current status of targeted therapy for advanced renal cell carcinoma

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the treatment of metastatic renal cell carcinoma (mRCC) has recently evolved from being predominantly cytokine-based treatment to the use of targeted agents, which include sorafenib, sunitinib, bevacizumab (plus interferon alpha [IFN-α]), temsirolimus, everolimus, pazopanib, and most recently, axitinib. Improved understanding of the molecular pathways implicated in the pathogenesis of RCC has led to the development of specific targeted therapies for treating the disease. in Korea, it has been 5 years since targeted therapy became available for mRCC. Thus, we now have broader and better therapeutic options at hand, leading to a significantly improved prognosis for patients with mRCC. however, the treatment of mRCC remains a challenge and a major health problem. Many questions remain on the efficacy of combination treatments and on the best methods for achieving complete remission. Additional studies are needed to optimize the use of these agents by identifying those patients who would most benefit and by elucidating the best means of delivering these agents, either in combination or as sequential single agents. Furthermore, numerous ongoing research activities aim at improving the benefits of the new compounds in the metastatic situation or their application in the early phase of the disease. This review introduces what is currently known regarding the fundamental biology that underlies clear cell RCC, summarizes the clinical evidence supporting the benefits of targeted agents in mRCC treatment, discusses survival endpoints used in pivotal clinical trials, and outlines future research directions.

key words: Molecular targeted therapy; mTOR protein; Renal cell carcinoma; Vascular endothelial growth factor A

INTRODUCTION

The resistance of renal cell carcinoma (RCC) to the traditional cytotoxic chemotherapy is now well established. For many years, the mainstream therapy modality of metastatic renal cell carcinoma (mRCC) was based on cytokine-mediated approaches using either interferon alpha (IFN-α) or interleukin-2 (IL-2) or both [1]. The results with these agents were less than satisfactory because they produced objective response rates on the order of only 10 to 20%, with long-term durable responses in 5 to 7% of cases, at least for high-dose IL-2 [2,3]. Since the availability of information regarding the aberrant activities of signal transduction pathways in RCC, specific molecular targets for potential therapies have been identified and analyzed pharmacologically in a variety of in vitro and preclinical studies. As a result, the treatment of mRCC has dramatically changed in recent years. This has been driven by two groups of targeted agents: namely, vascular endothelial growth factor (VEGF)-targeted therapies and mammalian target of rapamycin (mTOR) inhibitors [4]. Since 2005, seven targeted agents have been approved by regulatory authorities in the United States (US) and Europe (Axitinib is newly approved by the US Food and Drug Administration) for various uses in advanced RCC or mRCC patients. However, despite these advancements in treatment modalities, there

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Article History:
received 22 February, 2012
accepted 15 March, 2012

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Korean Journal of Urology
© the Korean Urological Association, 2012 217
Korean J Urol 2012;53:217-228
are many limitations when it comes to the treatment of mRCC.

The objectives of this article were to review the clinical evidence supporting the benefits of these agents, introduce the treatment guidelines, and identify limitations. Furthermore, future research directions with these targeted therapies are discussed.

EPIDEMIOLOGY OF RCC

RCC is the most common renal tumor and accounts for 3% of all adult cancers [5]. The incidence and mortality of renal malignancies have been on the rise worldwide over the past more than 30 years [6], particularly in the Western world, where kidney cancer has been among the highest of the tumors with an upward trend in incidence [6,7]. According to the recent report of the Korea Central Cancer Registry, RCC accounted for 1.8% of Korean cancers in 2008 [8]. Since then, the incidence of RCC in Korea has shown a steady increase.

Rising incidence rates are partly attributable to improvements in diagnostic imaging, but better detection does not explain the continued high number of advanced tumors and the increase in tumor size-specific mortality among RCC patients [7]. Surgery remains the mainstay therapy modality for those who present with clinically localized tumors and is an effective cure for the majority of patients. However, at least one-third of patients are diagnosed with metastases and an additional 20 to 40% of patients develop metastases after radical nephrectomy with curative intent [9-11]. The outcome of patients with mRCC is poor. The 5-year survival rate of RCC patients with metastatic lesions is 0 to 20% [12-14] and is 25 to 50% even if the metastatic lesion is solitary and can be completely resected [15-17].

IMMUNOTHERAPY FOR mRCC

Previously, systemic treatment was limited to cytokine therapy with IL-2 or IFN-α, because mRCC is largely resistant to chemotherapy [1]. Cytokine therapy is based on the rationale that stimulation of the immune system kills cancer cells. However, this modality in patients with mRCC is associated with low rates of response yet high rates of toxicity in the first-line setting [1]. In the second-line setting (in patients who have progressed on one cytokine), even fewer responses are observed, and toxicity remains similar to that of the first-line use [18]. In addition, patients’ median survival period is only about 13 months [19-21].

