Renal cell carcinoma with non-clear cell histology or sarcomatoid differentiation: recent insight in an unmet clinical need

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Renal cell carcinoma (RCC) is the 13th most common malignancy resulting in more than 400,000 new cases and 175,000 deaths per year worldwide (1). The histologic subset clear cell RCC (ccRCC), or conventional RCC, comprises 75% of cases. Clinical trials for 20 years in patients with metastatic RCC (mRCC) have therefore focused on patients with clear cell histology. Today 15 drugs have shown clinical benefit in this group of patients (Figure 1), leading to substantial, clinically meaningful improved outcomes for the vast majority of patients with mRCC (2). A wide availability of diagnostic CT has led to stage migration, with more than half of patients with RCC now identified as incidental findings (3). For patients with localized RCC, surgical options have improved with minimally invasive techniques, including robotic techniques and laparoscopic techniques (4), and renal cryoablation has also shown promise (5). Moreover, decline in tobacco use and increased physical activity (6) contribute to the improved results. In essence, mortality from RCC is declining (7); a recent nationwide study showed a 7% yearly reduction in mortality from RCC (8). With this achievement in the vast majority of patients, focus has shifted to the remaining RCC patients with area of needs. Progress has now been made in patients with non-clear cell histology (non-ccRCC), also called rare histologic variants, and patients with sarcomatoid histology.

Non-ccRCC comprises 25% of RCC and is a diverse group of tumors that includes papillary type I, papillary type II, chromophobe, medullary, collecting duct, TFE3 translocation, and unclassified RCC. Outcomes with targeted agents have generally been poor in patients with non-ccRCC (9-12). The ASPEN and ESPN trials, evaluating everolimus versus sunitinib, showed modest efficacy, with objective response rates (ORR) less than 20%, including less than 1% complete response (CR), and median progression free survival (PFS) less than 9 months (13,14). Interleukin-2 based immunotherapy did not show efficacy in non-ccRCC (15). In addition, both ccRCC and non-ccRCC may be associated with sarcomatoid differentiation (sarcomatoid RCC, sRCC). Only 10–20% of patients with mRCC have sRCC, but sRCC is associated with an aggressive phenotype, and an advanced stage at diagnosis (16); the rate of sarcomatoid differentiation was doubled in synchronous versus metachronous mRCC (17). These patients have a particularly poor prognosis, and systemic therapies in sRCC have been mostly ineffective (18,19).

In 2020/2021, three important studies have brought about greater insight in this specific area. Bradley A. McGregor and colleagues reported in the January 2020 edition of Journal of Clinical Oncology the results of an investigator-initiated multicenter Phase II study of atezolizumab and bevacizumab for patients with metastatic RCC with non-ccRCC and/or sarcomatoid features (20). The study included a total of 60 patients
at 4 academic institutions in the USA; 42 patients (70%) with non-ccRCC and 18 (30%) with ccRCC with ≥20% sarcomatoid differentiation. Most patients had prior nephrectomy, i.e., had metachronous metastatic disease (17) and the presence of bone metastases was unusually low (2%), therefore, selection of patients with less aggressive underlying biology may be suspected. The primary endpoint was ORR as the best overall response by RECIST version 1.1, by investigator assessment. Patients were treated with atezolizumab 1,200 mg and bevacizumab 15 mg/kg intravenously every 3 weeks. Median time of follow-up was 13 months. The ORR for the overall population was 33%, with 2 patients (3%) obtaining a CR, 50% of patients with ccRCC with sarcomatoid differentiation, and 26% in patients with non-ccRCC. The overall median PFS was 8.3 months (95% CI, 5.7 to 10.9 months), median OS was not reached; the PFS/OS results for subgroups with sarcomatoid or non-ccRCC was not provided. A baseline biopsy was mandatory per protocol. Tissue was, however, not suitable for PD-L1 staining, therefore, archival nephrectomy tissue was used instead in 36 (60%) patients. The authors used a PD-L1 score ≥1% tumor cells to determine positivity. Overall, 42% of patients were PD-L1 positive. Overall, ORR in PD-L1–positive vs. PD-L1–negative patients was 60% vs. 19%; in ccRCC with sarcomatoid differentiation 50% vs. 29%, respectively; in patients with non-ccRCC 67% vs. 14%. A summary of key findings is listed in Table 1.

