Biological Therapeutic Advances for the Treatment of Advanced Urothelial Cancers

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Abstract: In recent years, diagnostic and therapeutic advances have contributed to a reduction in mortality rates of patients with metastatic urothelial carcinoma (mUC). Immune checkpoint inhibitors have demonstrated efficacy and safety as both first-line and first-line switch maintenance therapy for mUC. For platinum-refractory patients, in addition to immunotherapy, other targeted agents (antibody–drug conjugates and fibroblast growth factor receptor inhibitors) have been approved after demonstrating a clinically relevant advantage in overall response rate, progression-free survival, and overall survival compared to standard of care. Sequential treatment strategies are finally feasible for patients with advanced urothelial carcinoma. This review will summarize the results of the most important phase II–III clinical trials for first-line, switch maintenance, second-line, and subsequent lines of therapy, and describe the most promising clinical trials currently ongoing in these treatment scenarios.

Keywords: metastatic urothelial carcinoma, targeted therapy, immune checkpoint inhibitors, first-line treatment, second-line treatment

Introduction

According to the GLOBOCAN 2020 estimates of cancer incidence and mortality, urothelial carcinoma (UC) is the tenth most commonly diagnosed cancer worldwide, being responsible for approximately 573,000 new cases and 213,000 deaths in 2020.1

As a whole, UC consists of both bladder cancer and upper urinary tract carcinomas arising from either the ureter or the renal pelvis. The main risk factors for these malignancies, which affect more males than females, are smoking, older age, family history of UC, infection with Schistosoma haematobium (in northern and sub-Saharan Africa), occupational exposure to aromatic amines and other chemicals affecting workers in the painting, rubber, or aluminum industries, and arsenic contamination of drinking water.2, 3

As a direct consequence of both the average age and the frequent smoking habit of these patients, subjects with UC are often frail and highly comorbid, with renal impairment being highly prevalent among them. In terms of natural history, UC initially occurs as superficial, often multicentric, non-muscle-invasive, lesions, which are treated by means of repeated transurethral resections (often followed by intracavitary therapies); however, these lesions tend to recur repeatedly over the years until they become muscle invasive, leading to the need for radical cystectomy or nephroureterectomy (in the case of UC arising in the upper urinary tract), preceded or followed by systemic chemotherapy.4, 5
Unfortunately, relapse after surgery occurs in approximately 30–50% of cases, often leading to the development of distant metastases; furthermore, 5% of all UC patients present with de novo metastatic disease. Both synchronous and metachronous metastatic urothelial carcinoma (mUC) patients have a very poor prognosis, although mortality rates have declined recently, at least in part as a consequence of therapeutic improvements, especially in more industrialized countries. Metastatic patients presenting with additional negative prognostic factors according to the Bajorin classification (ie low performance status and/or visceral metastases) have a particularly poor prognosis.6

In this review, we will summarize the most relevant phase II–III clinical trials conducted for first-line, maintenance of switch, second-line, and subsequent lines of therapy in advanced UC, and list ongoing trials.

**Cytotoxic Chemotherapy for Cisplatin-Eligible Metastatic UC Patients**

Locally advanced or metastatic (ie stage IV) UC has been shown to benefit from cytotoxic chemotherapy; despite therapeutic improvements made over the past few years, major international guidelines still recommend first-line cisplatin-containing combination chemotherapy for those patients who are fit enough to receive cisplatin.5,7,8

Among cisplatin-containing regimens, both the gemcitabine plus cisplatin doublet (GC) and the dose-dense MVAC (ddMVAC) polychemotherapy (which, together with cisplatin, also includes methotrexate, vinblastine, and doxorubicin), are commonly used as standard first-line treatments.

In 2000, a large, global, phase III study randomized 405 patients with locally advanced or metastatic UC to receive either GC or standard MVAC chemotherapy. At a median follow-up of 19 months, overall survival (OS) and time to progression (TTP) were similar in the two treatment arms; however, fewer toxic deaths were recorded among patients receiving GC compared with MVAC (1% vs 3%), a difference which was not statistically significant. A 5-year update analysis confirmed that GC was not superior to MVAC in terms of either OS (13.0% vs 15.3%, respectively) or progression-free survival (PFS) (9.8% vs 11.3%, respectively).9

A subsequent large, randomized, phase III trial compared ddMVAC to standard MVAC.10 At a median follow-up of 7.3 years, 24.6% of ddMVAC-treated patients were alive, compared with 13.2% of those treated with standard MVAC; furthermore, less overall toxicity was reported in the ddMVAC group.11 Accordingly, standard MVAC was almost completely abandoned.

