Systemic treatment for advanced urothelial cancer: an update on recent clinical trials and current treatment options

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After cisplatin-based chemotherapy became the standard treatment for metastatic urothelial cancer (mUC), very little progress has been made in the treatment landscape of this condition until recently. With increased knowledge about the molecular biology of mUC and advances in the field of cancer immunobiology, there has been an explosion in the number of clinical trials for mUC, and systemic treatment of mUC is rapidly changing. Despite the availability of several novel therapeutic agents, cisplatin-based cytotoxic chemotherapy remains the standard, first-line treatment option. Immune checkpoint inhibitors (ICIs), including programmed death-1 and programmed death ligand-1 inhibitors, are preferred second-line treatment options that are also used in first-line cisplatin-ineligible settings. For patients with actionable fibroblast growth factor receptor 2 (FGFR2) or FGFR3 genomic alterations, erdafitinib can be considered after platinum-based treatment. Enfortumab vedotin, a monoclonal antibody targeting nectin-4 conjugated to monomethyl auristatin E, has been approved for patients who do not respond to both cytotoxic chemotherapy and ICIs. In this review, we address the clinical trial data that have established the current standard treatments and ongoing clinical trials of various agents with different mechanisms as well as provide a brief overview of current practice guidelines and recommendations in patients with mUC.

Keywords: Carcinoma, transitional cell; Therapeutics; Drug therapy; Immunotherapy

INTRODUCTION

Bladder cancer is the 10th most common cancer worldwide, with an estimated 549,000 new cases and 200,000 deaths [1]. Urothelial cancer (UC) is the most common histology among cancers arising from the epithelium lining the urinary tract, any part of the renal pelvis, ureter, urinary bladder, and urethra [2]. Approximately 75% of patients with urothelial bladder cancer are diagnosed with non-muscle-invasive bladder cancer (NMIBC), which is usually managed by repeated local treatment with surveillance and is considered non-fatal. However, 25% of patients with bladder cancer have muscle-invasive bladder cancer (MIBC), which requires a multidisciplinary approach, including radical cystectomy and perioperative chemotherapy, and some of them have locally advanced unresectable, recurrent, or metastatic disease [3]. The prognosis of metastatic urothelial cancer (mUC) is poor, with a median overall survival (mOS) of approximately 15 months in large randomized trials [4,5];
the 5-year survival rates remained stable at approximately 10% during 1973 to 2005 [6].

Until mid-2010, there had been little progress in systemic treatments for mUC because of the wide usage of platinum-based combination chemotherapies in metastatic disease since the 1980s. However, mUC is one of the most rapidly progressing fields in medical oncology, with the recognition of actionable molecular alterations and introduction of novel agents, including immune checkpoint inhibitors (ICIs), molecularly targeted agents (MTAs), and antibody-drug conjugates (ADCs). In this review, we address the clinical trial data that have established the current standard treatments and ongoing clinical trials of various agents with different mechanisms, and also provide a brief overview of current practice guidelines and recommendations in patients with mUC.

**CYTOTOXIC CHEMOTHERAPY**

**First-line cisplatin-based chemotherapy**

Cytotoxic chemotherapy has been the mainstay of systemic treatment for mUC, and objective response rates (ORRs) to contemporary first-line combination cytotoxic chemotherapies range from 50% to 65% with complete response (CR) rates of 10% to 20% [7,8]. Among various chemotherapeutic agents, cisplatin has been the backbone of treatment, based on ORRs of approximately 33% as a single agent [9,10]. Thereafter, combination regimens containing cisplatin were developed. In a phase II trial, methotrexate, vinblastine, adriamycin, and cisplatin (MVAC) showed a promising response rate of 70% with an mOS of 13 months [11]. Subsequently, in two randomized phase III trials, MVAC outperformed cisplatin monotherapy [12] as well as cisplatin, cyclophosphamide, and adriamycin combination [13] and became a standard treatment for mUC. Despite MVAC being superior to other agents, it is associated with severe toxicities, including grade 3 or 4 leukopenia, thrombocytopenia, febrile neutropenia, sepsis, mucositis, nausea/vomiting, renal toxicity, hepatic toxicity, and therapy-related deaths [11,12]. Therefore, a less toxic regimen with similar or better efficacy was needed.

Gemcitabine demonstrated its efficacy in phase II trials as a second-line monotherapy and a first-line combination with cisplatin [14,15]. In a randomized phase III comparing gemcitabine plus cisplatin (GP) to MVAC, GP achieved comparable ORR (49% for GP vs. 46% for MVAC), median progression-free survival (mPFS, 7.7 months vs. 8.3 months), and mOS (14.0 months vs. 15.2 months) [5,8]. Although the study failed to prove a statistical difference in overall survival (OS), the primary endpoint, GP became the preferred first-line chemotherapy owing to its favorable toxicity profile. A 4-week schedule of GP was employed in the trial, but a 3-week schedule is widely used because of similar dose intensity and response rates with better compliance profile [16,17].

