Evaluating the safety and efficacy of axitinib in the treatment of advanced renal cell carcinoma

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Abstract: Axitinib is a tyrosine kinase inhibitor of vascular endothelial growth factor receptor, platelet-derived growth factor receptor-α, and c-kit. Phase I studies demonstrated 5 mg twice daily as the recommended starting dose with notable effects seen in renal cell carcinoma, an observation confirmed in Phase II trials. The trial of comparative effectiveness of axitinib versus sorafenib in advanced renal cell carcinoma (AXIS) was an international randomized Phase III study designed for registration purposes, compared axitinib to sunitinib. This trial randomized 723 patients with metastatic kidney cancer to axitinib or sunitinib in the second-line setting and demonstrated a median progression-free survival of 6.7 months for axitinib versus 4.7 months for sorafenib (P<0.0001). Clinical benefit was detected regardless of prior therapy, but no overall survival benefit has been observed. Axitinib is well tolerated without a significant effect on quality of life. The most common grade 3 toxicities are hypertension (16%), diarrhea (11%), and fatigue (11%), with other notable side effects being anorexia, nausea, hand–foot syndrome, and rash. Patients who developed diastolic blood pressure >90 mmHg were noted to have significantly longer median overall survival and overall response rates when compared to normotensive patients. Therefore, the manufacturer recommends escalating the twice-daily dose to 7 mg and 10 mg, as tolerated, if there is no significant increase in blood pressure on treatment. Currently, axitinib is approved for use in the second-line setting for patients with metastatic renal cell carcinoma. Research is ongoing in other disease settings.

Keywords: axitinib, renal cell carcinoma, side effects, drug safety

Review of pharmacology, mode of action, pharmacokinetics of axitinib

Axitinib is a substituted indazole derivative that potently inhibits all known vascular endothelial growth factor (VEGF) receptors (VEGFRs) at subnanomolar concentrations, and is able to inhibit platelet-derived growth factor receptor (PDGFR)-α and c-kit at nanomolar concentrations. In vitro, axitinib inhibits the VEGF signaling axis by preventing autophosphorylation in endothelial cells (ECs) leading to the inhibition of EC growth, survival, and proliferation.

The dramatic effect of axitinib on ECs has been demonstrated in preclinical studies. When axitinib was applied to a spontaneous islet cell tumor model of RIP-Tag2 transgenic mice, endothelial fenestrations disappeared, blood flow decreased, and new vessel sprouting was significantly diminished within 24 hours. Existing vascular density was reduced by almost 80% after 7 days, and there was significantly less expression of VEGFR-2 and VEGFR-3. The remaining vessels demonstrated ghost-like basement membranes that persisted after a loss of ECs, consistent with...
other models of vascular regression. When the experiment was extended to 21 days, investigators observed a significant concordant reduction in tumor mass.\(^2\)

Axitinib was initially evaluated in vivo in mice containing human renal cell carcinoma (RCC), melanoma, and colorectal cancer xenografts, and it inhibited tumor growth in a dose-dependent fashion for all three histologies. The investigators observed that the tumors with the most dramatic effects had the greatest reduction in CD31 staining, a marker expressed on ECs, suggesting activity through a reduction in microvessel density.\(^3\)

Axitinib is considerably more potent than the other available VEGFR tyrosine kinase inhibitors (TKIs). The inhibitory concentration (IC\(_{50}\)) of axitinib is at least 100 times lower than that of sorafenib, sunitinib, or pazopanib for VEGFR1, VEGFR2, and VEGFR3,\(^3–6\) and at least as potent at inhibiting PDGFR-\(\alpha\), PDGFR-\(\beta\), and c-kit. Axitinib appears to be considerably more selective for the VEGFR1–3 targets than for PDGFR, c-kit, and FLT3. It has almost no affinity for FLT3 and RET, unlike sunitinib and sorafenib.\(^3–6\)