Despite the fact that GC had standard MVAC, and not ddMVAC, as the control arm of its pivotal trial, both GC and ddMVAC (with growth factor support) are the present standard of care for metastatic disease.

Although first-line platinum-containing chemotherapy results in objective responses or stable disease in approximately 65–75% of patients, it usually provides limited long-term benefits, in terms of both PFS (6–8 months) and OS (9–14 months).

In order to improve these results, taxane-based combinations were also tested; in particular, a randomized phase III trial was conducted to compare GC and GC plus paclitaxel in 626 patients with locally advanced or metastatic UC. The addition of paclitaxel to GC resulted in higher response rates and a borderline OS advantage, which was not statistically significant in the intention-to-treat analysis; when eligible patients only were considered, the OS improvement for those patients treated with the three-drug regimen reached statistical significance (3.2 months), while no difference was recorded in terms of PFS. Furthermore, as expected, toxicity was worse for the three-drug combination, the incidence of neutropenic fever being 13.2% versus 4.3% for the doublet.12

Considering the limited benefit and the increase in severe toxicity observed, this combination ultimately did not replace the traditional standard of either GC or ddMVAC.

**Cytotoxic Chemotherapy for Cisplatin-Ineligible (or Unfit) Metastatic UC Patients**

Patients with renal impairment (estimated glomerular filtration rate <50–60 mL/min), poor performance status (ECOG 2 or Karnofsky ≤70), hearing loss (grade ≥2), peripheral neuropathy (grade ≥2), and/or symptomatic heart failure (NYHA class III/IV), who account for as many as 40% of all advanced UC patients, are considered cisplatin-ineligible or unfit;13 in this patient population, the doublet of carboplatin plus gemcitabine is the main recommended option.5,7,8

A phase II/III study assessed two carboplatin-containing regimens (carboplatin and gemcitabine or
methotrexate, carboplatin, and vinblastine) in cisplatin-unfit patients, mainly due to a performance status of 2; the overall response rate (ORR) was 42% and 30%, respectively, but ORR dropped to 26% and 20%, with an increased toxicity, among patients who had both a poor performance status and renal impairment.\textsuperscript{14}

However, in the most recent phase III studies with combinations of immune checkpoint inhibitors (ICIs) and chemotherapy, regimens containing cisplatin and carboplatin demonstrated similar OS results to the historical benchmarks.

### Switch Maintenance Immunotherapy Following Cytotoxic Chemotherapy

Since many chemotherapeutic agents, including platinum derivatives, may exert different immunogenic effects, the use of an ICI as a switch maintenance therapy following cytotoxic chemotherapy has been proposed.

Indeed, chemotherapy can induce immunogenic cell death, stimulate the antitumor immune responses through the release of “danger” signals from dying cells,\textsuperscript{15,16} deplete immunosuppressive populations (eg myeloid-derived suppressor cells and regulatory T cells),\textsuperscript{17,18} stimulate tumor infiltration by CD8\textsuperscript{+} cytotoxic T lymphocytes, activate natural killer (NK) cells, enhance antigen presentation, stimulate the generation of tumor neoantigens, and also upregulate the expression of programmed death ligand-1 (PD-L1).

This concept was the basis of JAVELIN Bladder 100, a randomized, phase III trial aimed at evaluating the anti-PD-L1 antibody avelumab as first-line switch maintenance therapy in 700 patients with stage IV UC. Enrolled patients had to have no progression after four to six cycles of first-line platinum-containing chemotherapy. After 4–10 weeks from the last dose of chemotherapy, patients were randomized to receive either avelumab (given at the dose of 10 mg/kg, i.v., every 2 weeks) plus best supportive care (BSC) or BSC alone. Patients were also stratified by best response to first-line chemotherapy (complete or partial response [CR or PR] vs disease stabilization [SD]) and site of metastases (visceral vs non-visceral). The primary endpoint of the study was OS, assessed in two primary populations: all randomized patients and patients with PD-L1-positive tumors, where PD-L1 positivity was evaluated by means of the Ventana SP263 assay, a subpopulation that accounted for 51% of the overall study group. The study succeeded, since avelumab plus BSC significantly prolonged OS vs BSC alone, median overall survival (mOS) being 21.4 months in the avelumab arm versus 14.3 months in the BSC-only arm, resulting in a hazard ratio (HR) of 0.69 (95% confidence interval [CI] 0.56–0.86; \(p=0.001\)); that is, a 31% reduction in the risk of death. Avelumab also significantly prolonged OS in the PD-L1-positive population; indeed, mOS was not reached in the avelumab arm versus 17.1 months in the control arm (HR=0.56, 95% CI 0.40–0.79; \(p<0.001\)).\textsuperscript{19}