Clinical trials continued to improve the efficacy of cytotoxic chemotherapy. As taxanes showed moderate efficacy as a single agent [18,19], they were evaluated in combination with cisplatin in randomized phase III trials (docetaxel plus cisplatin vs. MVAC [20] and larotaxel plus cisplatin vs. GP [21]), but failed to prove their superior efficacy. In a randomized phase III trial EORTC 30987, a combination of paclitaxel and GP was compared to GP in patients with locally advanced or metastatic UC. Even though the triplet group had a higher ORR (55.5% vs. 43.6%, p = 0.031), the study failed to show a statistically significant difference in OS, the primary endpoint [22].

Intensification of the standard treatments was also evaluated. The EORTC 30924 trial was a phase III randomized controlled trial that compared dose-dense MVAC (ddMVAC) plus prophylactic granulocyte colony-stimulating factor (G-CSF) with classic MVAC [4,7]. Although ddMVAC did not achieve a statistically significant difference in mOS (15.1 months vs. 14.9 months, p = 0.049), ddMVAC showed higher ORR (64% vs. 50%) and better long-term survival (22% vs. 14% of 5-year OS rate) with a more favorable toxicity profile [4,7]. Therefore, ddMVAC is recommended as a viable option in the treatment of mUC in guidelines [23-25]. Until now, despite considerable efforts, MVAC and GP are the standard first-line treatments for mUC for several decades.

**First-line chemotherapy in cisplatin-unfit patients**

Carboplatin is another platinum chemotherapeutic agent, which is devoid of nephrotoxicity, less emetogenic, and less neurotoxic [26]. There have been no adequately powered randomized trials comparing cisplatin- and carboplatin-based chemotherapy in mUC. However, some small studies and meta-analyses suggested that cisplatin-based chemotherapy significantly
increases the likelihood of achieving a response in mUC [27-29]. Therefore, cisplatin-based chemotherapy is recommended as the first-line treatment in current guidelines, while carboplatin is considered only when patients cannot tolerate cisplatin [23-25]. According to the expert consensus definition, cisplatin “unfit” is defined as the presence of at least one of the Eastern Cooperative Oncology Group performance status 2, creatinine clearance < 60 mL/min, peripheral neuropathy ≥ CTCAE v4.0 grade 2, hearing loss ≥ CTCAE v4.0 grade 2, or New York Heart Association class III heart failure [30]. Approximately 30% to 50% of patients with mUC are ineligible to cisplatin in the clinical practice [31]. For platinum unfit patients with mUC, large randomized phase III trials are scarce. EORTC 30986 is the only phase III randomized controlled trial in a platinum unfit population, which compared gemcitabine and carboplatin (GCb) versus methotrexate/carboplatin/vinblastine [32]. The ORR was higher in the GCb arm (41.2% vs. 30.3%), but OS and PFS were not significantly different between the two regimens. Although EORTC 30986 failed to prove the superiority of GCb, considering the lower incidence of severe acute toxicity in the GCb arm, GCb became the de facto standard treatment for platinum unfit mUC. Some other doublet combinations were evaluated in phase II trials [33-35], but evidence is insufficient to recommend any specific regimen in this population. Notably, in the COACH trial, gemcitabine plus oxaliplatin maintained its efficacy in patients with very poor renal function (chronic kidney disease stage 4–5), contrary to GCb [34]. Since renal impairment is the most common cause of cisplatin-ineligibility, additional investigation is needed to confirm this finding.

### Salvage chemotherapy after failure of first-line treatment

Although the first-line cisplatin-based chemotherapy has high ORRs, the response does not last long with an mPFS of 7 to 8 months [4,5]. For second-line therapy, many drugs were tested in phase II trials [36-47], but their efficacy was modest, with an ORR of 5-20%.

### Table 1. Food and Drug Administration-approved immune checkpoint inhibitors for patients with metastatic urothelial cancer in a platinum-refractory setting

| Study (phase) | Pembrolizumab | Atezolizumab | Nivolumab | Durvalumab | Avelumab |
|---------------|---------------|--------------|-----------|------------|----------|
| Number        | 270           | 497          | 265       | 191        | 249      |
| Dosing schedule | 200 mg q3 wk | 1,200 mg q3 wk | 3 mg/kg q2 wk | 10 mg/kg q2 wk | 10 mg/kg q2 wk |
| PD-L1 IHC assay | 22C3 pharmDx on DAKO, IC | SP142 on VENTANA, TC | 28-8 pharmDx on DAKO, TC or IC | SP263 on VENTANA, in tumor cells | 73-10 pharmDx on DAKO, in tumor cells |
| ORR (CR/PR) | 21.1% (9.3%/11.9%) | 13.4% (3%/10%) | 19.6% (2%/17%) | 17.8% (3.7%/14.1%) | 17% (6%/11%) |
| TTR | 2.1 mon | Not reported | Not reported | 1.41 mon | 11.4 wk |
| DOR | Not reached | 21.7 mon | Not reached | Not reached | Not reached |
| Median OS | 10.1 mon | 8.6 mon | 8.7 mon | 18.2 mon | 6.5 mon |
| Median PFS | 2.1 mon | 2.1 mon | 2.0 mon | 1.5 mon | 6.3 wk |
| TRAEs, any grade | 62.6%/16.5% | 69%/20% | 64%/18% | 60.7%/6.8% | 67%/8% |

