to Sunitinib with respect to PFS, with a similar overall survival (hazard ratio for death with Pazopanib, 0.91; 95% CI, 0.76 to 1.08). Also, Pazopanib was favored with respect to quality of life (QoL) measures in 11 of 14 health-related quality-of-life domains. Although, interpretation of the QoL results was thought to be difficult due to timing of the assessments and different dosing schedule for these two agents[27].

The newest multikinase TKI that was approved by FDA for RCC is Axitinib, a powerful inhibitor of VEGFR, PDGFR and c-KIT at very low concentrations (equivalent to 1% of other TKIs such as Sunitinib), without an effect on FLT-3 or RET[29,30]. In its first multicenter phase I trial in 36 patients, Axitinib was tolerated safely and demonstrated 2 partial responses among 6 patients with advanced RCC[29]. Investigated further in phase II trials, Axitinib delivered an overall response rate (ORR) of 44% with a Median Time to Progression of 15.7 months in 52 patients with cytokine refractory mRCC. These results showed superiority in efficacy compared to other TKIs in similar cohort of patients at the level of Phase II studies[29]. The side effect profile was similar to Sunitinib with diarrhea, fatigue, nausea and hypertension being the most commonly reported and what are now known as mostly VEGF inhibitor class effects[31]. Interestingly, several studies have reported a correlation between hypertension and response to treatment with Axitinib[32]. Axitinib was also effective at eliciting objective response in patients who had disease progression on treatment with Sorafenib, where it had an ORR of approximately 23%, with a median PFS of 7.4m. 74% of patients on this study had been through two or more systemic months therapies prior to Axitinib[29]. Axitinib was also effective at eliciting objective response in patients between hypertension and response to treatment with Axitinib[32].

### Table 1: FDA Approved Targeted Therapies for Renal Cell Carcinoma: Selected studies.

| Agent     | Setting                  | Comparison(s)                  | Phase/Design                      | PFS                  | Year | Ref. |
|-----------|--------------------------|--------------------------------|-----------------------------------|----------------------|------|------|
| Bevacizumab | Front-Line or Prior IL-2 Therapy | Placebo                        | Phase II, Randomized Controlled Trial | 4.8 m (Bev)          | 2003 | 9    |
|           |                          |                                |                                   | 2.5 m (Placebo)      |      |      |
|           | Frontline, no prior therapy | Bev + IFN vs IFN               | Phase III, Randomized controlled trial | 8.5 m (Bev+IFN)     | 2008 | 10   |
|           |                          |                                |                                   | 5.2 m (IFN)          |      |      |
|           | Frontline, no prior therapy | Bev + IFN vs IFN               | Phase III, Randomized controlled trial | 10.2 m (Bev+IFN)    | 2007 | 43   |
|           |                          |                                |                                   | 5.4 m (IFN)          |      |      |
|           | Frontline, or prior therapy | Placebo                        | Phase I, Randomized Discontinuation Trial | NR                  | 2006 | 14   |
| Sorafenib  | Front Line only           | IFN                            |Phase II, Open label               | 5.7 m(33)            | 2009 | 15   |
|           |                          |                                |                                   | 5.6 m (IFN)          |      |      |
| Sunitinib  | Front Line only           | IFN                            |Phase II, Open label               | 5.5 m (Sorafenib)    | 2007 | 16   |
|           |                          |                                |                                   | 2.8 m (Placebo)      |      |      |
|           | Front Line only           | IFN                            | Phase II, Open label               | 5.7 m(33)            | 2009 | 15   |
|           |                          |                                |                                   | 5.5 m (Sorafenib)    |      |      |
| Pazopanib  | Front Line only           | IFN                            | Phase II, Open label               | 5.5 m(33)            | 2009 | 15   |
|           |                          |                                |                                   | 5.5 m (Sorafenib)    |      |      |
|           | Front Line only           | IFN                            | Phase II, Open label               | 5.5 m(33)            | 2009 | 15   |
|           |                          |                                |                                   | 5.5 m (Sorafenib)    |      |      |
|           | Prior IL-2 Therapy        | Sunitinib                      | Phase II, Randomized Controlled Trial | 10.5 m(Sorafenib)    | 2013 | 27   |
|           |                          |                                |                                   | 10.5 m(Sorafenib)    |      |      |
|           | Prior Sorafenib           | Sunitinib                      | Phase II, Randomized Controlled Trial | 10.5 m(Sorafenib)    | 2013 | 27   |
|           |                          |                                |                                   | 10.5 m(Sorafenib)    |      |      |
|           | Second Line               | Placebo                        | Phase II, Randomized Controlled Trial | 8.3 m (Axitinib)     | 2013 | 45   |
|           |                          |                                |                                   | 5.7 m (Sorafenib)    |      |      |
|           | Second Line, after cytokine failure | N/A                           | Phase II, Randomized Controlled Trial | 8.3 m               | 2003 | 20   |
|           |                          |                                |                                   | 5.2 m (Sorafenib)    |      |      |
|           | Front Line only           | IFN                            | Phase II, Randomized Controlled Trial | 11 m (Sorafenib)     | 2007 | 21   |
|           |                          |                                |                                   | 5.5 m (IFN)          |      |      |
|           | Front Line only           | IFN                            | Phase II, Randomized Controlled Trial | 8.3 m               | 2013 | 20   |
|           |                          |                                |                                   | 5.7 m (Sorafenib)    |      |      |
|           | Front Line only           | IFN                            | Phase II, Randomized Controlled Trial | 8.3 m               | 2013 | 20   |
|           |                          |                                |                                   | 5.7 m (Sorafenib)    |      |      |
|           | Prior IL-2 Therapy        | Sunitinib                      | Phase II, Randomized Controlled Trial | 10.5 m(Sorafenib)    | 2013 | 27   |
|           |                          |                                |                                   | 10.5 m(Sorafenib)    |      |      |
|           | Prior Sorafenib           | Sunitinib                      | Phase II, Randomized Controlled Trial | 10.5 m(Sorafenib)    | 2013 | 27   |
|           |                          |                                |                                   | 10.5 m(Sorafenib)    |      |      |
|           | Second Line               | Sunitinib                      | Phase II, Randomized Controlled Trial | 8.3 m               | 2013 | 20   |
|           |                          |                                |                                   | 5.7 m (Sorafenib)    |      |      |