An OS benefit with avelumab plus BSC versus BSC alone was observed across a number of prespecified subgroups, including elderly (\(\geq 65\) years) patients (HR=0.63, 95% CI 0.47–0.83), patients with chronic kidney disease (creatinine clearance [CrCl] <60 mL/minute; HR=0.68, 95% CI 0.50–0.94), and patients with visceral, liver, or lung metastases (HRs=0.82, 95% CI 0.62–1.09; 0.92, 95% CI 0.54–1.56; and 0.86, 95% CI 0.56–1.30, respectively).\textsuperscript{20} Furthermore, the OS benefit yielded by avelumab switch maintenance therapy was achieved despite the fact that, among patients progressing on study, 53% of those originally randomized to BSC alone (vs just 9% of those randomized to avelumab plus BSC) received a subsequent PD-1 or PD-L1 inhibitor.\textsuperscript{19,21}

On the basis of the above results, avelumab was approved as a switch maintenance treatment for patients with locally advanced or metastatic UC whose disease had not progressed under first-line platinum-containing chemotherapy, and was thus granted the status of recommended treatment (level 1 evidence) in both the NCCN and ESMO guidelines.\textsuperscript{7,8}

Although in the JAVELIN Bladder 100 trial avelumab was administered at the weight-based dose of 10 mg/kg, the currently approved dose of avelumab, across different cancer types, is 800 mg flat dose, every 2 weeks. This flat dosing was approved based on pharmacokinetic and pharmacodynamic analyses showing minimal differences compared with weight-based dosing across various weight ranges.\textsuperscript{22}

In terms of safety, rates of grade \(\geq 3\) treatment-related adverse events (TRAEs) were 17% in avelumab-treated patients, with the most common grade \(\geq 3\) TRAEs being increased lipase (2.9%) and increased amylase (2.0%).\textsuperscript{19} Furthermore, any grade and grade \(\geq 3\) infusion reactions occurred in 10% and in 1% of avelumab-treated patients, respectively, despite the protocol requirement for premedication with an oral antihistamine and oral acetaminophen before the first four doses of the immunotherapeutic
Immune Checkpoint Inhibition for First-Line Treatment

For the past three decades, first-line systemic therapy for patients with locally advanced or metastatic UC has consisted of platinum-doublet chemotherapy. Beyond chemotherapy, several antiangiogenic agents (bevacizumab, sunitinib, sorafenib) and targeted therapy have been tested in unselected patients, including inhibitors of epidermal growth factor receptor (EGFR, eg gefitinib and cetuximab) and human epidermal growth factor-2 (anti-HER-1–2, eg lapatinib and trastuzumab), which failed to demonstrate any activity.

In this landscape, the recent approval of monoclonal antibodies targeting programmed death ligand-1 (PD-L1) has revolutionized treatment, particularly in patients unfit for cisplatin-based chemotherapy (Table 1).

The initial proof of principle that ICIs show activity in patients with mUC was demonstrated in phase II trials (IMvigor210 with atezolizumab and KEYNOTE-052 with pembrolizumab) in cisplatin-unfit patients with advanced/metastatic UC. Since then, atezolizumab and pembrolizumab have quickly moved into two randomized phase III trials in the first-line setting. Following the results of these two trials, the role of immunotherapy has become a universal component of treatment in fragile patients in Europe (EMA) and the USA (FDA).

Accelerated FDA approval of atezolizumab and pembrolizumab for cisplatin-ineligible chemotherapy-naïve patients with advanced UC was based on single-arm studies (the IMvigor210 trial and the KEYNOTE-052 trial, respectively), which demonstrated favorable ORRs and duration of response (DoR).

