FDA Approval Summary: Atezolizumab or Pembrolizumab for the Treatment of Patients with Advanced Urothelial Carcinoma Ineligible for Cisplatin-Containing Chemotherapy

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Disclosures of potential conflicts of interest may be found at the end of this article.

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ABSTRACT

The U.S. Food and Drug Administration (FDA) granted accelerated approval to atezolizumab and pembrolizumab in April and May 2017, respectively, for the treatment of patients with locally advanced or metastatic urothelial carcinoma who are not eligible for cisplatin-containing chemotherapy. These approvals were based on efficacy and safety data demonstrated in the two single-arm trials, IMvigor210 (atezolizumab) and KEYNOTE-052 (pembrolizumab). The primary endpoint, confirmed objective response rate, was 23.5% (95% confidence interval [CI]: 16.2%–32.2%) in patients receiving atezolizumab and 28.6% (95% CI: 24.1%–33.5%) in patients receiving pembrolizumab. The median duration of response was not reached in either study and responses were seen regardless of PD-L1 status. The safety profiles of both drugs were generally consistent with approved agents targeting PD-1/PD-L1. Two ongoing trials (IMvigor130 and KEYNOTE-361) are verifying benefit of these drugs. Based on concerning preliminary reports from these trials, FDA revised the indications for both agents in cisplatin-ineligible patients. Both drugs are now indicated for patients not eligible for any platinum-containing chemotherapy or not eligible for cisplatin-containing chemotherapy and whose tumors/infiltrating immune cells express a high level of PD-L1. The indications for atezolizumab and pembrolizumab in patients who have received prior platinum-based therapy have not been changed. This article summarizes the FDA thought process and data supporting the accelerated approval of both agents and the subsequent revision of the indications. The Oncologist 2018;23:1–7

Implications for Practice: The accelerated approvals of atezolizumab and pembrolizumab for cisplatin-ineligible patients with advanced urothelial carcinoma represent the first approved therapies for this patient population. These approvals were based on single-arm trials demonstrating reasonable objective response rates and favorable durations of response with an acceptable toxicity profile compared with available non-cisplatin-containing chemotherapy regimens. However, based on concerning preliminary reports from two ongoing phase III trials, the FDA revised the indication for both agents in cisplatin-ineligible patients. Both are now indicated either for patients not eligible for any platinum-containing chemotherapy or not eligible for cisplatin-containing chemotherapy and whose tumors have high expression of PD-L1.

INTRODUCTION

Urothelial carcinoma (UC) is the most common malignancy in the urinary tract and accounts for approximately 16,000 deaths annually in the U.S. [1, 2]. Although most UCs are non-muscle invasive at diagnosis and can be managed effectively with surgical resection and/or intravesical therapies, approximately 10%–15% of patients may develop...
invasive, locally advanced, and metastatic urothelial carcinoma [3]. Approximately 10% of patients have regionally advanced or metastatic disease at diagnosis [1]. The standard of care for patients with advanced disease is cisplatin-containing chemotherapy [2]. However, approximately 30%–50% of patients are considered ineligible for cisplatin-based chemotherapy because of comorbidities [4]. These patients are generally defined as those with a creatinine clearance less than 60 mL/minute, an Eastern Cooperative Oncology Group (ECOG) performance score of ≥2, or greater than or equal to grade 2 hearing impairment or peripheral neuropathy.

No approved therapies specifically for the cisplatin-ineligible population were available prior to April 2017. Non-cisplatin-based chemotherapy (e.g., carboplatin) was generally used but considered inferior to cisplatin [5]. Results from a few randomized trials of non-cisplatin-based regimens in this patient population along with several small single-arm trials of the most common regimens, including gemcitabine in combination with carboplatin or oxaliplatin, demonstrated objective response rates (ORRs) of approximately 30%–45% with median response durations of approximately 5–8 months. Overall survival (OS) in these patients is poor, ranging from 7 to 10 months [6–8]. Cytotoxic therapy is also poorly tolerated in these patients, with a high incidence of hematologic toxicities including neutropenic fever [6]. Thus, there is an unmet need for effective and tolerable treatments in cisplatin-ineligible patients with advanced urothelial carcinoma.