PD-L1, programmed death ligand 1; IHC, immunohistochemical staining; CPS, combined positive score; IC, immune cell; TC, tumor cell; ORR, objective response rate; CR, complete response; PR, partial response; TTR, time to response; DOR, duration of response; OS, overall survival; PFS, progression-free survival; TRAE, treatment-related adverse event.
tic agent that has been tested in a randomized phase III trial. In that trial, vinflunine did not reach statistical significance in improving mOS compared to BSC in the intention-to-treat (ITT) analysis (6.9 months vs. 43 months), although the use of vinflunine was independently correlated with improved survival in multivariable analysis [48,49]. Until mid-2010, there was no standard salvage treatment, and treatment guidelines recommended various agents such as taxanes (either paclitaxel or docetaxel), vinflunine, or sometimes pemetrexed [23-25].

**IMMUNE CHECKPOINT INHIBITORS**

**ICIs in platinum-pretreated population**

There are five U.S. Food and Drug Administration (FDA)-approved anti-programmed death 1 (PD-1)/programmed death ligand 1 (PD-L1) antibodies, nivolumab, pembrolizumab, durvalumab, avelumab, and atezolizumab, for mUC (Table 1) [50-54]. Among these drugs, pembrolizumab (KEYNOTE-045) and atezolizumab (IMvigor211) underwent phase III randomized controlled trials in a salvage setting [50,51]. In contrast, others have only phase I or II trial results.

In KEYNOTE-045, the efficacy of anti-PD-1 antibody pembrolizumab as a salvage treatment was compared with the investigator’s choice drug (paclitaxel, docetaxel, or vinflunine) in patients with mUC who had disease progression after platinum-based chemotherapy [50]. Co-primary endpoints were OS and PFS in the total population and in the patients with PD-L1 expressing tumors according to two thresholds (combined positive score [CPS] ≥ 1 and ≥ 10). A total of 542 patients were randomly assigned to the pembrolizumab and chemotherapy groups, and in the second interim analysis, co-primary endpoints were met. The mOS was 13.0 months with pembrolizumab and 7.4 months with chemotherapy (hazard ratio [HR], 0.73; \( p = 0.002 \)). There was no significant difference in mPFS (2.1 months vs. 3.3 months). ORR was significantly higher in pembrolizumab (21.1% vs. 11.4%, \( p = 0.001 \)). Long-term efficacy and safety results were consistent with the interim results [55].

IMvigor211 was a randomized phase III trial comparing atezolizumab, an anti-PD-L1 antibody with the investigator’s choice drug (paclitaxel, docetaxel, or vinflunine) [51]. Eligibility criteria were generally similar to those of KEYNOTE-045. The primary endpoint was OS tested hierarchically in prespecified populations; PD-L1 expression on ≥ 5% of tumor-infiltrating immune cells (IC2/3), followed by IC3/2 (PD-L1 expression on ≥ 1% of immune cells), followed by the ITT population. A total of 931 patients were randomized to the atezolizumab or chemotherapy arm. In the IC2/3 population (\( n = 234 \)), mOS did not differ significantly between patients in the atezolizumab group and in the chemotherapy group (11.1 months vs. 10.6 months; HR, 0.87; \( p = 0.41 \)), precluding further formal statistical comparisons. The ORR was similar between treatment arms (23.0% vs. 21.6% in IC2/3 population and 13.4% vs. 13.4% in ITT population), while mPFS was numerically shorter in the atezolizumab arm (2.4 months vs. 4.2 months in the IC2/3 population and 2.1 months vs. 4.0 months in the ITT population). Therefore, in contrast to the KEYNOTE-045 trial, IMvigor211 failed to prove the efficacy of atezolizumab over cytotoxic chemotherapy. However, in exploratory analysis of the ITT population of IMvigor211, the atezolizumab arm had a numerically improved OS compared to the chemotherapy arm. The mOS was 8.6 months versus 8.0 months, and the 1-year OS rate was 39.2% versus 32.4%, with an HR, 0.85 (95% confidence interval [CI], 0.73 to 0.99). The atezolizumab arm maintained the improved OS in an updated analysis, demonstrating a 2-year OS rate of 23% versus 13% and a 3-year OS rate of 18% versus 10% with an HR of 0.82 (95% CI, 0.71 to 0.94) [56]. This unexpected finding gives us a lesson that design and statistical analysis plan are important in clinical trials, especially when biomarkers, which have not been fully understood, are incorporated into trials. Based on these results, pembrolizumab became the standard treatment for platinum-pretreated mUC. However, as there is no clinical trial conducting head-to-head comparison of various ICIs in mUC, it is not evident whether pembrolizumab is superior to other ICIs. Considering real-world outcomes of ICIs were similar to those of KEYNOTE-045 or IMvigor211 [57,58], atezolizumab or other ICIs can also be used in platinum-pretreated populations [24].