Based on these promising preclinical results, a multi-center Phase I study of axitinib was conducted, enrolling 36 patients with advanced solid tumors.\(^1\) Six of those enrolled had metastatic RCC. Axitinib was dosed orally twice daily at doses between 5 mg and 30 mg. Hypertension was the most common dose-limiting toxicity (eleven patients experienced grade 3–4 toxicity, of which most were manageable with antihypertensive medications), while stomatitis, increased liver enzymes, and seizures occurred in two patients in each category. Five patients died while in the study, of whom three patients were considered to have died from disease progression. The remaining two patients had lung adenocarcinoma and developed fatal hemoptysis – one had a large centrally located tumor that bled while on treatment; the second developed grade 1 hemoptysis and was taken off study and then, some weeks later, developed disease progression and infection complicated by grade 4 hemoptysis. Following these events, the protocol was amended to exclude patients with centrally located lung tumors, as well as patients with squamous cell carcinoma of the lung. No subsequent episodes of hemoptysis have been reported.

The maximum tolerated dose was determined to be 5 mg twice daily, and axitinib was rapidly absorbed, reaching peak plasma concentrations after 2–6 hours of administration. Steady state was reached in 15 days with a terminal plasma half-life of 2–5 hours.\(^1\) Later work determined that axitinib is predominantly protein bound (>99%), primarily to albumin, but also to \(\alpha\)-1-acid glycoprotein.\(^7\) Pharmacokinetics analysis revealed that median plasma exposure was 49% higher in fasting subjects. The oral dose of 5 mg twice daily in the fasting state was recommended for Phase II dosing. In the Phase I evaluation, two of the six patients with RCC had an objective partial response (PR), foreshadowing the drug’s activity in this disease.\(^1\)

**Efficacy studies**

Two Phase II studies followed, confirming the antineoplastic activity for axitinib in RCC. The first study enrolled 52 patients who had failed cytokine therapy.\(^8\) Patients were excluded if they had received other VEGF-targeted therapy or if they had uncontrolled hypertension. All patients but three had undergone a prior nephrectomy. Thirty patients (58%) had at least one Memorial Sloan Kettering Cancer Center (MSKCC) adverse risk factor.\(^9\) All patients were treated with axitinib 5 mg twice daily in the fasting state, and therapy continued until unacceptable toxicity or progression. The study’s overall response rate (ORR) was 44.2% (95% confidence interval [CI]: 30.5–58.7), with two complete responses (CRs) and 21 PRs. An additional 13 patients had stable disease (SD) for more than 24 weeks, and nine had SD for more than 8 weeks, translating into an overall clinical benefit rate of 87%. Most responding patients had clear cell RCC. One patient had nonclear cell histology (papillary) and was determined to have SD with treatment.

Secondary endpoints included duration of response, time to progression, survival, quality of life, and safety. For those with CR or PR, the median response duration was 23.0 months (95% CI: 20.9– not estimable; range: 4.2–29.8). The median time to progression for the study cohort was 15.7 months (95% CI: 8.4–23.4; range: 0.03–31.5) and median overall survival (OS) was 29.9 months (95% CI: 20.3– not estimable; range: 2.4–35.8 months). The median duration of treatment was 9.4 months (range: 0.1–32.0 months). The ORR and response duration reported for axitinib compared favorably to Phase II studies of other similar TKIs.\(^10,11\) As with other single-arm Phase II studies, results may have been especially favorable due to subject selection bias, reflected in the observation that 42% of the patients had no MSKCC unfavorable risk factor. Nevertheless, this study demonstrated the potential for therapeutic efficacy with axitinib in the treatment of patients with metastatic RCC. Notably, a recent update of these data reported a 5-year survival rate of 20.6% (10.9%–32.4%) after a median follow up of 5.9 years.\(^12\)

The second Phase II study evaluated the activity of axitinib in patients previously treated with VEGF-targeted TKIs. Investigators enrolled patients with metastatic RCC who had
Axitinib and renal cell carcinoma