However, high-dose IL-2 remains the only agent with proven efficacy in producing complete and partial responses in patients with mRCC [21-23]. Furthermore, despite the use of a single-agent interferon, which has decreased significantly since the introduction of targeted therapy, it remains in the frontline setting in combination with bevacizumab as a result of 2 large phase III trials [24,25]. Lastly, improved understanding of immune regulation has led to the advancement of targeted immunotherapy using immune checkpoint inhibitors that have shown promising activity and that are moving forward in clinical development [26].

MECHANISMS OF TARGETED THERAPY IN mRCC

There are at least 5 histological forms of RCC. The most prevalent is the clear cell type, which accounts for 75% of cases [27,28]. Clear cell RCC is characterized by inactivation of a crucial tumor-suppressor gene, known as von Hippel Lindau (VHL) [29,30]. Understanding the biological processes that underlie the clear cell type RCC, in particular, the central role played by the VHL-hypoxia-inducible factor (HIF)-VEGF axis, is important. This is because the various members of this cascade are the therapeutic targets for most of the agents currently used in the management of clear cell type RCC. The concept of targeting these specific signaling molecules is the fundamental underpinning of the so-called “targeted therapies.” This principle has resulted in two fundamental ideologies that are nevertheless interrelated. These two principles underlie the categories of targeted therapeutics, i.e., those that block the VEGF pathway and those that block the mTOR pathway.

The typical mechanism of targeted therapy is shown in Fig. 1 [28]. VHL encodes the VHL protein. If VHL is inactivated, a defective VHL protein is produced and HIF is not degraded. Activated HIF then translocates into the nucleus and leads to the transcription of a wide repertoire of genes, including VEGF, platelet-derived growth factor (PDGF) [31], and transforming growth factor alpha [32], which have a central role in tumor angiogenesis and progression. Activation of the mTOR pathway also increases HIF levels [33]. This leads to increased transcription of genes, such as VEGF and PDGF, that control cell proliferation, glucose uptake, and angiogenesis [33]. Thus, increased HIF expression can promote angiogenesis in tumors.

EFFICACY OF TARGETED THERAPY IN mRCC

The targeted agents approved in the United States for the treatment of mRCC are sorafenib, sunitinib, bevacizumab (in combination with IFN-α), temsirolimus, everolimus, pazopanib, and, recently, axitinib. Table 1 summarizes the key efficacy data from the pivotal trials for these agents. Before the advent of targeted therapies, patients with mRCC treated with cytokines showed a median survival of 10 months [34]. Sunitinib, bevacizumab, and temsirolimus were compared with IFN-α in treatment-naïve patients and were found to be superior. In addition, sorafenib, everolimus, and pazopanib were shown to be superior to placebo in their defining trials, although the intent of many of these latter protocols was to focus on the patients who had already failed a cytokine or anti-VEGF therapy [24,25].

Korean J Urol 2012;53:217-228
FIG. 1. Biological pathways and therapeutic targeted agents for renal cell carcinoma. Reprinted from Rini BI, Atkins MB, Lancet Oncol 2009;10:992-1000, with permission of Elsevier [28].

TABLE 1. Targeted agents for metastatic renal cell cancer (approved): key results of phase III study

| Agent                        | Setting          | No. | ORR (%) | DCR\(^a\) (%) | CR + PR + SD | Median PFS (mo) | Median OS (mo) |
|------------------------------|------------------|-----|---------|---------------|--------------|----------------|---------------|
| Sunitinib vs. IFN-α [43]     | 1st-line         | 750 | 31 vs. 6\(^c\) | 0 + 31 + 48 vs. 0 + 6 + 49 | 11.0 vs. 5.1\(^c\) | 26.4 vs. 21.8 |
| Temsirolimus vs. IFN-α [41]  | 1st-line         | 626 | 8.6 vs. 4.8 | 32.1 vs. 15.5 | 3.8 vs. 1.9\(^c\) | 10.9 vs. 7.3c |
| Bevacizumab + IFN-α vs.      | Poor risk        | 649 | 31 vs. 13\(^c\) | 1 + 30 + 48 vs. 2 + 11 + 50 | 10.2 vs. 5.4\(^c\) | 23.3 vs. 21.3 |
| placebo + IFN-α (AVOREN) [24]| 1st-line         | 732 | 26 vs. 13\(^c\) | -              | 8.5 vs. 5.2\(^c\) | 18.3 vs. 17.4 |
| Bevacizumab + IFN-α vs.      | 1st-line         |     |         |               |              |                |               |
| IFN-α (CALBG 90206) [25]    |                  |     |         |               |              |                |               |
| Pazopanib vs. placebo [39]   | 1st-line         | 233 | 32 vs. 4\(^c\) | < 1 + 30 + 38 vs. 0 + 3 + 41 | 11.1 vs. 2.8\(^c\) | 22.9 vs. 20.5 |
| Sorafenib vs. placebo [42]   | 2nd-line         | 202 | 29 vs. 3\(^c\) | < 1 + 10 + 74 vs. 0 + 2 + 53 | 7.4 vs. 4.2\(^c\) | 17.8 vs. 15.2 |
| Everolimus vs. placebo [36]  | 2nd-line         | 903 | 10 vs. 2\(^c\) | < 1 + 10 + 74 vs. 0 + 2 + 53 | 5.5 vs. 2.8\(^c\) | 17.8 vs. 14.5\(^c\) |
| Axitinib vs. sorafenib [40]  | 2nd-line         | 416 | 1.8 vs. 0 | 0 + 27 + 67 vs. 0 + 0 + 32 | 4.9 vs. 1.9\(^c\) | 14.8 vs. 14.4 |