The incidence of treatment-related adverse events (AEs) of ICIs was lower than that of cytotoxic chemotherapy. For instance, any grade and grade 3-5 AEs occurred in 60.9% and 15.0%, respectively, of patients in the pembrolizumab arm, compared to 90.2% and 49.4%,
respectively, of patients in the chemotherapy arm in the KEYNOTE-045 trial [50]. Likewise, any grade and grade 3–5 AEs were reported in 69% and 20%, respectively, of patients in the atezolizumab arm, compared to 89% and
43%, respectively, of patients in the chemotherapy arm in the IMvigor211 trial [52]. The most common AEs for ICIs are pruritus, fatigue, nausea, diarrhea, decreased appetite, and rash. Although not typical, there are immune-related AEs that could sometimes be fatal. Thus, physicians need to be aware of the immune-related AEs and management [59,60]. In addition, there are distinct radiologic response and progression patterns of ICIs, such as pseudoprogression and hyperprogression [61]. The exact incidences of these phenomena in UC are not known, but these might be not uncommon, as 1.5% to 17% of pseudoprogression and 12% of hyperprogression are reported in mUC [62,63].

Maintenance ICI after first-line chemotherapy

Previous studies have investigated the efficacy of ICIs in mUC patients who experienced disease progression during or after platinum-based chemotherapy. In contrast, a phase III randomized trial, JAVELIN Bladder 100 trial (NCT02603432) investigated maintenance treatment with avelumab plus BSC versus BSC alone in patients with mUC whose disease did not progress after completion of first-line platinum-containing chemotherapy, and primary analysis results were presented at the American Society of Clinical Oncology meeting in 2020 [64]. A total of 700 patients were randomized, and a statistically significant improvement in OS was demonstrated in the avelumab arm in both ITT group (21.4 months vs. 14.3 months; stratified HR, 0.69 [95% CI, 0.56 to 0.86]; p < 0.001) and PD-L1 positive group (not reached vs. 17.1 months; stratified HR, 0.56 [95% CI, 0.40 to 0.79]; p < 0.001) [64]. Both PFS and ORR were also superior in the avelumab arm. Based on these results, avelumab maintenance in mUC patients whose disease has not progressed with first-line platinum-based chemotherapy would be a new standard of care.

First-line ICI in cisplatin-unfit patients

In cisplatin-unfit patients, GCb has been a de facto standard treatment after the EORTC 30986 trial [32], but there is an urgent need to improve treatment in this population. In IMvigor 210 (cohort 1) and KEYNOTE-052 trials, atezolizumab and pembrolizumab were tested as monotherapy in cisplatin-unfit chemotherapy-naïve patients with mUC [65-67]. In the IMvigor 210 trial, out of 123 patients who received atezolizumab, ORR, mPFS, and mOS were 23%, 2.7 months, and 15.9 months, respectively [65]. In the KEYNOTE-052 trial, out of 370 patients, ORR, mPFS, and mOS were 29%, 2.2 months, and 11.3 months, respectively [66,67]. Notably, patients with PD-L1 CPS ≥ 10 achieved more favorable outcomes with pembrolizumab compared to CPS < 10, with an ORR of 47.3% versus 20.3% and an mOS of 18.5 months versus 9.7 months in KEYNOTE-052. Based on these results, both atezolizumab and pembrolizumab were granted accelerated approval by the U.S. FDA.

For continued approval, confirmatory data in phase III trials are required from IMvigor 130 (NCT02807636) and KEYNOTE-361 (NCT02853305) trials. In the preliminary analyses of both trials, in patients with low PD-L1 expression, ICI arms had decreased survival compared to cisplatin- or carboplatin-based therapy. As a result, both trials stopped enrollment of patients with low PD-L1 status into monotherapy arms, and the indication for both agents was modified to include only patients who were not eligible for cisplatin-containing chemotherapy and who had high expression of PD-L1 or were not eligible for any platinum-containing chemotherapy regardless of the level of PD-L1 expression [68]. In the interim OS results of the IMvigor130 trial [69], although atezolizumab monotherapy group had numerically longer mOS than the chemotherapy group (15.7 months vs. 13.1 months), it was difficult to draw any conclusion because the stratified HR of 1.02 (95% CI, 0.83 to 1.24) and crossing survival curves suggested heterogeneous treatment effect across subgroups. Furthermore, formal statistical testing for comparing atezolizumab monotherapy with chemotherapy was not performed because of the hierarchical statistical design of the trial.

Ongoing first-line combination ICI trials

There are multiple ongoing clinical trials in the first-line setting, investigating whether the combination of ICIs with cytotoxic chemotherapy is superior to chemotherapy alone or ICI alone (Fig. 1). In addition, a clinical trial is evaluating first-line pembrolizumab in combination with lenvatinib versus pembrolizumab plus placebo in cisplatin-ineligible patients with CPS ≥ 10 or platinum-ineligible patients (LEAP-011 trial, NCT03898180). Among these trials, the final PFS and interim OS results of the IMvigor130 trial have been published [69]. After a median follow-up of 11.8 months, the mPFS was sig-
significantly superior in the combination arm: 8.2 months in the chemotherapy plus atezolizumab arm versus 6.3 months in the chemotherapy plus placebo arm (HR, 0.82; 95% CI, 0.70 to 0.96; \( p = 0.007 \)). mOS was 16.0 months versus 13.4 months, with a stratified HR of 0.83 (95% CI, 0.69 to 1.00) and a one-sided \( p \) value of 0.027. As the \( p \) value did not exceed the prespecified interim efficacy boundary, further follow-up for mature OS data is required. In addition, in the DANUBE trial, the primary endpoints were not attained [70]. A full report of the DANUBE trial has not been published yet.