progressive disease on treatment with sorafenib.13 This open-label, multicenter study enrolled 62 patients. Axitinib was given as 5 mg twice daily, with a provision for dose escalation to 10 mg twice daily in the absence of greater than grade 2 adverse effects, and in patients with no significant increase in blood pressure or hypertension, with a protocol amendment later introduced to prohibit dose escalation in patients receiving antihypertensive medications. Subjects were required to have received sorafenib, but they could have had other therapies as well; 74.2% of patients had received two or more lines of any systemic therapy prior to entry, and 22.6% of the cohort had received sunitinib in addition to sorafenib. This trial mainly enrolled patients with clear cell RCC. Only three subjects had nonclear cell histologies. All 62 subjects had had a prior nephrectomy. Fourteen patients experienced a PR for an ORR of 22.6% (95% CI: 12.9%–35.0%), and an additional eleven patients (17.7%) achieved SD. Approximately half of the patients (33 of 62) had their axitinib dose escalated to >5 mg twice daily, but the observed response rates did not differ by axitinib dose. Most (80%) of the evaluable patients demonstrated some degree of target lesion regression. The median duration of response was 17.5 months (95% CI: 7.4 – not estimable). The median progression-free survival (PFS) was 7.4 months (95% CI: 6.7–11.0) and the median OS was 13.6 months (range: 8.4–18.8 months) from the start of axitinib therapy. This Phase II study established the activity of axitinib in patients with clear cell RCC refractory to conventional VEGF-inhibitor therapy. PRs were seen in 26.7% of patients with one prior antiangiogenic therapy, and in 7.1% of patients who had received both sunitinib and sorafenib. The median number of prior therapies was two. Not surprisingly, response rates were lower in this study,11 when compared to the results in a population that had only received prior cytokine therapy.6 The authors hypothesized that axitinib’s efficacy in the second- and third-line setting after prior TKIs might have been due to axitinib’s high potency and high selectivity for VEGFR1, VEGFR2, and VEGFR3.

Following two successful Phase II studies, a large, international, multicenter, randomized Phase III study was designed for registration purposes (The trial of comparative Oncology Group performance status of 0 or 1. MSKCC risk stratification identified 28% of the patients as favorable risk, 37% as intermediate risk, and 33% as poor risk. Patients were randomized 1:1 to either sorafenib 400 mg twice daily or axitinib 5 mg twice daily. Randomization was stratified by performance status and by the type of first-line therapy. Axitinib dosing was titrated to 7 mg twice daily, and then to 10 mg twice daily in the absence of greater than grade 2 toxicity for 2 weeks, if their blood pressure was <150/90 mmHg and if the patient was not on antihypertensive medication. The primary endpoint was PFS, as assessed by a blinded independent review.

The trial demonstrated a statistically significant improvement in PFS for axitinib as compared with sorafenib (hazard ratio [HR] =0.67; 95% CI: 0.54–0.81;  P<0.0001, log–rank test).14 The median PFS was 6.7 months for patients treated with axitinib and 4.7 months for those randomized to sorafenib. Axitinib conferred clinical benefit in terms of PFS to patients regardless of prior therapy such as sunitinib (4.8 months versus 3.4 months, respectively;  P=0.0107) or prior cytokine therapy (12.5 months versus 6.5 months, respectively;  P<0.0001). Similar findings were observed after further segregating the groups by the specific cytokine therapies they had received previously. There were no CRs, but the PR rate was 19% for axitinib and 9% for sorafenib (P=0.0001), and the median duration of response was 11 months (95% CI: 7.4 – not estimable) for axitinib and 10.6 months (95% CI: 8.8–11.5) for sorafenib. No difference was observed between treatment arms in terms of OS. Based on the statistically significant increase in PFS seen for axitinib, on January 27, 2012, the United States Food and Drug Administration approved axitinib for the treatment of metastatic RCC after one prior systemic therapy, and other regulatory agencies followed suit (see Table 1 for a review on US FDA (Food and Drug Administration)-approved VEGF inhibitors for RCC). It is important to emphasize that given no observed difference in OS between axitinib and sorafenib, the overall benefit compared to sorafenib appears modest in the post-sunitinib setting.15

The updated results from the AXIS trial were reported in the spring of 201316 and showed that axitinib was associated with a median PFS of 8.3 months, as compared to 5.7 months for sorafenib (HR =0.66; 95% CI: 0.55–0.78; one-sided  P<0.0001) and a median OS of 20.1 months for axitinib compared to 19.2 months for sorafenib (HR =0.97; 95% CI: 0.80–1.17; one-sided  P=0.3744). The fact that there was no significant survival advantage is perhaps not surprising given
that 54% of the axitinib group and 57% of the sorafenib group received additional therapy, and that 23% and 25% of each drug group, respectively, went on to receive two or more subsequent treatments. The availability of multiple salvage therapies may limit the ability to interpret the efficacy of drugs in this setting.\textsuperscript{17}