In the IMvigor210 trial, with a median follow-up period of 17 months, the ORR was 23%, median PFS 2.7 months, mOS 15.9 months, 12-month survival rate 57%, and mOS after stable disease 19.1 months; OS was considerably longer than the long-standing result obtained with gemcitabine/carboplatin (9.3 months). In the KEYNOTE-052 trial, with a minimum follow-up of 24 months since the last patient enrolled, the ORR for patients with PD-L1-expressing and PD-L1-negative tumors was 47% and 20%, median DoR was not reached, and mOS was 18.5 months and 9.7 months, respectively, suggesting PD-L1 as a powerful predictive biomarker for the response to pembrolizumab in the first-line setting.

A subsequent large retrospective cohort study in a real-life cisplatin-ineligible population receiving pembrolizumab as first-line treatment confirmed lower short-term but superior long-term survival with ICIs and fewer adverse events compared to carboplatin-based chemotherapy.

The combination of higher efficacy in terms of potential for durable benefit, the improvement in quality of life, and the favorable safety profile compared to carboplatin-based chemotherapy initially led the FDA and EMA to approve pembrolizumab and atezolizumab as standard first-line treatment in unselected cisplatin-unfit patients with mUC.

To enrich the response to immunotherapy, and based on the results of phase II studies, three randomized clinical trials were designed to evaluate the combination of ICIs and chemotherapy for previously untreated locally advanced or metastatic UC. Pembrolizumab, atezolizumab, and durvalumab were further evaluated in combination with chemotherapy, although the results and implications were less straightforward compared to phase II trial results.

The first trial was the IM vigor130 trial, which evaluated atezolizumab plus platinum-based chemotherapy, atezolizumab alone, and placebo plus platinum-based chemotherapy in 1213 patients. The median PFS of atezolizumab plus platinum-based chemotherapy was not statistically superior to platinum-based chemotherapy (8.2 vs
6.3 months, respectively; HR=0.82, 95% CI 0.70–0.96; \( p=0.007 \). At a median follow-up of 11.8 months, mOS was 16 months for atezolizumab plus platinum-based chemotherapy and 13.4 months for platinum-based chemotherapy (HR=0.83, 95% CI 0.69–1.00; one-sided \( p=0.027 \)). The ORR was 47% for atezolizumab plus platinum-based chemotherapy, 23% for atezolizumab alone, and 44% for platinum-based chemotherapy. The prespecified dual primary endpoints of OS and PFS were not met, with a median progression-free survival (mPFS) in the pembrolizumab plus chemotherapy arm of 8.3 months compared with 7.1 months for chemotherapy alone (HR=0.78, 95% CI 0.65–0.93; \( p=0.003 \)) and mOS of 17.0 versus 14.3 months, respectively (HR=0.86, 95% CI 0.72–1.02; \( p=0.0407 \)).

Another phase III study of anti-PD-1 in combination with chemotherapy in first-line setting mUC was KEYNOTE-361, in which 1010 patients were randomly assigned to receive pembrolizumab in combination with platinum-based chemotherapy, pembrolizumab alone, or platinum-based chemotherapy. The ORR was 47% for atezolizumab plus platinum-based chemotherapy, 23% for atezolizumab alone, and 44% for platinum-based chemotherapy. The prespecified dual primary endpoints of OS and PFS were not met, with a median progression-free survival (mPFS) in the pembrolizumab plus chemotherapy arm of 8.3 months compared with 7.1 months for chemotherapy alone (HR=0.78, 95% CI 0.65–0.93; \( p=0.003 \)) and mOS of 17.0 versus 14.3 months, respectively (HR=0.86, 95% CI 0.72–1.02; \( p=0.0407 \)).

In subgroup analyses of both studies, in patients with low PD-L1 expression the ICI arms had reduced survival compared to chemotherapy regimens. On the other hand, in patients with high PD-L1 expression there was a trend toward a better outcome in favor of patients treated with immunotherapy alone. In May 2018, based on these results,
the FDA and EMA restricted the use of atezolizumab or pembrolizumab in the first-line setting to patients who are not eligible for cisplatin-containing chemotherapy and whose tumors express high PD-L1, as assessed by companion diagnostic tests, or patients who are considered unsuitable for any platinum-containing chemotherapy regardless of PD-L1 status.