**ORR**, objective response rate; **SD**, stable disease; **PFS**, progression-free survival; **OS**, overall survival; **IFN-α**, interferon alpha; **AVOREN**, Avastin and Roferon in Renal Cell Carcinoma; **CALBG**, Cancer and Leukemia Group B; **NA**, not available.

\(^a\)**DCR**(disease control rate), **CR**(complete response) + **PR**(partial response) + **SD**(stable disease) ≥ 3 mo, **CB**, clinical benefit; **OR**, objective response (CR + PR), **Statistically significant.**

In a recent study, as a second-line use, axitinib was found to be superior when compared with sorafenib [40]. An improvement in overall survival (OS) provides the most convincing evidence that a new therapy is superior to the existing standard modality. Only temsirolimus (for the patient with “poor-risk” RCC) had led to improved OS in randomized phase III trials [41]. However, other agents, including sunitinib, sorafenib, and pazopanib, are clearly active in RCC and constitute the comparator arm in several ongoing studies. Owing to the crossover of the patients and
TABLE 2. Results of targeted therapy for metastatic renal cell cancer (RCC) in Korea

| Agent | Total no. of patients | Median age (yr) | Clear cell RCC | Patient with lung meta | MSKCC 0 (favorable) | MSKCC 1,2 (intermediate) | MSKCC ≥ 3 (poor) | Median PFS (mo) | Median OS (mo) | Objective response | Complete response | Partial response | Stable disease | Disease control rate |
|-------|----------------------|----------------|---------------|----------------------|-------------------|----------------------|-------------------|----------------|----------------|------------------|-----------------|-----------------|---------------|-------------------|
| Sunitinib | 375 | 62 | 375 (100) | 292 (78) | 143 (38) | 290 (56) | 23 (6) | 11 (7.2) | 26.4 | 137 (37) | 1 (1) | 136 (6) | 176 (47) | 313 (81) |
| Sunitinib | 76 | 57.5 | 65 (85.5) | 56 (73.7) | 7 (11.6) | 47 (78.3) | 6 (10) | 7.2 | 22.8 | 21 (27.6) | 1 (1.3) | 20 (26.3) | 43 (56.6) | 64 (84.2) |
| Sunitinib | 65 | 58 | 55 (85) | 48 (74) | 14 (21) | 20 (31) | 28 (43) | 11.8 | 22.8 | 23 (43) | 0 (0) | 23 (43) | 23 (43) | 46 (86) |
| Sunitinib | 21 | 63.9 | 21 (100) | 15 (71.4) | 4 (19.0) | 10 (47.6) | 7 (33.3) | 13.4 | 28.1 | 11 (52.4) | 1 (4.8) | 10 (47.6) | 7 (33.3) | 18 (85.7) |
| Sunitinib | 132 | 57.0 | 109 (82.6) | 103 (78.0) | 35 (30.4) | 61 (53.0) | 19 (16.5) | 8.2 | 23.1 | 45 (34.1) | 2 (1.5) | 43 (32.6) | 59 (44.7) | 104 (80.3) |
| Bevacizumab | 88 | 56 | 71 (81.0) | 65 (74) | - | - | - | 5.0 | 31.3 |
| Pazopanib | 46 | 59 | 37 (80.4) | - | - | - | - | - |
| Temsirolimus | 88 | 56 | 14 (21) | 18 (21) | 1 (1) | 1 (1) | 28 (43) | 14 (30.4) |
| Sorafenib | 14 (30.4) | 35.0 |
| Everolimus | 31.3 | 17 (40.0) |

Values are presented as number (%).

MSKCC, Memorial Sloan-Kettering Cancer Center risk group; PFS, progression-free survival; OS, overall survival.