IMmotion151 was a randomized, open-label, phase III study evaluating the efficacy and safety of atezolizumab plus bevacizumab versus sunitinib in patients with previously untreated, inoperable RCC with a clear cell component (23). A prespecified subgroup analysis of patients with sRCC was recently published by Brian Rini and colleagues in Eur Urol (21). Of the 915 intention-to-treat (ITT) patients included in IMmotion151, a total of 142 (15.5%) patients with sRCC (81% had ccRCC, 19% had non-ccRCC) received atezolizumab + bevacizumab or sunitinib. Moreover, PD-L1 expression (≥1% vs. <1%) on tumor-infiltrating immune cells was assessed. In total, 86 patients (9.4% of ITT) had both sarcomatoid features as well as being PD-L1 positive. The median follow-up was 13 months. In patients with sarcomatoid features, results showed improved outcomes with atezolizumab plus bevacizumab versus sunitinib; median PFS 8.3 vs. 5.3 months (HR 0.52); median OS 21.7 vs. 15.0 months (HR 0.61); ORR 49% vs. 14% with CR rate 10% vs. 3% and ongoing response in 52% vs. 30% (Table 1). In the subset of patients with both sarcomatoid features and PD-L1 positive staining almost identical results was noted; median PFS 8.6 vs. 5.6 months (HR 0.45); median OS 19.3 vs. 15.0 months (HR 0.61); ORR 56% vs. 12% with CR rate 14% vs. 4% and ongoing response in 45% vs. 0%. Data in patients with both sarcomatoid features and PD-L1 negative staining was not provided. However, compared with
the intention to treat population where patients treated with atezolizumab plus bevacizumab had PFS of 11.2 months and OS of 33.6 months, outcomes for patients with sarcomatoid features or sarcomatoid features with PD-L1 expression were less favorable, and treatment with atezolizumab plus bevacizumab did not overcome the negative prognostic impact of sarcomatoid differentiation or PD-L1 expression.

CheckMate 214 was a randomized phase III study evaluating the efficacy and safety of nivolumab plus ipilimumab versus sunitinib (24). A post hoc analysis in patients with sRCC was recently published by Nizar Tannir and colleagues in Clinical Cancer Research (22). Of the 1,096 patients included in the phase III study, 139 (13%) patients had sRCC and were IMDC intermediate/

Table 1 Key study findings in patients with metastatic renal cell carcinoma with sarcomatoid differentiation and/or non-clear cell histology

| Study | ORR (%) | CR-rate (%) | PFS | OS | 42-mo OS (%) | Follow-up |
|-------|---------|-------------|-----|----|--------------|-----------|
| Bradley A. McGregor, JCO (20) (N=60) | 33 | 3 | 8.3 mo | NR | 13 mo |
| PD-L1+ (15/36; 42%) | 60 |
| PD-L1– (9/15; 58%) | 19 |
| sRCC overall (N=18/60) | 50 | | | | 13 mo |
| SRCC + PD-L1+ | 50 |
| SRCC + PD-L1– | 29 |
| Non-ccRCC overall (N=42/60) | 26 | | | | 13 mo |
| SRCC + PD-L1+ | 67 |
| SRCC + PD-L1– | 14 |
| Immotion 151 (N=915) | | | | | |
| Atezolizumab + bevacizumab | 37 | 5 | 11.2 mo | 33.6 mo | 13 mo |
| Sunitinib | 33 | 2 | 8.4 mo | 34.9 mo |
| Brian Rini, Eur Urol (21) | | | | | |
| sRCC overall (N=142/915; 15.5%) | | | | | |
| Atezolizumab + bevacizumab | 49 | 10 | 8.3 mo | 21.7 mo | 13 mo |
| Sunitinib | 14 | 3 | 5.3 mo | 15.4 mo |
| sRCC + PD-L1+ (N=86/142; 9%) | | | | | |
| Atezolizumab + bevacizumab | 56 | 14 | 8.6 mo | 19.3 mo | 13 mo |
| Sunitinib | 12 | 4 | 5.1 mo | 15.0 mo |
| CheckMate 214 I/P risk (N=847) | | | | | |
| Nivolumab + ipilimumab | 42.1 | 10.1 | 11.6 mo | 47.0 mo | 52 | 42 mo |
| Sunitinib | 26.3 | 1.4 | 8.3 mo | 26.6 mo | 39 |
| Nizar Tannir, Clin Cancer Res (22) | | | | | |
| sRCC overall (N=139/847; 16.4%) | | | | | |
| Nivolumab + ipilimumab | 60.8 | 18.9 | 26.5 mo | NR | 50.1 | 47.7 mo |
| Sunitinib | 23.1 | 3.1 | 5.1 mo | 14.2 mo | 22.6 |
| sRCC + PD-L1+ (N=69/133; 52%) | | | | | |
| Nivolumab + ipilimumab | 69.4 | 22.2 | NR | NR | 50 | 47.7 mo |
| Sunitinib | 24.2 | 3 | 4.4 mo | 20.9 mo | 30 |
| sRCC + PD-L1– (N=64/139; 48%) | | | | | |
| Nivolumab + ipilimumab | 54.3 | 17.1 | 10.9 mo | 40.4 mo | 48 | 47.7 mo |
| Sunitinib | 20.7 | 3.4 | 5.1 mo | 13.8 mo | 15 |