Axitinib has also been evaluated in the first-line setting in metastatic RCC. A multicenter trial randomized patients with treatment-naïve metastatic clear cell RCC, measurable disease, and an Eastern Cooperative Oncology Group PS of 0 or 1 to either axitinib 5 mg twice daily or sorafenib 400 mg twice daily (2:1 randomization).\textsuperscript{18} This trial enrolled 288 patients, primarily from outside the US, with 153 patients coming from the Ukraine, Russia, and India. The patients were well balanced for demographic and clinical variables, such as age (median age: 58 years in both groups), MSKCC

<table>
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<tr>
<th>Drug</th>
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<th>PFS Median (months)</th>
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<th>OS Median (months)</th>
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<td>III, First line</td>
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<td>0.57 (0.45–0.72)</td>
<td>23.3</td>
<td>0.86 (0.72–1.04)</td>
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<td>Bev + IFN\textalpha versus placebo + IFN\textalpha</td>
<td>First line</td>
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<td>III, First line</td>
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<td>Sunitinib versus placebo</td>
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<td>5</td>
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<td>Pazopanib versus placebo</td>
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<td>All patients:</td>
<td>0.46 (0.34–0.62)</td>
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<td>145</td>
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<td>Axitinib Motzer et al\textsuperscript{51}</td>
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Abbreviations: RCC, renal cell carcinoma; n, number; PFS, progression-free survival; HR, hazard ratio; CI, confidence interval; OS, overall survival; Bev, bevacizumab; IFN\textalpha, interferon-alpha; Tem, temsirolimus; NR, not reported.
favorable risk status (49% in the axitinib group versus 55% in the sorafenib group), and prior nephrectomy (85% in the axitinib group versus 90% in the sorafenib group). No statistically significant difference was observed between treatment groups in terms of the primary endpoint; the median PFS was 10.1 months for axitinib and 6.5 months for sorafenib (HR = 0.77; 95% CI: 0.56–1.05). The objective response rate was significantly higher with axitinib than sorafenib: 32% versus 15%, respectively (one-sided P = 0.0006).

Finally, axitinib has been studied in the neoadjuvant setting. A single-institution Phase II trial\(^\text{19}\) evaluated the efficacy of axitinib in terms of downsizing tumors in 24 patients with biopsy-proven clear cell RCC. The study protocol planned for treatment with axitinib for 12 weeks prior to surgery. Twenty-two patients completed therapy as planned, with one patient undergoing surgery after 11 weeks and one patient stopping therapy after 7 weeks due to acute kidney injury and taken to surgery at that time, earlier than anticipated. Notably, 100% of evaluable patients (number [n] = 23) showed tumor shrinkage with a median reduction in tumor diameter of 28.3%, and a 45.8% PR (n = 24). All patients had their axitinib dose up-titrated during therapy with a side effect profile similar to that seen in other studies.\(^\text{19}\)

**Dose titration**

Similar to other therapies that inhibit VEGF–VEGFR signaling, axitinib induces hypertension, which appears to be a biomarker for drug activity.\(^\text{20}\) In both Phase II studies described earlier,\(^\text{6,13}\) patients that developed a diastolic blood pressure (DBP) > 90 mmHg were noted to have significantly longer median OS and ORR when compared to those without elevated DBP (30 months versus 9.8 months, respectively; 50.8% versus 10.2%, respectively).\(^\text{19}\) Similar results were seen in a pooled analysis of data from three Phase II studies of 178 patients with metastatic RCC.\(^\text{22}\) According to that analysis, the median PFS was 14.6 months for patients with elevated DBP compared with 7.86 months for patients without elevated DBP (HR = 0.590; 95% CI: 0.402–0.866). The median OS for the same comparison was 29.5 months, as compared with 18.5 months (HR = 0.622; 95% CI: 0.411–0.942). Another recent pooled analysis included two Phase II trials in RCC, as well as three other single-agent axitinib Phase II studies in other tumor types. Among 238 patients in the analysis, those with at least one recorded elevated DBP level had a significantly lower risk of death (HR = 0.55; P < 0.001) and significantly greater ORRs (43.9% versus 12.0%; P < 0.001).\(^\text{20}\) Interestingly, in a recent meta-analysis,\(^\text{23}\) axitinib was found to be more likely to cause hypertension (40.1%) than sorafenib (23.4%; P < 0.0001), sunitinib (21.6%; P < 0.0001), pazopanib (35.5%; P = 0.34), and vandetanib (24.2%; P < 0.0001).\(^\text{21}\)