A third study that should be mentioned in this therapeutic context is the DANUBE trial, which explored the role of durvalumab with or without tremelimumab versus platinum-based chemotherapy alone in 1032 patients with untreated, locally advanced, or metastatic UC. The co-primary endpoints were OS in patients with high PD-L1 expression treated with durvalumab as monotherapy compared to patients treated with chemotherapy, and OS in the intention-to-treat population treated with durvalumab plus tremelimumab compared to the population treated with chemotherapy. With a median follow-up of 41.2 months, the study did not meet any of its co-primary endpoints: in the population with high PD-L1 expression, OS was 14.4 months with durvalumab versus 12.1 months with chemotherapy (HR=0.89, 95% CI 0.71–1.11; p=0.30), while in the intention-to-treat population OS was 15.1 months with durvalumab plus tremelimumab versus 12.1 months (95% CI 10.9–14.0) with chemotherapy (HR=0.85, 95% CI 0.72–1.02; p=0.075).32

Two additional ongoing studies are enrolling subjects to investigate the role of combination immunotherapy plus chemotherapy in this population. The NILE trial (NCT03682068) is exploring chemotherapy with or without durvalumab alone or in combination with tremelimumab, and the CheckMate 901 trial (NCT03036098) is comparing the combination of nivolumab and ipilimumab versus chemotherapy in patients ineligible for cisplatin and in patients with elevated PD-L1, along with a substudy evaluating standard cisplatin-based chemotherapy with or without nivolumab. Furthermore, the LEAP-011 trial (NCT03898180) is evaluating pembrolizumab in combination with lenvatinib versus pembrolizumab plus placebo for first-line treatment of cisplatin-ineligible patients with a combined positive score (CPS) ≥10 and platinum-ineligible patients.

### Immune Checkpoint Inhibitors and Targeted Therapies for Second-Line Treatment

Seventy percent of patients receiving platinum-based chemotherapy develop rapid progression. Several salvage therapies are currently approved for patients who have progressed to platinum-based chemotherapy, including checkpoint inhibitors (Table 2), fibroblast growth factor receptor (FGFR) inhibitors, and antiangiogenic agents.

Among PD-L1 checkpoint inhibitors, only pembrolizumab, in a randomized phase III trial, KEYNOTE-045, showed a survival benefit (OS 10.3 vs 7.4 months, p=0.002) versus chemotherapy with paclitaxel, docetaxel, or vinflunine, although there was no difference in PFS. The ORR for the pembrolizumab group was higher than for the chemotherapy group (21.1% vs 11.4%, p=0.001) and comparable between PD-L1 low and PD-L1 high expression subgroups. Treatment-related serious adverse events were reported in 15% of patients receiving pembrolizumab; the treatment discontinuation rate was 5.6%.33

Different results were seen in the phase III study IMvigor211, which showed no benefit in OS or PFS for atezolizumab compared to chemotherapy. Treatment-related serious adverse events were reported in 6% of patients receiving pembrolizumab; the treatment discontinuation rate was 3%.34

In addition, the results of the SAUL trial have recently been published. This was the first prospective single-arm study to confirm the safety and efficacy of atezolizumab in a large, real-world pretreated population with urinary tract carcinoma, including patients with symptomatic brain metastasis, autoimmune disease, ECOG >1, inadequate kidney function, or non-urothelial histology. The mOS was 8.7 months (95% CI 7.8–9.9), mPFS was 2.2 months (95% CI 2.1–2.4), and the ORR was 13% (95% CI 11–16%; 3% CR).35

Of note, contradictory results have been reported in these two large trials on the use of PD-L1 expression as biomarker-driven patient selection. The atezolizumab trial reported a lack of consistency on OS data in PD-L1-positive patients; in contrast, the pembrolizumab trial showed a relevant improved outcome in patients with high expression of PD-L1 (18.5 vs 9.7 months). Direct comparisons should not be made owing to the different definition of PD-L1 positivity in these two trials; thus, any conclusions are highly speculative and uncertain.

Nivolumab, durvalumab, and avelumab currently have no OS data in prospective randomized phase III trials, although in phase II trials similar proportions of patients (17–20%) achieve an overall response.

Another intriguing approach is to explore the molecularly targeted therapies in selected patients with UC. In particular, the dysregulation of fibroblast growth factor
signaling (FGF) can occur in around 20% of patients with mUC via FGF gene alteration, such as mutation, fusion, or amplification. The fibroblast growth factor-3 gene (FGFR3) has long been associated with bladder oncogenesis, and FGFR3 inhibition is becoming an interesting treatment approach for selected patients.