Monoclonal antibodies that disrupt the interaction of programmed death receptor-1 (PD-1) with programmed death-ligand 1 (PD-L1) release tumor-mediated inhibition of the immune response and have demonstrated antitumor activity in urothelial carcinoma. On May 18, 2016, the U.S. Food and Drug Administration (FDA) granted accelerated approval to atezolizumab (TECENTRIQ; Genentech, Inc., South San Francisco, CA), a monoclonal antibody that binds PD-L1, for use in patients with urothelial carcinoma that had progressed during or after platinum-based chemotherapy or within 12 months of neoadjuvant or adjuvant treatment with platinum-containing chemotherapy [9]. This approval was based on a similar ORR and improved duration of response (DoR) compared with historical data for available chemotherapy. Additional accelerated approvals of anti-PD-1 and anti-PD-L1 antibodies in the second-line urothelial carcinoma setting were subsequently granted to atezolizumab and pembrolizumab in this novel cisplatin-ineligible patient population, as well as FDA’s rationale for revising the indications based on emerging data from two ongoing trials.

**Clinical Trial Design**

Cohort 1 of IMvigor210 (ClinicalTrials.gov Identifier NCT02108652) and KEYNOTE-052 (ClinicalTrials.gov Identifier NCT02335424) were similar in design. Both studies were single arm, open-label trials that enrolled patients with advanced urothelial carcinoma who were ineligible for cisplatin-containing chemotherapy. Both trials enrolled patients regardless of PD-L1 expression levels. Cisplatin ineligibility was defined as one of the following: (a) impaired renal function (glomerular filtration rate >30 but <60 mL/minute), (b) greater than or equal to grade 2 hearing loss, (c) greater than or equal to grade 2 peripheral neuropathy, or (d) ECOG performance score of ≥2.

KEYNOTE-052 additionally defined patients with New York Heart Association Class III heart failure as ineligible to receive cisplatin and thus eligible for the trial. Both trials required patients to either be previously untreated or have disease progression more than 12 months from their last dose of platinum-containing neoadjuvant or adjuvant chemotherapy. Key exclusion criteria for both trials included a history of autoimmune disease requiring administration of systemic immunosuppressive medications.

IMvigor210 included two patient cohorts; Cohort 1 enrolled patients who were cisplatin ineligible. For the approval described herein, the FDA review focused primarily on patients enrolled in Cohort 1.

Patients received either an intravenous infusion of 1,200 mg of atezolizumab or 200 mg of pembrolizumab every 3 weeks. Patients on atezolizumab could continue treatment until unacceptable toxicity or radiographic or symptomatic progression. Patients on pembrolizumab could continue treatment until unacceptable toxicity, confirmed radiographic disease progression, investigator’s decision to withdraw the subject, completion of 24 months of treatment, or development of an intercurrent illness that prevented further treatment.

Patients receiving atezolizumab were scanned every 9 weeks for the first 54 weeks and every 12 weeks thereafter. Patients receiving pembrolizumab were scanned at 9 weeks, then every 6 weeks for 48 weeks and every 12 weeks thereafter. The primary endpoint of both trials was confirmed ORR using RECIST 1.1 as determined by blinded independent central review (BICR). Duration of response was also assessed.
PD-L1 expression status was prospectively assessed in IMvigor210 at a central laboratory using the Ventana PD-L1 (SP142) Assay. Although PD-L1 expression status was prospectively assessed in KEYNOTE-052 using the Agilent PD-L1 IHC22C3 pharmDx assay, a marketing application for the diagnostic test was not initially submitted with the pembrolizumab application.