One hypothesis to explain the finding of varying degrees of blood pressure response to axitinib treatment is that patients who do not develop hypertension may not have sufficient plasma exposure to axitinib. A retrospective analysis of 590 subjects who received axitinib in a total of 17 trials was performed,\(^\text{22}\) which included 181 patients with metastatic RCC. This trial found, by logistic regression, a significant (P < 0.0001) relationship between axitinib plasma exposure and the probability of a response (ie, a 1.5-fold increase in the probability of achieving a PR for every 100 h×ng/mL increase in the area under the curve [AUC]). They also explored the relationship between AUC and clinical endpoints such as PFS and OS. By stratifying the patients into groups of AUC greater than or equal to the axitinib total daily therapeutic exposure of 300 h×ng/mL (high AUC) or < 300 h×ng/mL (low AUC), they found that PFS was prolonged in the high AUC group compared with the low AUC group (13.8 months versus 7.4 months, respectively; HR = 0.558; P = 0.003). Similarly, these investigators also found prolonged OS in the high AUC group compared with the low AUC group (37.4 months versus 15.8 months, respectively; HR = 0.489; P < 0.001). This trial also identified a relationship between blood pressure and response to therapy. The patients that were noted to have DBP ≥ 90 had an improved PFS compared with patients who did not (14.6 months versus 7.9 months, respectively; HR = 0.590; P = 0.006) and OS (29.5 months versus 18.5 months, respectively; HR = 0.622; P = 0.024). These findings opened the door to further investigation regarding the relationship between axitinib exposure and outcomes.

Dose titration was formally evaluated in a randomized Phase II study of patients with untreated metastatic RCC.\(^\text{24}\) In this trial, 213 patients received axitinib 5 mg twice daily for 4 weeks, and the 112 patients that did not develop hypertension or significant adverse events (AEs) were randomized to titration to axitinib 10 mg twice daily or they continued 5 mg twice daily with placebo titration. Fifty-four percent (95% CI: 40–67) of patients in the axitinib titration group and 34% (95% CI: 22–48) of patients in the placebo group had an objective response (one-sided P = 0.19), while 59% (95% CI: 49–70) of nonrandomized patients had an objective response. There was no significant PFS difference reported between the groups (titration: 14.5 months; placebo: 15.7 months; nonrandomized: 16.6 months). The HR for PFS with axitinib titration when compared to placebo did not reach statistical significance (HR = 0.85; 95% CI: 0.54–1.35; one-sided P = 0.24). AEs were similar as in other studies,\(^\text{8,13,14}\) (both
phase II studies and the phase III study), but hypertension, hand–foot syndrome, and vomiting were more common in the axitinib titration group when compared with the placebo group. For the subset of 73 patients with pharmacokinetic sampling, the patients who were eligible for dose titration had both lower area under the plasma concentration–time curve (AUC24; 176 ng•h/mL and 187 ng•h/mL for the axitinib group and placebo group, respectively) and lower maximum observed plasma concentration at baseline (Cmax; 28.6 ng/mL and 22.5 ng/mL for the axitinib and placebo groups, respectively) when compared with the group not eligible for dose titration (AUC24 = 432 ng•h/mL; Cmax = 38.7 ng/mL). The authors concluded that the patients eligible for dose titration likely derived clinical benefit, although the results did not reach statistical significance.

The final statement as to whether or not dose titration is useful may have come from the AXIS study. Selected patients appropriate for dose titration received axitinib 7 mg twice daily and then 10 mg twice daily. The presentation of the data at the American Society for Clinical Oncology (ASCO) conference in 2012 did not report an improved survival rate in patients who were able to receive at least one dose escalation. However, when the OS analysis and updated results from the AXIS trial were presented in 2013, an evaluation of patients 12 weeks after the start of treatment showed that patients with a DBP ≥90 mmHg had an improved OS of 20.7 months compared with 12.9 months for patients with lower blood pressure values (HR = 0.716; 95% CI: 0.537–0.957; $P = 0.0329$), further supporting the concept of using dose titration when indicated.