**When switched to Sorafenib on progression. ^NS- Not significant. Legend: IFN: Interferon, m – months.**
renal cell carcinoma along with others.[37] In a phase II randomized discontinuation trial it showed impressive efficacy in the overall trial population (n=272), almost half of which were treatment naïve (54%) with predominantly clear cell histology (83%). This trial reported an Objective Response Rate of 24% and PFS of 11.7 months.[38]

Tivozanib reached the crucial phase III TIVO-1 trial, where it compared head to head with Sorafenib. Of this multicenter trial of 517 patients, the overwhelming majority of patients were from Eastern Europe.[39] This was also an open-label trial and patients who progressing on Sorafenib were given the opportunity to cross over to Tivozanib.[39] Although the trial met the primary endpoint of demonstrating superior PFS with Tivozanib (11 months vs. 9.1 months), Sorafenib showed superior OS; perhaps as a byproduct of a cross over effect, where almost 150 patients on Sorafenib had crossed over to Tivozanib. Unfortunately, the conduct of this trial in some resource limited settings meant that patients who were on the Tivozanib arm did not have access to other VEGF directed therapies to switch over to, likely creating an imbalance in VEGF antagonist exposure between the two arms.

mTOR Targeting in Renal Cell Carcinoma

As described above, the PI3K /AKT/mTOR pathway has been a pro-growth signaling pathway activated in many cancers and in the majority of RCCs due to its regulation via HIFs as well. The FDA approved targeted agents developed towards this pathway include Temsirolimus and Everolimus.

Oral mTOR Inhibitors

Temsirolimus was developed in parallel with the earlier VEGF TKIs, in the early 2000s, where it was found to be effective in RCC in its initial phase I studies. Temsirolimus demonstrated confirmed partial response in a patient with metastatic treatment refractory RCC and a patient with breast cancer among 24 total patients in an early Phase I study.[42] However, Its FDA approval for mRCC came from a Phase III randomized clinical trial of 626 patients with mRCC with poor and Intermediate MKSCC prognostic criteria. In this high to intermediate risk population the median survival was 7.3 months in the Interferon group, 10.9 months in the Temsirolimus group, and 8.4 months with a combination-therapy group.[50] This trial forms the basis of the FDA approval of this agent and also for the NCCN’s recommendation of Temsirolimus as a first line agent in patients with poor prognostic criteria.

In addition to Temsirolimus, Everolimus is the other mTOR inhibitor approved for treatment of mRCC. Everolimus was evaluated in the RECORD-1 trial, a double blind randomized controlled phase III trial conducted in 410 patients who had progressed on first line Sunitinib or Sorafenib. There was a significant improvement with Everolimus, with a doubling of median progression free survival to 4.0 months [95% CI 3.7-5.5] compared with 1.9 months [95% CI 1.8-1.9] on best supportive care. Stomatitis, fatigue and rash, a class effect of mTOR inhibitors occurred in a large minority of patients (40%, 25% and 20% respectively). Pneumonitis was the most common severe side effect; observed in 8 out of 272 patients on Everolimus reporting grade III events.[51] mTOR pathway inhibition is a promising strategy and continues to be evaluated in renal cell carcinoma. mTOR inhibition has also been studied in non-clear cell renal cell cancer with evidence for modest benefit.[52]

First Generation Systemic Immunotherapies

Interleukin and Interferon Therapy

Although cancer immunotherapy has become the center of attention in the recent years, the early evidence of its effectiveness in cancer treatment and in RCC comes from the use of interferon-alpha (IFN-α) and interleukin 2 (IL-2) in the earlier trials[53]. IL-2 stimulates a stress response to infection in the human body such as through the stimulation of T-cell proliferation. The first demonstration of efficacy was in a young woman with metastatic melanoma in 1984, where IL-2 caused diffuse shrinkage of her tumors and a complete response.
that persisting three decades[14]. IL-2 was also the first successful demonstration that extraneous activation of the immune system can have spectacular effects on cancer control. Renal Cell Carcinoma is an immunogenic tumor and has also benefited from IL-2 treatment. High dose IL-2 was developed for human use and approved by FDA in 1996 for metastatic Renal Cell Carcinoma on the basis of encouraging and durable responses in a small fraction of patients (5-10%). Response rates as high as 28% have been reported in recent studies and expression of biomarkers such as PD-L1 or CA9 may serve as predictors of overall response rates[59, 60]. Significant acute toxicity and lack of benefit to a vast majority of treated patients continue to remain the main issue with this agent.

**SECOND GENERATION TARGETED IMMUNOTHERAPIES: CHECK POINT INHIBITION**

First generation immune therapy with Interferon-α and IL-2 only provided limited benefit due to the ability of cancer cells to escape an extrinsic cellular anti-tumor response—a process now recognized as immunoediting[25]. This process, driven by both the innate and adaptive immune systems comes into play when intrinsic cellular checkpoints such as tumor suppressors fail to contain abnormal growth, leading to nascent cancer cell formation Figure 2. Initial stages involve elimination of the tumor, but on rare occasions, the tumor cells that cannot be completely eliminated enter the ‘equilibrium’ phase, during this phase, the constant fight between the immune system and the tumor achieves an evolution of the final form of the tumor which is able to ‘escape’ immune surveillance. One of these mechanisms is through the expression of immune checkpoint modulators[26, 27].