**MOLECULARLY TARGETED AGENTS**

Owing to next-generation sequencing and bioinformatics, genomic landscape and actionable mutations in UC have been unveiled [71-74]. According to the updated results of The Cancer Genome Atlas (TCGA) report [74], the most common mutations in MIBC are TP53, PIK3CA, CDKN1A, ERCC2, fibroblast growth factor receptor 3 (FGFR3), and ERBB3. Most recently conducted trials have focused on actionable mutations among the aforementioned alterations.

**Targeting fibroblast growth factor receptor**

FGFR3 alteration is a common event in UC, with a spectrum of mutations, including point mutation and fusion, having been reported [75]. Although FGFR3 alterations are more common in NMIBC (up to 80% in Ta and 30% in T1) [76], 12% to 15% of MIBC patients have FGFR3 alterations [71-74]. Activating point mutations in exons 7, 10, and 15 (S249C, R248C, and Y373C [Y375C]) are the most common FGFR3 alterations, and less commonly, gene fusions (FGFR3-TACC3, FGFR3-BAIAP2L1, and FGFR3-JAKMIP1) have also been observed [75]. In addition to mutations, FGFR3 protein or mRNA overexpression is also present in MIBC [77], probably by epigenetic regulation [78]. As FGFR3 alteration plays an important oncogenic role in UC [75], FGFR3 signaling is an attractive target, and many drugs have been tested in UC [79-85]. The outcomes of select trials of FGFR inhibitors in mUC are described in Table 2, and ongoing trials of MTAs are listed in Table 3.

Erdafitinib is a potent tyrosine kinase inhibitor (TKI) of FGFR1–4. In the BLC2001 trial, a phase II trial evalu-
| Drug | Clinical trial, phase | Treatment arm(s) | Main inclusion criteria |
|------|----------------------|------------------|------------------------|
| FGFR inhibitor | | | |
| **Erdafitinib** | THOR (NCT03939504), phase III | Erdafitinib vs. chemotherapy or pembrolizumab | Platinum-treated with/without ICI-treated mUC with FGFR mutation or fusions/translocations |
| | FIGHT-201 (NCT02837741), phase II | Erdafitinib |铂金治疗，与/无ICI治疗的mUC与FGFR突变或融合/移位 |
| | FUZE (NCT03413220), phase I | Pembrolizumab | Pembrolizumab |
| | Delibl147 | Pembrolinib | Pembrolinib |
| **Pemigatinib** | FIGHT-25 (NCT0409610), phase II | Pemigatinib vs. pembrolizumab | Platinum-treated or unfit mUC with FGFR3 mutation/alteration |
| | NORSE (NCT03473743), phase Ib/II | Pemigatinib | Pemigatinib or pembrolizumab |
| | FUZE (NCT03413220), phase I | Pembrolizumab | Pembrolizumab |
| **Debio1347** | FUZE (NCT03834220), phase II | Debio1347 | Treatment refractory solid tumors harboring FGFR3 fusions/ translocations |
| **FGFR inhibitor in combination with ICI** | | | |
| **Rogaratinib** | FORT-2 (NCT03473756), phase Ib/II | Rogaratinib + atezolizumab vs. placebo + atezolizumab | Treatment-naïve cisplatin-unfit mUC with high FGFR3 mutation/fusion |
| | FORT-2 (NCT03473756), phase II | Rogaratinib | Treatment-naïve unfract mUC with high FGFR3 mutation/fusion |
| | FUZE (NCT03413220), phase I | Pembrolizumab | Pembrolizumab |
| **Erdafitinib + atezolizumab** | NORSE (NCT03473743), phase Ib/II | Erdafitinib + atezolizumab | Treatment-naïve cisplatin-unfit mUC with high FGFR3 mutation/fusion |
| | FIGHT-201 (NCT02837741), phase II | Erdafitinib | Erdafitinib |
| **Pemigatinib + pembrolizumab** | NORSE (NCT03473743), phase Ib/II | Pemigatinib + pembrolizumab | Treatment-naïve cisplatin-unfit mUC with high FGFR3 mutation/fusion |
| | FIGHT-201 (NCT02837741), phase II | Pemigatinib | Pemigatinib |
| | FUZE (NCT03413220), phase I | Pembrolizumab | Pembrolizumab |
| **Debio1347 + pembrolizumab** | NORSE (NCT03473743), phase Ib/II | Debio1347 + pembrolizumab | Treatment-naïve cisplatin-unfit mUC with high FGFR3 mutation/fusion |
| | FIGHT-201 (NCT02837741), phase II | Debio1347 | Debio1347 |
| | FUZE (NCT03413220), phase I | Pembrolizumab | Pembrolizumab |
| **Afatinib** | LUX-Bladder1 (NCT02780687), phase II | Afatinib | Treatment-refractory solid tumors with FGFR3 fusions/alterations (cohort B) |
| | LUX-Bladder1 (NCT02780687), phase I | Afatinib | Afatinib |
| **Pertuzumab/trastuzumab** | MyPathway (NCT02091149), phase II basket | Pertuzumab + trastuzumab | Treatment-refractory solid tumors with HER2 amplification (by NGS, FISH or CISH) and/or HER2 actionable mutation (cohort B) |
| | MyPathway (NCT02091149), phase I | Pertuzumab | Pertuzumab |
| **Lenvatinib + pembrolizumab** | PembroCl (NCT03833842), phase II | Lenvatinib + pembrolizumab | Treatment-refractory solid tumors with HER2 amplification (by NGS, FISH or CISH) and/or HER2 actionable mutation (cohort B) |
| | PembroCl (NCT03833842), phase I | Lenvatinib | Lenvatinib |
| **Cabozantinib** | ARCADIA (NCT03833842), phase I | Cabozantinib | Treatment-refractory solid tumors with HER2 amplification (by NGS, FISH or CISH) and/or HER2 actionable mutation (cohort B) |
| | ARCADIA (NCT03833842), phase I | Cabozantinib | Cabozantinib |