TABLE 3. Evidence-based clinical guidelines for systemic therapy for metastatic renal cell carcinoma

| Setting | MSKCC risk or prior treatment | Histology | Recommended (option) |
|---------|------------------------------|-----------|----------------------|
|         |                              |           | NCCN\textsuperscript{a} | EAU\textsuperscript{b} |
| First-line | Good or intermediate risk | Clear cell | Sunitinib | Sunitinib |
|          |                              | Non-clear cell | Bevacizumab + IFN-α | Bevacizumab + IFN-α |
|          |                              |           | Pazopanib (HD IL-2) (sorafenib) | Pazopanib (HD IL-2) |
|          |                              |           | Clinical trial (sorafenib) (sunitinib) (temsirolimus) | - |
| Poor risk |                              | Clear cell | Temsirolimus | Temsirolimus |
|          |                              | Non-clear cell | Temsirolimus | Temsirolimus |
| Second-line | Prior cytokine | Clear cell | Sorafenib | Sorafenib |
|          |                              | Non-clear cell | Sunitinib | Sunitinib |
|          |                              |           | Pazopanib (temsirolimus) (bevacizumab) | Pazopanib (sunitinib) (temsirolimus) |
| Prior VEGFRI |                              |           | Everolimus (sorafenib) (sunitinib) | Everolimus |
| Prior mTORi |                              |           | Clinical trial | Clinical trial |

MSKCC, Memorial Sloan-Kettering Cancer Center; NCCN, National Comprehensive Cancer Network; EAU, European Association of Urology; IFN-α, interferon alpha; HD, high-dose; IL-2, interleukin-2; VEGFRI, vascular endothelial growth factor receptor inhibitor; mTORi, mammalian target of rapamycin inhibitor.

\textsuperscript{a}: NCCN: Kidney Cancer, NCCN 2012 Clinical Practice Guidelines in Oncology V.1. 2012, Cat 1 & 2A, \textsuperscript{b}: EAU: Guideline on Renal Cell Carcinoma 2010, Grade A.

a widespread use of active agents in patients who progress on their assigned therapy, the OS analysis in randomized trials may be confounded. In a recent update, with censoring of crossover data, the median overall survival with sunitinib and sorafenib was significantly longer than for their respective control groups [42,43]. Therefore, the OS benefit of these drugs seems to be relatively clear.

Currently, several retrospective studies and one prospective trial using tyrosine kinase inhibitors (TKIs) have been published in Korea (Table 2). According to these studies, TKIs, especially sunitinib, seem to be efficacious in a manner similar to or even better than in Western patients with slightly different treatment-related adverse events, such as a high incidence of hematological toxicities [44].
CURRENT GUIDELINES FOR THE TREATMENT OF mRCC

The treatment guidelines for mRCC are experiencing a rapid evolution to incorporate the new molecular-targeted therapies that have been approved recently by the US and European regulatory authorities. Table 3 summarizes the current updated recommendations contained in the National Comprehensive Cancer Network (NCCN) and European Association of Urology (EAU) guidelines.

In terms of clear cell mRCC, the NCCN guidelines (ver. 1.2012) recommend a number of options for first- and second-line treatment of mRCC. For first-line therapy, the recommended modalities for patients with a good or intermediate prognosis are sunitinib, bevacizumab plus IFN-α, and pazopanib. Temsirolimus has a recommendation for the treatment of patients with poor-prognosis mRCC. In addition, the NCCN also suggests alternatives for selected patients in the first-line setting, such as high-dose IL-2 or sorafenib, temsirolimus, and enrollment in a clinical trial.

For second-line therapy, everolimus is the only agent to have an NCCN category 1 recommendation for the treatment of patients who have failed TKI treatment. Sorafenib, sunitinib, and pazopanib are recommended for the treatment of patients after cytokine failure. For the treatment of patients after the failure of other TKIs, sorafenib, sunitinib, and pazopanib are also recommended as an alternative treatment (category 2B or 3). Temsirolimus is also considered as the treatment for patients after cytokine failure (category 2A) in addition to the treatment for patients after TKI failure (category 2B). The EAU guidelines also recommend treatment options for patients with predominantly non-clear cell RCC. As a first-line therapy, sunitinib, bevacizumab plus IFN-α, and pazopanib are recommended for the treatment of low- and intermediate-risk patients. For the treatment of high-risk patients, temsirolimus is recommended. IFN-α monotherapy is no longer recommended. However, high-dose bolus IL-2 is recommended as a first-line treatment for mRCC in patients with clear cell histology and good prognostic factors. As a second-line therapy, everolimus has a recommendation for the treatment of patients after TKI failure; sorafenib and pazopanib are each recommended for the treatment of patients after cytokine failure; and enrollment in clinical trials is recommended for the treatment of patients after mTOR inhibitor failure.

For patients with non-clear cell mRCC, enrollment in clinical trials of systemic therapy has been the preferred strategy. However, the 2012 NCCN guidelines provide first-line therapy recommendations for patients with non-clear cell mRCC. Temsirolimus has a category 1 recommendation for the treatment of patients with poor risk non-clear cell mRCC.

In Korea, sunitinib and sorafenib have been used as a first-line treatment for patients with clear cell mRCC under the support of national health insurance since 2007. Furthermore, pazopanib has also been used since May 2011. For the treatment of non-clear cell mRCC or clear cell mRCC, with poor risk group patients, national health insurance has reimbursed the use of temsirolimus since June 2011. Most recently, everolimus was approved by the multidisciplinary committee for patients with sunitinib or sorafenib or pazopanib failure under insurance coverage. Table 4 shows the coverage of national health insurance in Korea for patients with mRCC.