**Table 2** Examples of ongoing trials evaluating new therapies in metastatic renal cell carcinoma.

| Investigational Agent(s) | Mechanism of Action | Study Name / Description | ClinicalTrials.gov Identifier |
|--------------------------|---------------------|--------------------------|-------------------------------|
| HyperAcute-Renal Immunotherapy | Cancer vaccine | A Phase I Study of HyperAcute-Renal (HAR) Immunotherapy In Patients With Metastatic Renal Cell Cancer | NCT02035338 |
| Nivolumab, Ipilimumab | Anti PD1 ab + Anti CTLA4 ab | Nivolumab+Ipilimumab + Bevacizumab or Nivolumab + Ipilimumab in Metastatic Renal Cell Carcinoma (mRCC) | NCT02210117 |
| Nivolumab | Anti PD1 ab | A Study of Anti-PD1 (Nivolumab) Therapy as Pre- and Post-operative Therapy in Metastatic Renal Cell Cancer (ADAPTeR) | NCT02446680 |
| Intuvax( vaccine) | Cancer Vaccine | Intratumoral Vaccination With Intuvax Pre-nephrectomy Followed by Sunitinib Post-nephrectomy vs Sunitinib Post-nephrectomy in Newly Diagnosed Metastatic Renal Cell Carcinoma (mRCC) (MERECA) | NCT02432846 |
| Varilimumab | anti-CD27 Ab | A Study of Varilimumab (Anti-CD27) and Sunitinib in Patients With Metastatic Clear Cell Renal Cell Carcinoma | NCT02886111 |
| PT2385 | HIF-2A Inhibitor | A Phase 1, Dose-Escalation Trial of PT2385 Tablets In Patients With Advanced Clear Cell Renal Cell Carcinoma | NCT02923980 |
| Atezolimumab | Anti-PD-L1 Ab | A Study of MPDL3280A (Anti-Programmed Death Ligand 1 [PD-L1] Antibody) in Combination With Bevacizumab Versus Sunitinib in Patients With Untreated Advanced Renal Cell Carcinoma | NCT02420821 |
| Avelumab | anti PD-L1 ab | Avelumab in Metastatic or Locally Advanced Solid Tumors [JAVELIN Solid Tumor] | NCT01772004 |
| CAR-T cells for RCC | Anti-VEGFR2 Engineered CD8 + Cells | CAR-T Cell Receptor Immunotherapy Targeting VEGFR2 for Patients With Metastatic Cancer | NCT01218867 |

CTLA-4 inhibitors

The process of immune surveillance involves the identification of cancerous cells and then phagocytosis by antigen presenting cells (APCs). A protein that regulates this process is the CTLA-4; or Cytotoxic T lymphocyte antigen 4. CTLA-4 behaves like a brake for the immune system, preventing excessive activation. However, this brake is utilized by the cancer cells to override the immune response. CTLA-4 antibodies have been developed which release these brakes from the immune system. Figure 1 describes the mechanism of action of these drugs. Drugs in this category such as Ipilimumab have now obtained FDA approval in treatment of cancers such as melanoma.

Some of the earliest clinical evidence for Ipilimumab came from renal cell carcinoma patients, in 2007, Yang et al reported a phase II study where 5/40 patients on Ipilimumab at 3 mg/kg had a partial response to therapy with the drug[64]. These responses were in patients who had progressed on IL-2 therapy; however it appeared that responses were highly correlated with the frequency of immune related adverse events (irAEs) with 30% of those with adverse effects having responses while no patients without irAEs had any responses[64]. However, Ipilimumabmonotherapy for renal carcinoma remained on the backburner until the advent of the newer checkpoint inhibitors.

PD-1/PD-L1 checkpoint inhibitors

One of the most promising agents to arrive on the horizon for cancer immunotherapy have been the PD-1 checkpoint inhibitors. PD-1/PD-L1 stand for programmed death receptor and programmed death ligand respectively. The PD-1 receptor is present on T cells and the PD-L1 ligand is present on the surface of antigen presenting cells (APC) or on Tumor cells. The interaction is immunosuppressive, and tumors expressing high concentrations are able to subdue the cellular immune response against tumors[65]. PD-1 inhibitors had showed significant activity in the recent years in tumors such as melanoma but data presented in June 2015 now demonstrates significant success in the treatment of a large number of tumor types including NSCLC, liver cancer, advanced head and neck cancers, and particular subsets of colorectal cancer patients.