Safety and tolerability of axitinib

Axitinib is well tolerated. Published experience in the Phase II and III settings confirms a toxicity profile comparable to other agents in the class. In a Phase II study in the post-cytokine population, 15 patients (28.8%) required dose reductions and six (11.5%) discontinued therapy due to AEs such as hypertension, stomatitis, fatigue, diarrhea, or joint pain. The most frequent grade 3–4 toxicities were diarrhea (9.6%), hypertension (15.4%), and fatigue (7.7%). Thirty (57.7%) patients experienced any grade hypertension, of which 18 patients had a history of pre-existing hypertension. Hypertension resolved after adjusting or starting antihypertensive medications in 22 of these 30 patients. Seven of the remaining eight patients were hypertensive prior to starting therapy. Hand–foot syndrome was only noted in four (7.7%) patients, and eleven (21.1%) patients experienced bleeding complications, but these were predominantly epistaxis and only one (hematuria) was grade 3. The second Phase II study, conducted in patients refractory to sorafenib, again suggested that the drug is generally well tolerated, as the majority of AEs were grade 1–2. The most common grade 3 AEs were hand–foot syndrome (16.1%), fatigue (16.1%), hypertension (16.1%), and diarrhea (14.5%).

The AXIS trial confirmed the overall safety profile of axitinib. Diarrhea, hypertension, fatigue, anorexia, nausea, and dysphonia all occurred to some degree in more than 30% of patients receiving axitinib. Hypothyroidism occurred in 19% of patients with only one grade 3 hypothyroidism event. This is important, as changes in thyroid-stimulating hormone levels appear to correlate with fatigue in patients taking axitinib. The most common grade 3 or greater AEs in the AXIS trial were hypertension (16% axitinib; 11% sorafenib), fatigue (11% axitinib; 5% sorafenib), and diarrhea (11% axitinib; 7% sorafenib). Cutaneous toxicities were infrequent with axitinib, with only 5% of patients experiencing hand–foot syndrome (compared to 16% in the sorafenib arm) and <1% of patients receiving axitinib reporting rash. The rates of grade 3 thrombocytopenia and neutropenia were similar for both drugs (0%–1%), as was grade 3 lymphopenia (3% for axitinib versus 4% for sorafenib). Grade 3 anemia was more common for sorafenib (4%) compared with axitinib (<1%). No treatment toxicity deaths were seen in the axitinib group. There were two in the sorafenib group—one due to retroperitoneal bleeding associated with tumor necrosis in a patient on anticoagulation, and the other due to gastrointestinal bleeding. It should be noted that although poorly controlled hypertension can lead to serious cardiovascular events, cardiotoxicity has rarely been reported in any of the reported axitinib studies. Three patients in the AXIS trial discontinued axitinib therapy due to a transient ischemic attack (TIA), while no TIAs were reported for sorafenib.

Effect of axitinib on quality of life and functional assessment

Health-related quality of life has been assessed in various studies. In the Phase II study conducted by Rixe et al, axitinib responders were more likely to have diarrhea than nonresponders, but they were less likely to have deterioration in social function and global quality of life. The second Phase II study of axitinib presented by Rini et al evaluated patient-reported disease-related symptoms at baseline and
after 20 weeks (day 141) of treatment. Significant adverse changes in appetite, weight, energy, and bone pain were noted, and patients reported being increasingly bothered by side effects compared to baseline. Finally, in the AXIS trial, quality of life was assessed as a component of a composite endpoint of the first occurrence of death, progression, or deterioration of symptoms. Axitinib was found to confer a 16% lower risk for this composite endpoint compared to sorafenib (one-sided \( P=0.0203 \)). Discontinuation due to treatment-related side effects was lower with axitinib (4% of patients; most common causes were fatigue and TIAs) compared to sorafenib (8% of patients; most common causes were hand–foot syndrome, diarrhea, and asthenia), which suggests better overall tolerability of axitinib compared to sorafenib. Further supporting the tolerability of axitinib was the fact that dose reduction was more frequent with sorafenib treatment (52%) than with axitinib (31%).