In April 2019, the US FDA approved erdafitinib for locally advanced or metastatic platinum-refractory UC with at least one FGFR3 alteration or FGFR2/3-containing fusion. BLC2001, an open-label phase II clinical trial, enrolled 99 patients with prespecified FGFR2/3 alterations which progressed during or following at least one line of prior chemotheraphy for locally advanced UC or unresectable/metastatic UC. After a median follow-up of 11.2 months, 40% of patients had an objective response to erdafitinib. A higher ORR (59%) was observed among patients who had previously received ICIs. With a median follow-up of 24 months, the mPFS was 5.52 months (95% CI 4.2–6.0) and the mOS was 11.3 months (95% CI 9.8 to not reached).

Other FGFR inhibitors, including rogaratinib, infigratinib, and pemigatinib, were tested in phase I and II studies and showed an ORR of around 25%. The results of a phase II study of vofatamab, a selective inhibitor of FGFR3, as salvage therapy in mUC, have recently been presented, with an ORR of 10% in monotherapy and 33% in combination with docetaxel. In the phase I/II FIERCE-22 study, the combination of vofatamab and pembrolizumab in previously pretreated mUC with and without FGFR3 alterations resulted in an ORR of 33% in wild-type tumors and 43% in tumors with FGFR3 alterations. Phase Ib/II trials with combinations of erdafitinib/certrelimab (NCT03473743), pemigatinib/pembrolizumab (NCT04003610), and rogaratinib/atezolizumab (NCT03473756) are currently ongoing.

In addition, in this landscape even angiogenic agents have been investigated in UC, based on encouraging results of phase II trials; however, the two phase III trials did not confirm the preliminary results when used alone or in combination with chemotherapy.

The phase III trial CALGB90601 compared bevacizumab, a monoclonal antibody against vascular endothelial growth factor-A (VEGF-A), plus chemotherapy (gemcitabine and cisplatin, GC) versus placebo plus chemotherapy as first-line therapy. The addition of bevacizumab to chemotherapy did not result in improved OS (14.5 vs 14.3 months; HR=0.87, 95% CI: 0.71–1.07).

### Table 2 Phase II/III Trials of Second-Line Targeted Therapy

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<th>Study (Phase)</th>
<th>Treatment (No. of Patients)</th>
<th>mPFS (Months)</th>
<th>mOS (Months)</th>
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**Abbreviations:** mPFS, median progression-free survival; mOS, median overall survival; ORR, objective response rate; AE, adverse event; ICS, immune cell score.
CI 0.72–1.05; two-sided stratified log-rank \( p = 0.14 \). The phase III RANGE evaluated ramucirumab, a monoclonal antibody to VEGFR-2, plus docetaxel versus placebo plus docetaxel in patients with locally advanced or metastatic UC who progressed on platinum-based chemotherapy. Ramucirumab plus docetaxel improved PFS (4.1 months, 95% CI 3.3–4.8, vs 2.8 months, 95% CI 2.6–2.9; HR=0.696, 95% CI 0.573–0.845; \( p = 0.0002 \)), without a significant improvement in OS (9.4 months; 95% CI 7.9–11.4, in the ramucirumab group versus 7.9 months, 95% CI 7.0–9.3, in the placebo group; stratified HR=0.887, 95% CI 0.724–1.086; \( p = 0.25 \)), for patients with platinum-refractory advanced UC.\(^3\)

In conclusion, although the combinations of anti-VEGFR and chemotherapy showed a prolonged PFS, the inhibition of VEGFR signaling was not effective in improving OS in patients with mUC.

**Beyond Second-Line Treatment**

Despite the recent advances, both chemotherapy and immunotherapy have had limited success, with long-term response rates of 20–30% in patients with mUC. Up to 70–80% of patients experience disease progression or relapse after chemotherapy and immunotherapy. Therefore, there is an urgent need for effective new agents to overcome resistance to cancer treatment and delay the relapse of the tumor. Antibody–drug conjugate (ADC) agents are a promising new therapeutic option that utilizes antibody-mediated delivery of a potent cytotoxic payload directly to the cancer cell, and represents a shift in the treatment of advanced UC (Table 3).