Nivolumab, a well-tolerated anti-PD-1 antibody was investigated in mRCC settings and is first immune checkpoint inhibitor now
approved for metastatic renal carcinoma. Phase I trials showed good tolerability, with some responses in patients with mRCC in the cohort[69]. In a recent phase II study patients with mRCC refractory to treatment with VEGF inhibition were randomized 1:1 to Nivolumab in increasing doses of 0.3mg/kg, 2 mg/kg or 10 mg/kg. There was an ORR around 20%; but with a median OS of 25.5 months in patients being treated with Nivolumab 3 mg/kg group and 24.7 months with Nivolumab 2 mg/kg[67]. Side effects with PD-1 blockade is a stark contrast to those with agents such as IL-2 and Interferon, with 19/168 patients in the trial experiencing a grade III/IV adverse events, most commonly fatigue[67]. A smaller phase II trial with 34 patients and Nivolumab dosed at either 1 mg/kg or 10 mg/kg found an ORR of 29%, with an additional 27% with disease stabilization, and a median OS of 22.4 months[69]. This was finally followed by the CheckMate025 study, which compared Nivolumab to Everolimus in patients with advanced clear cell renal carcinoma that had progressed on first and second line anti-angiogenic therapies. 821 patients participated, and were randomized to receive either Nivolumab at 3mg/kg every 2 weeks or Everolimus 10 mg tablets daily. Median overall survival with Nivolumab was 25.0 months (95% CI 21.8 – Not estimable) and 19.6 months (95% CI 17.6 to 23.1) with Everolimus resulting in a Hazard Ratio [HR] of 0.73 (98.5% CI, 0.57 to 0.93; P=0.002), indicating superiority of the Nivolumab regimen. As expected, this benefit also came with minimal risks, with Grade III/IV toxicity rates being 19% in the immune arm in comparison to 37% in patients on Everolimus[69]. This highly encouraging data has led to the FDA approval of Nivolumab for metastatic renal cancer patients.

Similar to PD1 inhibition Blockade of the PD-L1 ligand is also being investigated in kidney cancer. This includes agents such as MPDL3280A (now called atezolizumab). In a Phase I trial of 277 patients, including 69 patients with metastatic renal cell carcinoma administered MPDL3280A, there was an objective response rate of 14% with 8 out of 58 evaluable RCC patients with partial responses. The median response was 54 weeks, a very robust duration – and a characteristic of this therapy type, where benefit to those patients tends to have durable responses. PFS at 24 weeks was 53% in this cohort[69].

### COMBINATIONS OF CHECKPOINT INHIBITORS

Given the relatively mild side effect profile of the newer targeted treatments and their clear efficacy, there is growing interest in evaluating combination therapies of immune agents, either with each other or with anti-angiogenic agents in mRCC.

Recent updated results from the Checkmate 016 study evaluated combination of the CTLA-4 and anti-PD1 checkpoint inhibitors Ipilimumab and Nivolumab respectively. Three groups were compared, with Nivolumab and Ipilimumab one low and high doses of 1mg/kg and 3 mg/kg each. The combined higher dose combination was not tolerated secondary to toxicity. Nivolumab (3mg/kg) + Ipilimumab (1mg/kg) had 34% patients with Grade 3-4 Adverse effects while it was 64% for the Nivolumab (1mg/kg) + Ipilimumab (3mg/kg) dosing. The Objective response rates in both arms were similar at around 40%, with a median PFS of 30-36 weeks[69]. Patients were intermediate-favorable per MSKCC criteria and approximately half had been treatment naïve.

The anti-PDL1 antibody MPDL3280A described earlier was also investigated in combination with Bevacizumab in a recent phase Ib study where it demonstrated good tolerability, with no grade 3-4 adverse events reported in the limited sample of 12 patients. Also, there was an objective response rate of 40% in first –line patients. The data is still early, but encouraging for further follow up. A phase II was ongoing when last reported[70].

### SEQUENCING OF AGENTS AND ROLE OF ADJUVANT THERAPY

#### Adjuvant Therapy for Renal Cell Carcinoma

The concept of adjuvant chemotherapy for renal cell carcinoma was a subject of great interest due to the in-principle plausibility of benefit. A recent large trial (ASSURE) was conducted on 1943 patients with subjects randomized to 1:1:1 of Sunitinib, Sorafenib and placebo for locally advanced renal cancer in an adjuvant setting. Most recent updates from the trial unfortunately suggest no statistically significant benefit in either group[73]. The trial was also subject to significant rates of attrition due to side effects, with 26% in the experimental arms and thus had dose reductions (e.g. Sunitinib at 37.5 mg and Sorafenib at 400 mg ) reducing the attrition rate to 14%, however this is unlikely to have had an impact on the overall hazard ratio[72]. Although this has been a negative trial, it has not shut the door on the concept of adjuvant therapy for Renal Cell Carcinoma. It is still to be determined if new generation checkpoint inhibitors will be more tolerable and improve outcomes in this population.

#### Sequencing of therapies

The relatively recent approval of the newer mTOR and VEGF inhibitors has resulted in some confusion on the most efficacious sequence of the use of these agents. There are now 7 FDA approved agents spanning both classes, all developed and approved within a few years. In first line settings, apart from Interleukin and Interferon-Sunitinib, Temsirolimus, Pazopanib, Axitinib and Sorafenib have all shown activity. The COMPARZ trial of Sunitinib and Pazopanib referred above did not show any significant benefit to either but sets up a case for patient-physician discussion based on side effect profiles[74]. A trial of Everolimus followed by Sunitinib or Vice versa has been explored and presented recently as the RECORD-3 trial. The trial analyzed individual PFS benefits as well as the OS benefit for each of the sequences in treatment. Median OS was 22.4 months for Everolimus followed by Sunitinib and 29.5 months for Sunitinib followed by Everolimus (HR, 1.09; 95% CI, 0.87-1.37), this suggests that the current practice of starting with VEGF targeted therapy maybe the most effective approach[75].