Quality of life was further evaluated within AXIS by Cella et al, where investigators used the Functional Assessment of Cancer Therapy (FACT) Kidney Cancer Symptom Index (FKSI), FKSI disease-related symptoms (FKSI-DRS), and the European Quality of Life self-report questionnaire (EQ-5D). This study found that the quality of life of patients on axitinib or sorafenib was similar to that of the general US population at baseline, and it was maintained until the end of treatment, when meaningful worsening of patient-reported outcomes was seen for both drugs. There was no significant difference noted in terms of quality of life for those taking axitinib or sorafenib.

**Axitinib in other diseases**

Axitinib is currently being evaluated as a possible treatment in multiple different settings, such as recurrent glioblastoma, melanoma, prostate cancer, pheochromocytoma/paraganglioma, hepatocellular carcinoma, nasopharyngeal, and other head and neck cancers. Thus far, it has shown promising results in a Phase II study in thyroid cancer enrolling all histologies, with PR in 30% of patients and SD in 38% of patients. Combining axitinib with traditional chemotherapy, however, has not yielded good results. In colorectal carcinoma, it was inferior to bevacizumab when given with FOLFOX (folinic acid–fluorouracil–oxaliplatin) in a Phase II study. In metastatic breast cancer, docetaxel with axitinib versus placebo did not result in a significantly prolonged time to progression of disease. Gemcitabine with axitinib or placebo did not yield any significant improvement in OS in advanced pancreatic cancer. In lung cancer, axitinib has been compared with cisplatin and pemetrexed versus placebo for nonsquamous non-small-cell lung cancer without significant difference in PFS, the primary endpoint. Comparing the addition of axitinib versus bevacizumab to paclitaxel/carboplatin also yielded no significant PFS improvement for axitinib.

**Future directions**

Research in regards to the effects of axitinib in the RCC setting and other select non-RCC indications is ongoing. Currently, axitinib is only approved for metastatic RCC, but further evaluation in regards to its potential neoadjuvant utility is ongoing with a Phase II trial, the Axipan study, which is currently active in France with results pending. Axitinib is currently being examined in the adjuvant setting to decrease the recurrence risk for high-risk RCC patients after nephrectomy (NCT01599754). It is being studied in patients with metastatic RCC unsuitable for nephrectomy (NCT01693822), and studies are ongoing for patients with nonclear cell disease, such as for those previously treated with temsirolimus (NCT01798446). Phase IB studies of axitinib with other molecularly targeted agents such as everolimus and temsirolimus have completed accrual and await presentation (NCT01334073 and NCT01529138, respectively). Combination studies with various antibodies also are underway, including a Phase IB study of axitinib and TRC105 for patients with RCC who have progressed after nephrectomy (NCT01806064). Finally, axitinib also is being examined in advanced RCC in combination with inhibitors of the PD-1/PD-L1 axis, such as pembrolizumab (MK-3475, a PD-1 inhibitor) (NCT02133742).

**Conclusion**

As a highly potent inhibitor of the VEGF–VEGFR signaling axis that is tolerable, axitinib has emerged as an important therapeutic option for use in the treatment of patients with refractory metastatic clear cell RCC. Axitinib has been found to confer clear-cut benefits to patients in terms of prolonging the PFS in the second-line setting over sorafenib. Its toxicity profile is no worse than that of other drugs in its class, and a useful strategy that may enhance clinical benefit has been developed to increase the dose in the absence of treatment-related AEs. Because of the choice of sorafenib as a comparator in the AXIS study, questions remain regarding axitinib’s place in the treatment of patients with refractory disease relative to everolimus that likely will remain unanswered. Additionally, because the randomized first-line study was underpowered, questions remain as to whether there is a
place for axitinib in the first-line setting. In the end, axitinib makes an important and useful contribution to management in second-line metastatic clear cell RCC for use when VEGFR-inhibitor therapy is desired, and the potential exists for greater utility in the future as a combination partner with emerging novel therapies, such as with targeted immunomodulatory agents.

Disclosure
The authors report no conflicts of interest in this work.

References