FGFR, fibroblast growth factor receptor; ICI, immune checkpoint inhibitor; mUC, metastatic urothelial cancer; HER, human epidermal growth factor receptor; EGFR, epidermal growth factor receptor; NGS, next generation sequencing; FISH, fluorescent in situ hybridization; CISH, chromogenic in situ hybridization; VEGF, vascular endothelial growth factor; PD-L1, programmed death ligand-1.
The ErbB family of interest in mUC includes HER-directed therapy with pertuzumab plus trastuzumab for patients with mUC with HER2 amplification (by next-generation sequencing, fluorescence in situ hybridization, or chromogenic in situ hybridization) and/or immunohistochemical (IHC)3+ and/or HER2 actionable mutation (Table 3) [102].

Targeting vascular endothelial growth factor signaling
Bladder cancer produces pro-angiogenic factors, including vascular endothelial growth factor (VEGF), and high expression of pro-angiogenic factors were found to be correlated with disease progression and poor survival [103]. Several VEGF receptor TKIs have been investigated in mUC, as a monotherapy in phase II trials [104-109] or as a combination with cytotoxic chemotherapy [110-112]. All these agents were not investigated further owing to a lack of efficacy and/or excessive toxicities.

Monoclonal antibodies targeting VEGF signaling were also tested. Ramucirumab, a monoclonal antibody to VEGFR-2, was evaluated in a phase III randomized controlled trial (RANGE) comparing the efficacy of ramucirumab plus docetaxel with placebo plus docetaxel in previously treated mUC [113,114]. Bevacizumab, a monoclonal antibody to VEGF-A, was investigated in a phase III randomized controlled trial (CALGB90601) comparing the efficacy of GP plus bevacizumab with GP plus placebo in chemotherapy-naïve patients with mUC [115]. Both trials showed that monoclonal VEGF antibodies combined with cytotoxic chemotherapy failed to improve OS in patients with mUC, although the combinations prolonged PFS.

Besides the antiangiogenic effect, there is preclin-
ical evidence that VEGF inhibition also facilitates anti-tumor immunity [116]. VEGF inhibition enhances T cell infiltration and activation and inhibits suppressive immune cells. Therefore, the combination of VEGF inhibitors and ICIs can be synergistic, and there are several ongoing clinical trials evaluating the efficacy of a combination of VEGF inhibitors and ICIs in mUC (Table 3).

**ANTIBODY-DRUG CONJUGATES**

ADCs are a novel class of drugs that are rationally designed to deliver effective cytotoxic drugs directly and selectively to cancer cells. ADCs comprise a monoclonal antibody that recognizes tumor-associated antigens and to which a potent cytotoxic agent is conjugated via chemical linkages [117]. There are several promising ADCs under evaluation for mUC.

**Enfortumab vedotin (ASG-22CE; ASG-22ME)**

Enfortumab vedotin is an ADC that comprises a fully human monoclonal antibody targeting nectin-4 conjugated to monomethyl auristatin E (MMAE) via a protease-cleavable linker [118]. Nectin-4 is a member of the transmembrane protein nectin family cell adhesion molecules that are involved in various cellular processes, including carcinogenesis [119]. Nectin-4 is highly expressed in various solid tumors including UC [120]. In a phase 2, open-label, single-arm study (EV-201), the efficacy of enfortumab vedotin was tested in patients with mUC who previously received an ICI with or without prior platinum chemotherapy. Results of patients who received both ICI and chemotherapy were published [118]. The ORR was 44% and CR was 12%; mPFS and mOS were 5.8 and 11.7 months, respectively. Treatment response was observed in all subgroups, including ICI non-responders and those with liver metastases. Common treatment-related AEs included fatigue (50%), peripheral neuropathy (50%), alopecia (49%), rash (48%), decreased appetite (44%), and dysgeusia (40%). Grade 3 or more AEs occurred in over 5% of patients, with only fatigue occurring in 6%. No fatal treatment-related AEs were reported. Based on these data, the FDA approved enfortumab vedotin for mUC following chemotherapy and ICI treatment. Currently, a randomized phase 3 trial (EV-301, NCT03474107) is ongoing to compare enfortumab vedotin with the investigator’s choice (docetaxel, paclitaxel, or vinflunine) in patients with mUC who showed disease progression with platinum-based chemotherapy and an ICI (Table 4).