In a second line setting the INTORSECT trial evaluated Temsirolimus and Sorafenib inpatients who had previously progressed on Sunitinib therapy. Although the objective response rate for both arms were 8%, and Temsirolimus had a similar median PFS of 4.3 months versus 3.9 months for Sorafenib, there was an OS benefit for Sorafenib (16.6 months) compared with Temsirolimus (12.3 months) demonstrating that even disease progressive with VEGF blockade with one agent can respond to subsequent therapy[69]. Challenges to interpreting this data as a verdict on VEGF vs mTOR blockade in the second line setting include the proven efficacy of Axitinib in the AXIS trial and Everolimus in the RECORD-1 trial and there remains scope for further research. Another question for oncologists has been potential Interchangeability between the FDA approved 2 mTOR inhibitors. A trial has now compared Everolimus and Temsirolimus as second line after therapy with an anti-VEGF TKI, which seems to suggest that Everolimus may lead to a superior OS (24.2 months vs 12.1months) [76]. A meta-analysis extrapolating data from available studies also supports the direction of this effect[77].
INVESTIGATIONAL THERAPIES FOR KIDNEY CANCER

Apart from the anti-angiogenic and immune checkpoint therapies discussed so far, there are several other promising targeted therapies that under development. Some of the strategies employed are either medications that can target resistance mechanisms to VEGF inhibitors, or act on novel molecular targets of renal cell carcinoma.

Resistance mechanisms to the commonly targeted processes such as VEGF inhibition are under intense investigation, a next generation of molecules are being developed which may target these resistance mechanisms[79]. Endoglin is an accessory receptor for TGF-Beta1 and has been shown to be expressed in higher amounts in tumors with advanced stage renal carcinoma, it is assumed to be an important piece in developing resistance to VEGF inhibitors[77]. In a recent phase I study of TRC105 in 18 mRCC patients refractory to multiple agents including prior Axitinib, an anti-Endoglin monoclonal antibody co-administered with Axitinib showed 18% RECIST response rate with almost half (47%) having a >10% decrease in tumor burden[76].

Another agent that shares a similar mechanism, by decreasing resistance to VEGF therapies is Dalantercept, which is an activating receptor kinase-1 (ALK1) inhibitor: a recent phase I trial on 29 patients’ refractory to previous mTOR, VEGF or immune therapies has shown very encouraging results. Dalantercept was combined with Axitinib at 5 mg PO twice a day dosing, there were no severe dose limiting toxicities with incremental dosing. While the ORR was 25% for the whole group, the median PFS was not reached at the maximum tolerated dose as of data reported until May 2015[79]. The study will now expand to a randomized phase II setting comparing Dalantercept versus placebo in combination with Axitinib[79].

There has been an observation of increased c-MET expression in many renal cell carcinomas that are resistant to VEGF therapy, Cabozantinib is a small molecule tyrosine kinase inhibitor with action only on the VEGF receptor but also inhibitory action on Met, RET, AXL, KIT and FLT 3, however the c-MET inhibition alongwith VEGF inhibition are likely key elements of efficacy in renal cell carcinomia[79]. A Phase I study of Cabozantinib in heavily pretreated patients with RCC (n = 25) demonstrated that it was tolerated well, with a partial response rate of 28%, and median progression free survival of 12.9 months, and mean overall survival of 15.0 months[81]. This was followed by a pivotal phase III randomized controlled trial of Cabozantinib versus Everolimus in 658 patients, in this trial median progression free survival with Cabozantinib was 7.4 months in comparison to 3.8 months with Everolimus- for a Hazard Ratio [HR] of 0.58 (95% CI 0.45-0.75) in favor of Cabozantinib. PFS also translated into an improvement in overall survival as well. Adverse events requiring discontinuation of therapy were similar across both arms ( 9% for Cabozantinib vs 10% Everolimus)[82]. Cabozantinib has been granted breakthrough therapy designation by the FDA and may likely be the 9th targeted therapy to be approved for kidney cancer.

While the checkpoint inhibitors have been recently discovered to augment the immune response against cancers, thus improving disease control, an alternative approach that preceded this discovery was the development of peptide vaccines for cancer. There are certain proteins expressed by malignant cells that can help the body identify the underlying process. These antigens, called Tumor Associated Antigens (TAAs), have been thought of as antigenic stimulants for an immune response. IMA901 is a synthetic RCC-specific, multiantigen peptide vaccine that contains tumor associated peptides derived from the antigens overexpressed in RCC. In phase I, among 28 patients, 20 patients had a T cell response to at least one tumor associated antigen and 8 had responses to more than one. Studies in RCC showed a correlation of multi-tumor associated antigen responses with OS, and there are phase III trials now ongoing to further evaluate its efficacy[83]. Similar such agents include Hyperacute- Renal Immunotherapy, and Intuvax. Unpublished communication from the maker suggests that of 11 patients on the phase I/II trial of this agent in 2012, 5 patients continue to survive until 2015 for a median overall survival of 29.8 months versus expected 15.2 months[84].

Finally, as we discussed before, hypoxia inducible factors form the final common product of the hypoxia axis activation due to various mutations associated with renal cell cancer. Until recently there were no potent inhibitors of HIF. However, currently some trials are recruiting to evaluate efficacy of newly developed oral HIF inhibitors (refer table).

DISCUSSION

Metastatic renal cell cancer has become a much more “treatable disease” in the past 10 years than it has ever been in the past. The last decade has seen an explosion of therapeutic options in patients with metastatic renal cell carcinoma. These developments have been led by our deeper understanding of the underlying biology of mRCC, ultimately leading to the development of targeted agents for the VEGF/angiogenesis pathway. More recently achieved is our improved understanding of immune mechanism in the setting of cancer, leading to approval of checkpoint inhibitors in not only kidney cancer but also in multiple other cancers including lung and melanoma. Other immunotherapy approaches including vaccines as well as combination approaches are currently undergoing investigation at this time with promising early results. Importantly, studies are also ongoing to identify biomarkers for response and toxicity to all therapies involving kidney cancer as well as cancer therapy as a whole in pursuit of “personalizing” cancer care. Going forward, as a result of the rapid development of these agents, several questions will need to be answered which include optimal combination or sequencing of therapeutic agents, effectiveness of available agents in an adjuvant or neoadjuvant setting, and the development of biomarkers to identify those who may or may not respond and with what degree of tolerance. Overall, however, it is without a doubt that the landscape of renal cell carcinoma has changed significantly for the better and there are definitely more exciting times ahead.