DRUG RESISTANCE AND TREATMENT STRATEGIES

Some patients are inherently resistant to these targeted agents. In addition, most, if not all, patients acquire resistance over time. In general, the development of resistance has been observed after a median of 6 to 11 months of treatment [28]. Complete or durable responses have only rarely been noted, necessitating chronic therapy for most patients. Several strategies emerge from the above considerations regarding the mechanisms of resistance to the currently available therapeutics in mRCC. The first involves the use of an agent that blocks a resistance mechanism. Such an agent could be used as a monotherapy at the time of resistance to targeted therapy, could be added to continued VEGF blockade at the time of resistance, or could be used as a first-line treatment for patients with clear cell mRCC under the support of national health insurance since 2007. Furthermore, pazopanib has also been used since May 2011. For the treatment of non-clear cell mRCC or clear cell mRCC, with poor risk group patients, national health insurance has reimbursed the use of temsirolimus since June 2011. Most recently, everolimus was approved by the multidisciplinary committee for patients with sunitinib or sorafenib or pazopanib failure under insurance coverage. Table 4 shows the coverage of national health insurance in Korea for patients with mRCC.

| No. | Systemic agent              | Treatment target       | Treatment setting          |
|-----|-----------------------------|------------------------|----------------------------|
| 1   | Aldesleukin (IL-2)          | Stage 4 RCC            | First line or more         |
| 2   | Aldesleukin + IFN-α         | Stage 4 RCC            | First line or more         |
| 3   | Aldesleukin + IFN-α + 5FU   | Stage 4 RCC            | First line or more         |
| 4   | Sunitinib                   | Metastatic or recurrent| First line or more         |
| 5   | Sorafenib                   | Metastatic or recurrent| First line or more         |
| 6   | Pazopanib                   | Metastatic or recurrent| First line or more         |
| 7   | Temsirolimus*               | Metastatic or recurrent| First line                 |
| 8   | Everolimus                  | Metastatic or recurrent| Second line or more        |

*Only for non-clear cell carcinoma or poor risk clear cell carcinoma.

RCC, renal cell carcinoma; IL-2, interleukin-2; IFN-α, interferon alpha; 5FU, 5-fluorouracil.

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be added initially in combination with VEGF blockade with the hope of delaying the onset of resistance. Alternatively, more effective blockade of the initial target could involve increasing the dose of an existing agent or measuring the drug concentrations to ensure an adequate dosing, or alternatively, switching to an agent that more potently inhibits the target.

The current strategies to maximize the effectiveness of treatment are as follows.

1. Sequential therapy

Targeting different pathways may offer benefit in terms of overcoming resistance to individual agents. Sequential therapy has the potential to change mRCC into a chronic disease that can be managed for a long term through the administration of targeted agents in sequence. Although clinicians are currently using targeted agents in a sequential manner for patients with mRCC in practice, concerns remain regarding cross-resistance between the different agents. Thus, there are many questions regarding the optimal sequence for obtaining maximum clinical benefit from the available targeted therapies. In addition, adequate management of treatment-related toxicity can allow patients to remain on treatment for long periods and can help to maximize the clinical benefit of targeted agents. Recently, strategies to manage treatment-related adverse events are being refined [45].

The first randomized phase III study to investigate sequential targeted therapy in mRCC showed clinical efficacy for the sequence of sunitinib or sorafenib, followed by everolimus (RECORD-1) [36,46]. In this study, patients who had failed an earlier anti-VEGF therapy, 71% of whom had received sunitinib previously, were treated with either everolimus (RECORD-1) [36,46]. In this study, patients who had failed an earlier anti-VEGF therapy, 71% of whom had received sunitinib previously, were treated with either everolimus or placebo. The median progression-free survival (PFS) was 4.9 months vs. 1.9 months for those treated with sorafenib (p=0.011). Furthermore, the shorter median PFS was 4.8 months with axitinib and 3.4 months with sorafenib in patients who had progressed after previous treatment with sunitinib, bevacizumab+IFN-α, temsirolimus, or cytokines. The AXIS trial was these patients’ first exposure to a VEGFR-TKI, whereas 54% of patients had received previous sunitinib. In the subgroup of AXIS patients who had received previous sunitinib, the median PFS was 4.8 months with axitinib and 3.4 months with sorafenib (p=0.011). Furthermore, the shorter median PFS observed in both treatment arms in the sunitinib-refractory patients relative to those who received cytokines is suggestive of at least partial cross-resistance with sequential VEGF-targeted therapy [40]. Axitinib was shown to be more effective than sorafenib in patients who had progressed after previous treatment with sunitinib, bevacizumab+IFN-α, temsirolimus, or cytokines. The AXIS trial was these patients’ first exposure to a VEGFR-TKI, whereas 54% of patients had received previous sunitinib. In the subgroup of AXIS patients who had received previous sunitinib, the median PFS was 4.8 months with axitinib and 3.4 months with sorafenib (p=0.011). Furthermore, the shorter median PFS observed in both treatment arms in the sunitinib-refractory patients relative to those who received cytokines is suggestive of at least partial cross-resistance with sequential VEGF-targeted therapy. Table 5 summarizes the results of several prospective trials of sequential therapy with TKIs to TKI/mTOR inhibitors.