Another ongoing study (EV-103, NCT03288545) is a phase 1b-2 trial evaluating enfortumab vedotin combined with pembrolizumab and/or chemotherapy in patients with metastatic UC in multiple cohorts. The preliminary results of cohort A, in which 45 cisplatin-ineligible patients were treated with enfortumab vedotin plus pembrolizumab showed that the ORR was 73.3% with 15.6% CR and the response was not associated with PD-L1 status [121]. Based on this promising result, a phase 3 study, EV-302 (NCT04228956) is designed and currently recruiting patients. This trial will evaluate first-line enfortumab vedotin in combination with pembrolizumab with or without chemotherapy versus chemotherapy in patients with advanced UC.

**Sacituzumab govitecan (IMMU-132)**

Sacituzumab govitecan is an ADC in which SN-38 (an active metabolite of irinotecan) is conjugated to the humanized anti-trophoblast cell-surface antigen 2 (Trop-2) monoclonal antibody via a cleavable linker. Trop-2, a transmembrane calcium signal transducer, is overexpressed in many epithelial cancers [122], and its expression correlates with disease severity in UC [123].

In a phase I/II basket trial, sacituzumab govitecan was assessed in a UC cohort that included patients with mUC who progressed after one or more prior systemic therapy [124]. In the cohort of 45 patients, the ORR was 31% with two patients with CR, and the mPFS and mOS were 7.3 and 18.9 months, respectively. Grade 3 or more AEs observed were neutropenia (38%), anemia (11%), hypophosphatemia (11%), diarrhea (9%), fatigue (9%), and febrile neutropenia (7%). TROPHY-U-01 (NCT03547973) is an open-label, single-arm, phase 2 trial evaluating sacituzumab govitecan in 140 patients with mUC who failed both platinum-based chemotherapy and ICI (cohort 1, 100 patients) or failed ICI in cisplatin-ineligible patients (cohort 2, 40 patients). In pre-planned interim analysis in cohort 1 (n = 35) [125], the ORR was 29%, which surpassed the prespecified futility endpoint; therefore, further enrollment is being continued (Table 4).
Other ADCs actively investigated in mUC are listed in Table 4. RC48-ADC is a novel humanized high-affinity anti-HER2 antibody hertuzumab conjugated with MMAE via a cleavable linker [126]. A phase II trial (NCT03507166) of HER2-overexpressed (IHC2+ or 3+) pretreated advanced UC completed accrual, and its result is awaited [127]. Trastuzumab deruxtecan (DS-8201a) is another trastuzumab-based ADC linked to deruxtecan, a derivative of topoisomerase I exatecan. The phase I/II trial combining DS-8201a with nivolumab in advanced HER2-expressing breast or UC is underway (NCT03523572) [128].

TREATMENT RECOMMENDATIONS AND SUGGESTED TREATMENT SEQUENCE

Before starting the systemic treatment of patients with UC, it is crucial to determine whether the patient is recurrent or metastatic UC. The median age of newly diagnosed UC patients is 73 years, and most patients are over 65 years [129]. For example, in an observational study of real-world practice involving elderly patients with newly diagnosed UC, 42% of patients received first-line chemotherapy and only 27% of first-line-treated patients received cisplatin-based chemotherapy [131]. In contrast, for a well-selected population, chemotherapy can be similarly effective between young and elderly patients [132]. Therefore, to avoid excessive toxicity and to select potential beneficiaries, a thorough examination, including geriatric assessment, is recommended for the elderly population [133]. Furthermore, patients with UC usually have multiple comorbidities. In one study, patients with bladder cancer had a median of 8 (interquartile range, 5 to 11) chronic conditions, including chronic kidney disease, coronary artery disease, and diabetes mellitus [34].

Table 4. Ongoing clinical trials of antibody-drug conjugates in patients with metastatic urothelial cancer

| ADC Name | Trial Design | Trial Identifier | Notes |
|----------|--------------|-----------------|-------|
| EV-301 (NCT03474107), phase III | Enfortumab vedotin vs. chemotherapy (docetaxel, paclitaxel, or vinflunine) | Prior platinum-treated and ICI-treated mUC |
| EV03 (NCT0328845) | Enfortumab vedotin ± pembrolizumab ± chemotherapy (cisplatin, carboplatin, or gemcitabine) | Treatment-naïve cisplatin eligible/ineligible or platinum in eligible mUC |
| EV-302 (NCT4233866), phase Ib-II | Enfortumab vedotin ± pembrolizumab vs. gemcitabine or carboplatin vs. enfortumab vedotin ± pembrolizumab ± cisplatin or carboplatin | Platinum-treated mUC |
| Sacituzumab govitecan (anti-Trop-2 antibody conjugated to SN-38) | Sacituzumab govitecan (cohort 1, 2) | Prior platinum-treated and ICI-treated (cohort 1) or treatment-naïve platinum-ineligible mUC (cohort 2) |
| RC48-ADC (anti-HER2 antibody hertuzumab conjugated to MMAE) | Failure of at least 1 systemic chemotherapy, HER-2 IHC2+ or 3+ |
| NCT04507166, phase II | RC48-ADC | Prior platinum-treated (cohort 3) |
| Trastuzumab deruxtecan (anti-HER2 antibody trastuzumab conjugated to deruxtecan) | Failure of platinum-based chemotherapy, HER2 IHC2+ or 3+ |
| NCT0353572, phase Ib | Trastuzumab deruxtecan ± nivolumab |