CONFLICT OF INTERESTS

There are no conflicts of interest with regard to the present study.

REFERENCES

1. Ferlay J, Soerjomataram I, Dikshit R, et al. Cancer incidence and mortality worldwide: sources, methods and major patterns in GLOBOCAN 2012. International journal of cancer Journal international du cancer 2015; 136(5): E359-86.
2. Siegel RL, Miller KD, Jemal A. Cancer statistics, 2015. CA: a cancer journal for clinicians 2015; 65(1): 5-29.
3. Siegel R, Naishadham D, Jemal A. Cancer statistics, 2012. CA: a cancer journal for clinicians 2012; 62(1): 10-29.
4. Sanfilippo KM, McGtigue KM, Fidler CJ, et al. Hypertension and obesity and the risk of kidney cancer in 2 large cohorts of US men and women. Hypertension 2014; 63(5): 934-41.
5. Motzer RJ, Rini BI, McDermott DF, et al. Nivolumab for Metastatic Renal Cell Carcinoma: Results of a Randomized Phase II Trial. Journal of clinical oncology: official journal of the American Soci-
6. Linehan WM, Srinivasan R, Schmidt LS. The genetic basis of kidney cancer: a metabolic disease. Nature reviews Urology 2010; 7(5): 277-85.

7. Ohh M, Kazin WG Jr. VHL and kidney cancer. Methods in molecular biology 2003; 222: 167-83.

8. Bukowski RM. Cytokine therapy for metastatic renal cell carcinoma. Seminars in urologic oncology 2001; 19(2): 148-54.

9. Yang JC, Haworth L, Sherry RM, et al. A randomized trial of bevacizumab, an anti-vascular endothelial growth factor antibody, for metastatic renal cancer. The New England journal of medicine 2003; 349(5): 427-34.

10. Rini BI, Halabi S, Rosenberg JE, et al. Bevacizumab plus interferon alpha compared with interferon alpha monotherapy in patients with metastatic renal cell carcinoma: CALGB 90206. Journal of clinical oncology: official journal of the American Society of Clinical Oncology 2008; 26(33): 5422-8.

11. Gollob JA. Sorafenib: scientific rationales for single-agent and combination therapy in clear-cell renal cell carcinoma. Clinical genitourinary cancer 2005; 4(3): 167-74.

12. Larkin JM, Eisen T. Renal cell carcinoma and the use of sorafenib. Therapeutics and clinical risk management 2006; 2(1): 87-98.

13. Strumberg D, Voliotis D, Moeller JG, et al. Results of phase I pharmacokinetic and pharmacodynamic studies of the Raf kinase inhibitor BAY 43-9006 in patients with solid tumors. International journal of clinical pharmacology and therapeutics 2002; 40(12): 580-1.

14. Ratain MJ, Eisen T, Stadler WM, et al. Phase II placebo-controlled randomized discontinuation trial of sorafenib in patients with metastatic renal cell carcinoma. Journal of clinical oncology: official journal of the American Society of Clinical Oncology 2006; 24(16): 2505-12.

15. Escudier B, Szczylik C, Hutson TE, et al. Randomized phase II trial of first-line treatment with sorafenib versus interferon Alfa-2a in patients with metastatic renal cell carcinoma. Journal of clinical oncology: official journal of the American Society of Clinical Oncology 2009; 27(8): 1280-9.

16. Escudier B, Eisen T, Stadler WM, et al. Sorafenib in advanced clear-cell renal-cell carcinoma. The New England journal of medicine 2007; 356(2): 125-34.

17. Bukowski RM. How I treat renal cell carcinoma. Journal of oncology practice / American Society of Clinical Oncology 2008; 4(3): 150-2.

18. Faivre S, Delbaldo C, Vera K, et al. Safety, pharmacokinetic, and antitumor activity of SU11248, a novel oral multitarget tyrosine kinase inhibitor, in patients with cancer. Journal of clinical oncology: official journal of the American Society of Clinical Oncology 2006; 24(1): 25-35.

19. Motzer RJ, Michaelson MD, Redman BG, et al. Activity of SU11248, a multitargeted inhibitor of vascular endothelial growth factor receptor and platelet-derived growth factor receptor, in patients with metastatic renal cell carcinoma. Journal of clinical oncology: official journal of the American Society of Clinical Oncology 2006; 24(1): 16-24.

20. Motzer RJ, Rini BI, Bukowski RM, et al. Sunitinib in patients with metastatic renal cell carcinoma. JAMA 2006; 295(21): 2516-24.

21. Motzer RJ, Hutson TE, Tomczak P, et al. Sunitinib versus interferon alpha in metastatic renal-cell carcinoma. The New England journal of medicine 2007; 356(2): 115-24.

22. Lee CH, Motzer RJ. Sunitinib as a paradigm for tyrosine kinase inhibitor development for renal cell carcinoma. Urologic oncology 2014.

23. Jae-Lyun Lee MKK, Inkeun Park, Jin-Hee Ahn, Dae Ho Lee, Hoon-Mo Ryoo, Cheryn Song, Bum-Sik Hong, Jun Hyuck Hong, Hanjung Ahn. Randomized phase II trial of sunitinib four-week on and two-week off versus two-week on and one-week off in metastatic clear cell type renal cell carcinoma: RESTORE trial. ASCO GU 2015; Orlando; 2015.

24. Hurwitz HI, Dovlati A, Saini S, et al. Phase I trial of pazopanib in patients with advanced cancer. Clinical cancer research: an official journal of the American Association for Cancer Research 2009; 15(12): 4220-7.