### Table 5. Prospective trials of sequential therapy (TKIs to TKI/mTOR inhibitors)

| Agent | Phase | Population | N  | ORR (%) | PFS (mo) | OS (mo) |
|-------|-------|------------|----|---------|----------|---------|
| Axitinib vs. sorafenib [40] | III | Sunitinib, bevacizumab + IFN, Temsirolimus, or cytokine | 717 | 19.4 vs. 9.4 | 6.7 vs. 4.7 | NA |
| Sunitinib [51] | II | Bevacizumab-refractory | 62 | 23 | 7.1 | 11.8 |
| Axitinib [80] | II | Sorafenib-refractory | 62 | 22.6 | 7.4 | 13.6 |
| Axitinib [81] | II | Sunitinib-refractory | 14 | 7 | 7.1 | 11.5 |
| Sorafenib [82] | II | Bevacizumab or sunitinib-R | 31 | 3 | 308 | NA |
| Sorafenib [49] | II | Sunitinib-refractory | 52 | 9.6 | 4 | 8 |
| ABT-869 [83] | II | Sunitinib-refractory | 53 | 9 | 5.4 | NA |
| mTOR inhibitors | III | TKI-refractory | 416 | 2 vs. 0 | 4.9 vs. 1.9 | 14.8 vs. 14.43 |
| Everolimus vs. placebo [36] | II | TKI-refractory | 26 | 0 | 6.5+ | 16.3+ |
| Everolimus [84] | II | TKI-refractory | 37 | 18 | 5.3 | NA |
| Temsirolimus + bevacizumab [85] | II | TKI-refractory | 30 | 23 | 7.1 | NA |

TKIs, tyrosine kinase inhibitors; mTOR, mammalian target of rapamycin; ORR, objective response rate; PFS, progression-free survival; OS, overall survival; INF, interferon; NA, not available.
Recently, several studies evaluating the efficacy of a second VEGFR-TKI, following a VEGFR-TKI→mTOR inhibitor treatment sequence, have been reported with encouraging results (Table 6). These results suggest that re-introduction of a VEGFR-TKI following progression on a VEGFR-TKI→mTOR inhibitor treatment sequence is an effective strategy. However, patients appear to derive a lesser degree of clinical benefit from VEGFR-TKI repeat challenge than that obtained in the first-line setting, which suggests at least partial cross-resistance. Interestingly, transient resistance to the same agent has also been observed. In a recent retrospective review of 23 patients, repeat challenge with sunitinib in patients with disease progression on sunitinib and other therapies resulted in 5 patients (22%) achieving a partial response and 17 patients (74%) achieving stable disease [52]. Repeat challenge was associated with a median PFS of 7.2 months compared with 13.7 months on the initial treatment (p=0.04). The results described here indicate the potential for re-treating with an agent, despite the occurrence of resistance from the first treatment, and have implications for achieving a continuum of treatment in these patients. So far, no therapies are approved for the third-line treatment of mRCC. In clinical practice, however, a strategy that is growing is the re-introduction of a VEGFR-TKI following the progression on a VEGFR-TKI and an mTOR inhibitor.

### 2. Combination therapy

The effectiveness of combination therapies is much more promising than that of single targeted therapies. It is hoped that combination therapies may induce better responses because they interfere with sequential steps in a single pathway or attack a tumor from two sides. However, for any potential clinical benefit, there must be a balance between the potential increases in toxicity associated with combining therapeutic agents. Because most combination data to date are preliminary, no combination can be said to fulfill this requirement. To determine clinical applicability, further studies are required of targeted agent combinations, if any. It is also important to consider the therapeutic options that are possible or available following the use of combination-targeted agent therapy [53].

#### 3. Adjuvant therapy

Adjuvant therapy has been used in the treatment of several malignancies with favorable results. To date, however, no medical therapies have been shown to improve outcomes in RCC when used in the adjuvant setting [54-56]. In the field of targeted therapy, the S-TRAC, ASSURE, and SORCE trials are ongoing for patients at high risk of recurrence (Table 7). The results of these trials are eagerly awaited to determine the role of targeted therapy in the adjuvant setting.

#### 4. Neoadjuvant (presurgical) therapy

Nowadays, we have yet to understand how targeted agents can be integrated with surgical approaches to maximize clinical benefit [57-60]. Cytokines do not affect the primary tumor. However, targeted therapies have an effect on the primary tumor, even in the absence of metastatic disease.