sRCC, sarcomatoid renal cell carcinoma; ccRCC, clear cell renal cell carcinoma; ORR, objective response rate; CR, complete response; PFS, median progression free survival; OS, median overall survival.
poor-risk; the ratio of sRCC vs. non-ccRCC histology was not given. The baseline tumor cell PD-L1 expression was assessed and approximately half of patients had a positive tumor PD-L1 expression. Minimum follow-up was 47.7 months. Nivolumab plus ipilimumab showed notable benefits over sunitinib in patients with sRCC; median OS not reached vs. 14.2 months (HR 0.45); 42-month probability of survival 50.1% vs. 22.6%; median PFS 26.5 vs. 5.1 months (HR 0.54); 36-month probability of PFS 48.2% vs. 20.3%; ORR 60.8% vs. 23.1%, with CR-rate of 18.9% vs. 3.1%, respectively (Table 1). Moreover, ongoing response was observed in 69% vs. 53%. Efficacy outcomes were notably better with nivolumab plus ipilimumab versus sunitinib, regardless of PD-L1 expression; 22.2% of patients with sRCC and tumor PD-L1 expression ≥1% achieved CR with NIVO+IPI, and 17.1% of patients with sRCC and tumor PD-L1 expression <1% achieved CR with NIVO+IPI.

The data presented here indicates with a 42 months’ minimum follow-up, that nivolumab plus ipilimumab offers patients with IMDC intermediate/poor risk and sRCC the potential of ORR in two thirds of cases; CR in approximately one fifth; durable responses in two thirds of cases obtaining response; 3-year PFS-plateau of 48%; identical OS-plateau in sRCC as in non-sRCC patients; and identical OS-plateau in sRCC patients regardless of PD-L1 expression. This supports nivolumab plus ipilimumab as a standard of care for first-line treatment in patients with sRCC disease. In essence, the negative prognostic impact of sRCC as well as PD-L1 expression were overcome with use of nivolumab plus ipilimumab. We are awaiting long-term follow-up data from each of the three trials; KEYNOTE-426 trial assessing pembrolizumab plus axitinib versus sunitinib, JAVELIN Renal 101 trial assessing avelumab plus axitinib versus sunitinib, and CheckMate 9ER assessing nivolumab plus cabozantinib versus sunitinib.

Sarcomatoid transformation in RCC is characterized by a transformative growth pattern of the epithelial neoplasm into malignant spindle-shaped cells. Histologically, sRCC contains both epithelial (carcinomatous) and mesenchymal (sarcomatoid) components, unlike a true sarcoma of the kidney. The correct diagnosis of sRCC is based on the experience of the pathologist and is based on the degree of assessment. Assessment of a core biopsy versus thorough assessment of the entire nephrectomy specimen may impact findings. Also the timepoint of assessment has impact; assessment at the time of metastasis or at the time of primary diagnosis will impact the rate of sRCC diagnosis. Finally, is the correct cutoff 20% or 1%? These dilemmas were all clearly illustrated by the articles reviewed (20-22). In most cases, an underestimation of sRCC may occur. Nevertheless, whereas sRCC used to be a marker of severe poor prognosis and resistant to therapy, treatment with nivolumab and ipilimumab now offers hope of a potential cure in a substantial subset of patients with sRCC. This must be considered a truly remarkable and monumental achievement within the research of sRCC.

PD-L1 staining is without consistency in mRCC, as illustrated by the above studies. Three different antibodies were used in the three studies: 405.9A11 mouse monoclonal antibody (20), VENTANA PD-L1 SP142 assay (21), and Dako PD-L1 IHC 28-8 pharmDx test (22), while staining was on either tumor cells (20,22) or tumor-infiltrating immune cells (21). This highly illustrates the lack of consistency in assessment methods. PD-L1 expression as a predictive marker has been assessed for a decennium. In patients with mRCC this marker only has a role as a negative prognostic factor, not a predictive factor for therapy selection, as both patients with PD-L1 positive as well as PD-L1 negative tumors benefit from therapy. Treatment with nivolumab in combination with ipilimumab now represents the new backbone of therapy for patients with IMDC intermediate/poor risk or sRCC, irrespective of PD-L1 expression. Moreover, for patients with sRCC, the negative prognostic impact of sRCC as well as PD-L1 expression were overcome with use of nivolumab plus ipilimumab.

Other areas of need for patients with mRCC may now hopefully be the focus of further intense investigations, especially patients with brain metastasis. Other areas of need include: patients with poor performance status, patients with synchronous metastatic disease, patients with considerable co-morbidity, patients with rapidly growing tumors and concomitant symptoms, and patients with bone metastases or lymph node metastases. Therefore, clinical trials within the aforementioned areas are greatly welcomed.

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