25. Sternberg CN, Davis ID, Mardiak J, et al. Pazopanib in locally advanced or metastatic renal cell carcinoma: results of a randomized phase III trial. Journal of clinical oncology: official journal of the American Society of Clinical Oncology 2010; 28(6): 1061-8.

26. Hutson TE, Davis ID, Machiels JP, et al. Efficacy and safety of pazopanib in patients with metastatic renal cell carcinoma. Journal of clinical oncology: official journal of the American Society of Clinical Oncology 2010; 28(3): 475-80.

27. Motzer RJ, Hutson TE, Celli D, et al. Pazopanib versus sorafenib in metastatic renal-cell carcinoma. The New England journal of medicine 2013; 369(8): 722-31.

28. Motzer RJ, McCann L, Deen K. Pazopanib versus sorafenib in renal cancer. The New England journal of medicine 2013; 369(20): 1970.

29. Gunmarsson O, Pfanzelter NR, Cohen RB, Keefe SM. Evaluating the safety and efficacy of axitinib in the treatment of advanced renal cell carcinoma. Cancer management and research 2015; 7: 65-73.

30. Rugo HS, Herbst RS, Liu G, et al. Phase I trial of the oral antiangiogenesis agent AG-013736 in patients with advanced solid tumors: pharmacokinetic and clinical results. Journal of clinical oncology: official journal of the American Society of Clinical Oncology 2005; 23(24): 5474-83.

31. Rixe O, Bukowski RM, Michaelson MD, et al. Axitinib treatment in patients with cytokine-refractory metastatic renal-cell cancer: a phase II study. The Lancet Oncology 2007; 8(11): 975-84.

32. Rini BI, Cohen DP, Lu DR, et al. Hypertension as a biomarker of efficacy in patients with metastatic renal cell carcinoma treated with sunitinib. Journal of the National Cancer Institute 2011; 103(9): 763-73.

33. Rini BI, Melichar B, Fishman MN, et al. Axitinib dose titration: analyses of exposure, blood pressure and clinical response from a randomized phase II study in metastatic renal cell carcinoma. Annals of oncology: official journal of the European Society for Medical Oncology / ESMO 2015.

34. Joosten SC, Hamming L, Soetekouw PM, et al. Resistance to sunitinib in renal cell carcinoma: From molecular mechanisms to predictive markers and future perspectives. Biochimica et biophysica acta 2015; 1855(1): 1-16.

35. Welti J-C, Gourhounen M, Powles T, et al. Fibroblast growth factor 2 regulates endothelial cell sensitivity to sunitinib. Oncogene 2011; 30(10): 1183-93.

36. Mehta A, Sonpavde G, Escudier B. Tivozanib for the treatment of renal cell carcinoma: results and implications of the TIVO-1 trial. Future oncology 2014; 10(11): 1819-26.

37. Eskens FA, de Jonge MJ, Bhargava P, et al. Biologic and clinical activity of tivozanib (AV-951, KRN-951), a selective inhibitor of VEGF receptor-1, -2, and -3 tyrosine kinase activity of tivozanib (A V-951, KRN-951), a selective inhibitor of VEGF receptor-1, -2, and -3 (1751): 1819-26.

38. Nosov DA, Esteves B, Lipatov ON, et al. Antitumor activity and pharmacokinetic and pharmacodynamic results. Journal of the American Society of Clinical Oncology / ESMO 2015.

39. Motzer RJ, Nosov D, Eisen T, et al. Tivozanib versus sorafenib as initial targeted therapy for patients with metastatic renal cell carcinoma: results from a phase III trial. Journal of clinical oncology: official journal of the American Society of Clinical Oncology 2010; 28(3): 1678-85.

40. Motzer RJ, Nosov D, Eisen T, et al. Tivozanib versus sorafenib as initial targeted therapy for patients with metastatic renal cell carcinoma: results from a phase III trial. Journal of clinical oncology: official journal of the American Society of Clinical Oncology 2013; 31(30): 3791-9.

41. Hutson TE, Escudier B, Esteban E, et al. Randomized phase III trial of temsirolimus versus sorafenib as second-line therapy after sunitinib in patients with metastatic renal cell carcinoma. Journal of
clinical oncology: official journal of the American Society of Clinical Oncology 2014; 32(8): 760-7.

41. Daniel Yick Chin Heng CW, Frede Donskov, Brian I. Rini, Jae-Lyun Lee, Georg A. Bjarnason, Benoit Beuselinck, Martin Smorgoniewicz, Ajijai Shivaram Alva, Sandy Srinivas, Lori Wood, Haru Yamamoto, D. Scott Ernst, Sumanta Kumar Pal, Takeshi Yuasa, Reuben James Broom, Ravindran Kanveswaran, Aristotelis Biamis, Jennifer J. Knox, Toni K. Choueiri . Third-line therapy in metastatic renal cell carcinoma: Results from the International mKCC Database Consortium. ASCO GU; 2015; Orlando: JCO; 2015.

42. Raymond E, Alexandre J, Faivre S, et al. Safety and pharmacokinetics of escalated doses of weekly intravenous infusion of CCI-779, a novel mTOR inhibitor, in patients with cancer. Journal of clinical oncology: official journal of the American Society of Clinical Oncology 2004; 22(12): 2356-47.

43. Escudier B, Pluzanska A, Koralewski P, et al. Bevacizumab plus interferon alfa-2a for treatment of metastatic renal cell carcinoma: a randomised, double-blind phase III trial. Lancet 2007; 370(9605): 2103-11.