The first ADC drug approved by the FDA was enfortumab vedotinib (EV), a humanized monoclonal antibody linked to the chemotoxic microtubule-disrupting agent monomethyl auristatin E (MMAE), which targets nectin-4, a transmembrane protein widely expressed and associated with poor prognosis in mUC tumors.

After the phase I/II study yielded promising results, the phase III study EV-301 compared EV to investigator-chosen chemotherapy (docetaxel, paclitaxel, or vinflunine) in patients with locally advanced or metastatic UC who had previously been treated with ICI and platinum-based chemotherapy. This trial was stopped early owing to positive interim analysis. The results demonstrated a significant improvement in mOS, which was 12.88 months (95% CI 8.05–10.74) with EV versus 8.97 months (95% CI 8.05–10.74) with chemotherapy; mPFS was 5.55 months (95% CI 5.32–5.82) with EV versus 3.71 months (95% CI 3.52–3.94) with chemotherapy (HR=0.62, 95% CI 0.51–0.75; \( p = 0.001 \); ORR was 40.6% with EV versus 17.9% with chemotherapy (\( p = 0.001 \)); and the median DoR to EV was 7.39 months.\(^4\) The most common side effects were rash, peripheral neuropathy, fatigue, and alopecia. It is important to note the ocular toxicity and hyperglycemia.

These promising data and preliminary results of synergic effects when combined with immunotherapy have led to research evaluating the efficacy of EV in other earlier treatment scenarios, with two studies currently underway: the EV302 study is evaluating EV in combination with pembrolizumab in the first-line setting and the KEYNOTE-915 study is evaluating the same combination in the perioperative setting.

Other exciting ADCs are currently under development are sacituzumab govitecan (a conjugate formed by an antibody against trop2 and SN-38 and an active metabolite of irinotecan), RC48-ADC (a conjugate formed by an antibody against HER-2 and MMAE), and trastuzumab deruxtecan (a conjugate formed by an antibody against HER-2 and a topoisomerase-1 inhibitor). The response rates observed with these ADCs in phase I trials are extremely promising.

Intriguing new results have been reported from phase I/II data on sacituzumab, which showed an ORR of 31% in 45 patients with advanced UC.\(^4\) A phase II study (NCT03547973) is currently ongoing.

**Conclusions**

The therapeutic landscape of advanced urothelial carcinoma has been enriched with additional classes of agents.

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**Table 3: Phase II/III Trials with Targeted Therapies Beyond Second-Line Treatment**

<table>
<thead>
<tr>
<th>Study (Phase)</th>
<th>Treatment (No. of Patients)</th>
<th>mPFS (Months)</th>
<th>mOS (Months)</th>
<th>ORR (%)</th>
<th>G3–4 Treatment-Related AEs</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Beyond second line</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>EV-301 (phase III)</td>
<td>Enfortumab vedotinib (301)</td>
<td>5.55</td>
<td>12.88</td>
<td>40.6</td>
<td>51.4</td>
</tr>
<tr>
<td></td>
<td>Docetaxel/paclitaxel/vinflunine (301)</td>
<td>3.71</td>
<td>8.97</td>
<td>17.9</td>
<td>49.8</td>
</tr>
<tr>
<td>THROPY-U-01 (phase II)</td>
<td>Sacituzumab (113)</td>
<td>5.4</td>
<td>10.9</td>
<td>27</td>
<td></td>
</tr>
</tbody>
</table>

**Abbreviations:** mPFS, median progression-free survival; mOS, median overall survival; ORR, objective response rate; AE, adverse event.
Cisplatin-based chemotherapy remains the first-line standard treatment for cisplatin-eligible patients. For cisplatin-ineligible patients, ICIs can be used in PD-L1-high patients.

For second-line treatment, among the FDA-approved ICIs, pembrolizumab is the only one with an OS benefit observed in a randomized phase III trial (KEYNOTE-045).

The ORR of targeted therapies is significantly higher than for cytotoxic drugs in patients refractory to platinum-based chemotherapy.

For patients with actionable FGFR2 or FGFR3 genomic alterations, erdafitinib is the first-choice treatment option after platinum-based treatment.

The results (benefit in OS, PFS, and ORR) obtained with EV in patients who failed both platinum and ICI have never been achieved before.

Since the therapeutic armamentarium for advanced urothelial carcinoma is finally being enriched, we need to identify the clinical and molecular factors that can help to select the optimal treatment sequence for a specific patient.

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References


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