MMAE, monomethyl auristatin-E; ICI, immune checkpoint inhibitor; mUC, metastatic urothelial cancer; MIBC, muscle-invasive bladder cancer; HER2, human epidermal growth factor receptor 2; IHC, immunohistochemical staining.
morbidities are important in determining the “fitness” for cisplatin-based chemotherapy and seem to affect prognosis [135]. Besides age and comorbidities, socioeconomic status is another important factor affecting chemotherapy administration [136].

Although ICIs are generally well-tolerated and have more favorable toxicity profiles than cytotoxic chemotherapy, there are multiple conditions that need special concern, including autoimmune diseases, ongoing immunosuppressant use, or chronic viral infection. These populations can be cautiously treated with ICIs [137], but shared decision-making and monitoring is required.

**Prognostic factors for mUC**

Well-established prognostic models have essential roles in the management of patients with cancer; they enable physicians to predict life expectancy, guide treatment selection, analyze results of clinical studies, and educate patients and their families. There are several prognostic models available for mUC and are used in different clinical situations (Fig. 2).

For patients receiving first-line cisplatin-based chemotherapy, the Bajorin prognostic model can be applied [138]. This model has also been validated in patients receiving first-line carboplatin-based treatment [32]. Additionally, in platinum-refractory settings, the Bellmunt prognostic model can be used [139].

The above models are all based on data from clinical trials of cytotoxic chemotherapy; therefore, whether they can be applied for ICI is not certain. There is no well-validated prognostic model specific to ICI.

**First-line treatment for patients with mUC**

As of 2020, cisplatin-based combination chemotherapy remains the standard treatment for cisplatin-eligible patients (Fig. 3). Both GP and MVAC can be used,
and ddMVAC with prophylactic G-CSF can also be used [4,5,7,8]. All of these regimens are deemed to have similar efficacy, but their toxicity profiles differ from one another. For cisplatin-ineligible patients, there is no universally accepted standard treatment. GCb has been the most widely used regimen [32]. Considering longer survival in atezolizumab and pembrolizumab arms among cisplatin-ineligible patients compared to the historical control in IMvigor210 and KEYNOTE-052 trials [65,66], and numerically longer survival of atezolizumab monotherapy arm compared to the chemotherapy arm in interim results of the IMvigor130 trial [69], ICIs can also be used in only PD-L1 high patients [68].

When platinum cannot be used (platinum-ineligible), ICIs can be used regardless of PD-L1 expression, even though the criteria for “platinum-ineligibility” are not yet well defined [140]. If ICIs are not available or patients have conditions to avoid ICIs, single-agent gemcitabine can be considered, although there is no convincing evidence. If patients cannot tolerate any systemic treatment, offer BSC only.

**Second- or later-line treatment**

For patients who progressed after platinum-based chemotherapy, ICIs are the standard treatment option (Fig. 4) [50-54]. Among FDA-approved ICIs (atezolizumab, avelumab, durvalumab, nivolumab, and pembrolizumab), pembrolizumab is the preferred choice based on the KEYNOTE-045 trial [50]. For patients whose tumors have susceptible FGFR2 or 3 mutations, erdafitinib can be considered [80].

When patients progress after first-line ICI, cytotoxic chemotherapy can be administered. There are no prospective trial data in this situation, but platinum-based chemotherapies would be preferred, if we take account of their efficacy in a first-line setting. Similar to first-line setting, the regimen can be chosen according to cisplatin-eligibility.

For patients who failed both platinum and ICI, enfurterumb vedotin was proven effective [118]. Erdafitinib can also be used if there are susceptible FGFR alterations. Under certain conditions, when all available treatment fails or novel drugs are unavailable, classical cytotoxic chemotherapy (paclitaxel, docetaxel, pemetrexed, or vinflunine) might be considered for palliation.

At any point of time, if the general condition deteriorates or the patient seems not to tolerate systemic treatment, offer BSC only.

**CONCLUSIONS**

Systemic treatment of mUC has been stagnant for many decades, but revolutionary breakthroughs have now occurred. Guidelines have now included ICIs (atezolizumab, avelumab, durvalumab, nivolumab, and pembrolizumab), FGFR3 inhibitors (erdaftitinib), and ADCs (enfortumab vedotin) in treatment protocols. Furthermore, multiple clinical trials, including but not limited to those mentioned above, are ongoing. We believe that the prospect of the therapeutic landscape is promising with this armamentarium. However, great effort is needed to fill the gap between treatment guidelines and real-world practice, as patients with mUC are more likely to be elderly and comorbid than those with other cancers, and there are growing concerns about the affordability and availability of novel treatments.

**Conflict of interest**

No potential conflict of interest relevant to this article was reported.

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