44. Rini BI, Wilding G, Hades G, et al. Phase II study of axitinib in sorafenib-refractory metastatic renal cell carcinoma. Journal of clinical oncology: official journal of the American Society of Clinical Oncology 2009; 27(27): 4462-8.

45. Motzer RJ, Escudier B, Tomczak P, et al. Axitinib versus sorafenib as second-line treatment for advanced renal cell carcinoma: overall survival analysis and updated results from a randomised phase 3 trial. The Lancet Oncology 2013; 14(6): 552-62.

46. Sun Y, Rha S, Lee SH, et al. Phase II study of the safety and efficacy of temsirolimus in East Asian patients with advanced renal cell carcinoma. Japanese journal of clinical oncology 2012; 42(9): 836-44.

47. Hainsworth JD, Spigel DR, Burris HA, 3rd, Waterhouse D, Clark BL, Whorf R. Phase II trial of bevacizumab and everolimus in patients with advanced renal cell carcinoma. Journal of clinical oncology: official journal of the American Society of Clinical Oncology 2010; 28(13): 2131-6.

48. Motzer RJ, Escudier B, Oudard S, et al. Phase 3 trial of everolimus for metastatic renal cell carcinoma: final results and analysis of prognostic factors. Cancer 2010; 116(18): 4256-65.

49. Motzer RJ, Escudier B, McDermott DF, et al. Nivolumab versus Everolimus in Advanced Renal-Cell Carcinoma. The New England journal of medicine 2015; 373(19): 1803-13.

50. Hades G, Carducci M, Tomczak P, et al. Temsirolimus, interferon alfa, or both for advanced renal-cell carcinoma. The New England journal of medicine 2007; 356(22): 2271-81.

51. Motzer RJ, Escudier B, Oudard S, et al. Efficacy of everolimus in advanced renal cell carcinoma: a double-blind, randomised, placebo-controlled phase III trial. Lancet 2008; 372(9637): 449-56.

52. Martin Henner Voss YC, Joshua Chaim, Devyn Taylor Coskey, Katlin Woo, Sujata Patil, Ana M. Molina, James Hsieh, Robert J. Motzer, Darren Richard Feldman. A phase II trial of everolimus and bevacizumab in advanced non-clear cell renal cell cancer. 2015 genitourinary cancers symposium: jco; 2015.

53. Rosenberg SA, Mule JJ, Spiess PJ, Reichert CM, Schwarz SL. Regression of established pulmonary metastases and subcutaneous tumor mediated by the systemic administration of high-dose recombinant interleukin 2. The Journal of experimental medicine 1985; 161(5): 1169-88.

54. Rosenberg SA. IL-2: the first effective immunotherapy for human cancer. Journal of immunology 2014; 192(12): 5451-8.

55. Fyfe G, Fisher RJ, Rosenberg SA, Sznol M, Parkinson DR, Louie AC. Results of treatment of 255 patients with metastatic renal cell carcinoma who received high-dose recombinant interleukin-2 therapy. Journal of clinical oncology: official journal of the American Society of Clinical Oncology 1995; 13(3): 688-96.

56. Rosenberg SA, Yang JC, Topalian SL, et al. Treatment of 283 consecutive patients with metastatic melanoma or renal cell cancer us-
72. Naomi B. Haas JM, Robert G. Uzzo, Michael B. Atkins, George Wilding, Michael Pins, Michael A. S. Jewett, Christopher J. Kane, David Cell, Lynne I. Wagner, Bob Coomes, Christopher G. Wood, Janice P. Dutcher, Keith Flaherty, Robert S. DiPaola; Dose analysis of ASSURE (E2805): Adjuvant sorafenib or sunitinib for unfavorable renal carcinoma, an ECOG-ACRIN-led, NCTN phase 3 trial. ASCO Annual Meeting; 2015; Chicago: JCO; 2015.

73. Jennifer J. Knox CHB, Tae Min Kim, Thomas Cosgriff, Vickie Srimuninnimit, Kenneth B. Pittman, Roberto Sabbatini, Sun Young Rha, Thomas W. Flagg, Ray D. Page, J. Thaddeus Beck, Foon yia Cheung, Sunil Yadav, Poulam M. Patel, Lionel Geoffrois, Edward Schiff, Julie Niolat, Noah Berkowitz, Maurizio Vo, Robert Motzer,. Final overall survival analysis for the RECORD-3 study of first-line everolimus followed by sunitinib versus first-line sunitinib followed by everolimus in metastatic RCC (mRCC). ASCO Annual Meeting 2015; 2015; Chicago: J Clin Oncol 33, 2015 (suppl; abstr 4554); 2015.

74. Shiven B. Patel NA, JoAnn Hsu, Srinivas Kiran Tantravahi, David Gill, Winston Vuong, Julia A. Batten, David D. Stenehjem, Sumanta Kumar Pal. Everolimus (EVE) versus temsirolimus (TEM) after first-line treatment with VEGF TKI in patients with metastatic renal cell carcinoma. ASCO GU cancers symposium; 2015; Orlando: JCO; 2015.

75. Iacovelli R, Santoni M, Verzoni E, et al. Everolimus and temsirolimus are not the same second-line in metastatic renal cell cancer. A systematic review and meta-analysis of literature data. Clinical genitourinary cancer 2015; 13(2): 137-41.

76. Dubinski W, Gabril M, Iakovlev VV, et al. Assessment of the prognostic significance of endoglin (CD105) in clear cell renal cell carcinoma using automated image analysis. Human pathology 2012; 43(7): 1037-43.

77. Ardelean DS, Jerzic M, Yin M, et al. Endoglin and activin receptor-like kinase 1 heterozygous mice have a distinct pulmonary and hepatic angiogenic profile and response to anti-VEGF treatment. Angiogenesis 2014; 17(1): 129-46.