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**Table 6. Efficacy of treatment with a third-line TKI following treatment with sequential VEGFR-TKI→TOR inhibitor therapy**

| Sequence | Design | No. | Median PFS 1st line TKIs (mo) | Median PFS 2nd line mTORi (mo) | Median PFS 3rd line TKIs (mo) | ORR n (%) | SD n (%) | DCR n (%) | OS (mo) |
|----------|--------|-----|-----------------------------|-----------------------------|-----------------------------|-----------|--------|----------|--------|
| TKIs→Eve→TKIs | Retro | 40 | 11.3 | 5.9 | 5.5^a (0.4-22.3) | 4 (10) | 22^b (55) | 26 (65) | 11.3^c (0.8-22.3) |
| Su→Various→Su | Retro | 26 | 13.7 | 26 | 7.2^d (1.2-28.5+) | 5 (21) | 17 (71) | 22 (92) | NA |
| Su→mTORi→So | Retro | 34 | 10 | 4 (Eve)/2 (Tem) | 4 (3-6) | 8^e (23.5) | 7 (20.5) | 15 (44) | 7^f (6-10) |
| TKIs→Eve→TKIs | Retro | 36 | 11.4 | 8.9 | 8.2 | 3 (8.6) | 24 (68.6) | 27 (77.1) | 29.1^g |
| So→Eve→Su | Retro | 14 | 11.7 | 5.1 | 9.1 | 21.9 |
| Su→Eve→So | Pros | 26 | 14.4 | 4.3 | 3.9 | 22.8 |
| TKIs→Eve→Dov | Pros | 31 | 5.5 | 2 (3.4) | 29 (49.2)^f | 11.8^f |

TKI, tyrosine kinase inhibitor; VEGFR, vascular endothelial growth factor receptor; mTORi, mammalian target of rapamycin inhibitor; PFS, progression-free survival; ORR, objective response rate; SD, stable disease; DCR, disease control rate; OS, overall survival; Eve, everolimus; Tem, temsirolimus; Su, sunitinib; So, sorafenib; Dov, dovitinib; Retro, retrospective; Pros, prospective.

^a: SD ≥3 mo, DCR: PR+SD. ^b: PFS: treatment of sorafenib +/-, PFS after everolimus 3.7 vs. 11.3 mo, p=0.036 in uni-variate analyses, ^c: OS: from start of third-line TKIs, OS: PFS first-line VEGF treatment ≥6 mo vs. <6 mo, OS 53.4 vs.19.3 mo, p=0.002. ^d: Median PFS: sunitinib rechallenge interval ≥6 mo vs. ≤6 mo, 16.5 vs. 6.5 mo, p=0.03. ^e: Response rate: first-line sunitinib responder vs. non-responder, 47% vs. 0%, p=0.0027. ^f: OS: everolimus→sunitinib vs. everolimus→sorafenib=30.5 mo vs. 17.6 mo (p=0.102). ^g: OS: SD≥2 mo 29 (49.2%), SD>4 mo 16 (27.1%). ^b: OS: from start of sorafenib treatment, ^f: OS: from start of dovitinib treatment.
TABLE 7. Ongoing adjuvant trials with targeted agents in renal cell carcinoma (RCC)

| Treatment | Clinical trials identifier | No. | Sponsor | Start | Projected completion |
|-----------|---------------------------|-----|---------|-------|----------------------|
| SORCE     | NCT00492258               | 1,656 | MRC/EORTC | June 2007 | August 2012 |
| ASSURE    | NCT00326898               | 1,923 | ECOG    | May 2006 | April 2016 |
| S-TRAC    | NCT00375674               | 600   | Pfizer  | July 2007 | June 2017 |
| EVEREST   | NCT01120249               | 1,218 | SWOG    | February 2010 | August 2013 |
| PROTECT   | VEG113387                 | 1,500 | GSK     | November 2010 | December 2012 |

SORCE, A phase III randomised double-blind study comparing Sorafenib with placebo in patients with Resected primary renal cell carcinoma at high or intermediate risk of relapse; RCC, renal cell carcinoma; ASSURE, Adjuvant Sunitinib versus Sorafenib versus Placebo in Patients with Resected Renal Cell Carcinoma; S-TRAC, Sunitinib Treatment of Renal Adjuvant Cancer; EVEREST, EVErolimus for Renal Cancer Ensuing Surgical Therapy, A phase III study; PROTECT, Patient Related Outcomes with Endeavor versus Cypher Standing Trial; MRC, Medical Research Council; EORTC, European Organization for Research and Treatment of Cancer; ECOG, Eastern Cooperative Oncology Group; SWOG, Southwest Oncology Group; GSK, Glaxo Smith Kline.

In locally advanced non-mRCC, Karakiewicz et al. [61] reported an atrial tumor thrombus that was downstaged to the level of the infrahepatic vena cava after two cycles of sunitinib. However, another study showed that only 4 of 17 patients with advanced RCC taking sunitinib, without a previous nephrectomy, experienced partial responses in their primary tumors and 1 patient progressed [62]. Neoadjuvant targeted therapy can only downstage or improve the resectability of the primary tumor or associated lesions in 20 to 25% of patients [63]. To date, there are no established criteria for discriminating between patients who will and those who will not benefit from neoadjuvant setting.

Furthermore, it is not known which patients with mRCC might benefit from an upfront targeted therapy followed by cytoreductive nephrectomy [57,64]. To clarify these critical issues, several studies are ongoing. The CARMENA and EORTC trials will provide evidence-based information regarding the benefits and the timing of cytoreductive nephrectomy in the targeted therapy era. Also in Korea, the Korean Urological Oncology Society recently led a prospective trial of “neoadjuvant sunitinib treatment for metastatic clear cell RCC” (NCT01069770).

Although studies have demonstrated the general tolerability of targeted agents, data are still limited on the safety of surgical resection following treatment with these agents, and several studies have shown increased perioperative complications after treatment [65,66].

5. Targeted therapy for non-clear cell RCC

Temsirolimus represents the only targeted therapy for which superior efficacy was confirmed in advanced non-clear cell RCC in a prospective and randomized fashion. In addition, Hudes et al. [67] demonstrated better responses in patients with advanced non-clear cell RCC than in patients with clear cell disease in their temsirolimus data, which suggests that temsirolimus is an excellent first-line option in patients with non-clear cell mRCC. Despite the absence of prospective data for the efficacy of sunitinib or sorafenib, several investigators have demonstrated the efficacy of these two agents in patients with non-clear cell mRCC [68-70]. These data indicate that targeted therapies are clearly effective in patients with non-clear cell mRCC. Therefore, we need to look at the results of large prospective studies in the future.

Present and Future of Targeted Therapy in mRCC

All seven agents (sunitinib, sorafenib, temsirolimus, bevacizumab + IFN-α, pazopanib, everolimus, and axitinib) have shown efficacy and safety in phase III randomized controlled clinical trials. However, mRCC patients have to undergo chronic treatment, because targeted therapies rarely achieve durable and complete remissions. Drug resistance is the underlying reason for the growth and spread of tumors in the presence of systemic treatment. Further-
more, it remains the main barrier against long-term tumor control. In general, better tolerance of the targeted therapies than of chemotherapies is experienced by cancer patients. Nevertheless, the targeted agents still act as cytotoxic substances in a broader sense and can, therefore, cause several side effects that must not be neglected [71,72].

One therapy modality is not likely to benefit all patients, but rather a therapy modality is indicated on an individual, case-by-case basis. Treatment should be tailored to meet individual circumstances and needs, and achieving this is a considerable clinical challenge. Thus, many investigators emphasize the value of clinical judgment and experience to support treatment decisions for an individual [73]. Most patients today will receive several targeted therapies in a treatment sequence. Before starting first-line therapy, the whole potential therapeutic sequence should be considered for the individual patient.

An optimal balance between quality of life and prolongation of survival will only be achieved by considering both the benefits and the risks of the new targeted therapies [74]. We must overcome several treatment challenges to maximize the potential of targeted agents, and these include the identification of predictive molecular markers, drug resistance, the identification of the most effective sequence or combination of targeted agents, efficient clinical trial design, and the provision of cost-effective access to treatment for all patients with mRCC.

The development of targeted agents has substantially improved the prognosis of patients with mRCC. To further these advances, the following assignments are proposed for the future [53]. Identifying and optimizing the most appropriate sequence or combination of agents should be considered first. Second, overcoming drug resistance through newer agents or sequential or combination therapies should be addressed. Third, imaging techniques as part of the whole potential therapeutic sequence should be considered for the individual patient.

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**CONCLUSIONS**

Since the advent of targeted therapies, the modality of mRCC treatment has changed and the OS for these patients is now greater than 2 years in prospective studies. Currently, the first-line standard of care for patients with clear cell mRCC includes sunitinib, bevacizumab combination with IFN-α, pazopanib, and temsirolimus, all of which are options for patients with high-risk characteristics. Everolimus has proven efficacy as a second-line targeted therapy. The sequential use of targeted therapies can improve PFS and OS. Additional progression-free survival benefits might be derived from cytoreductive nephrectomy while we await for the results of ongoing phase III trials. However, we have much work to be completed. As more information regarding mechanisms of disease and drug resistance becomes available, new targets, new targeted agents, and new combinations will be studied with the goal of providing maximal efficacy with manageable toxicity.

**CONFLICTS OF INTEREST**
The authors have nothing to disclose.

**ACKNOWLEDGEMENTS**

This study was supported by a Korean National Cancer Center Grant, No. 1110560.

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