Each drug has an ongoing randomized trial that could be used for confirmation of clinical benefit: IMvigor130 (NCT02807636) and KEYNOTE-361 (NCT02853305). The designs of both trials were similar. Both trials enrolled patients with locally advanced or metastatic urothelial cancer who were newly diagnosed or had received neoadjuvant/adjuvant therapy more than 12 months prior to study entry. Patients were enrolled regardless of PD-L1 status.

Patients enrolled in IMvigor130 were stratified by PD-L1 staining in tumor-infiltrating immune cells (IC), IC0, IC1, or IC2/3, where IC2/3 corresponds to ≥5% staining. Patients enrolling in KEYNOTE-361 were stratified by a Combined Positive Score (CPS) reflecting staining of both tumor and immune cells ≥10 or < 10. In IMvigor130, patients were randomized to atezolizumab plus chemotherapy, chemotherapy alone, or monotherapy with pembrolizumab. Chemotherapy consisted of gemcitabine plus cisplatin or carboplatin in both studies. Prior to randomization, investigators determined whether the patient would receive cisplatin or carboplatin. Strict criteria for cisplatin ineligibility were not prespecified; however, investigators were required to provide the reason for cisplatin ineligibility. The coprimary endpoints of IMvigor 130 are progression-free survival (PFS) and OS in the combination arm versus the chemotherapy-alone arm and OS in the atezolizumab monotherapy versus chemotherapy arms, all in the intent-to-treat (ITT) populations, while OS in PD-L1-high patients will be evaluated if OS is positive in both the combination versus chemotherapy ITT populations and atezolizumab monotherapy versus chemotherapy ITT populations. KEYNOTE-361 incorporates multiple hierarchical hypotheses with initial primary endpoints of PFS and OS in the combination versus chemotherapy arms and subsequent noninferiority and superiorly endpoints of OS in the pembrolizumab versus chemotherapy arm.

### Results

#### Atezolizumab

IMvigor210 enrolled 119 patients in Cohort 1. Key demographic and disease characteristics of these patients are summarized in Table 1. The median age of Cohort 1 was 73 years. Approximately 26% of patients had lymph node-only disease, whereas 21% had liver involvement. The most common reason for cisplatin ineligibility was impaired renal function.

As of the data cutoff, 18% of patients were on study treatment and 82% of patients discontinued study treatment. Of the discontinued patients, 77% were due to disease progression and 10% due to adverse events. The median treatment duration was 3.6 months (range: 0.02–20 months).

#### Efficacy

Efficacy results are shown in Table 2. With a median follow-up time of 14.4 months, BICR-confirmed ORR was 23.5% (95% confidence interval [CI]: 16.2%–32.2%) in all treated patients. Both complete responses (CRs) and partial

| **Table 1. Key baseline characteristics of patients in Cohort 1 of IMvigor210 and KEYNOTE-052** |
|---------------------------------|-----------------|
| **Characteristics**             | IMvigor210      | KEYNOTE-052      |
| **No. of patients**             | 119             | 370             |
| **Age in years, median (range)**| 73 (51–92)      | 74 (34–94)      |
| **Sex**                         |                 |                 |
| **Male**                        | 96 (81)         | 286 (77)        |
| **Female**                      | 23 (19)         | 84 (23)         |
| **Race**                        |                 |                 |
| **White**                       | 108 (91)        | 328 (89)        |
| **Black**                       | 3 (3)           | 8 (2)           |
| **Asian**                       | 2 (2)           | 26 (7)          |
| **Other**                       | 6 (5)           | 8 (2)           |
| **ECOG score**                  |                 |                 |
| **0**                           | 45 (38)         | 80 (22)         |
| **1**                           | 50 (42)         | 133 (36)        |
| **2**                           | 24 (20)         | 156 (42)        |
| **3**                           | 0               | 1 (<1)          |
| **Reasons for cisplatin ineligibility** |             |                 |
| **ECOG of 2**                   | 24 (20)         | 120 (32)        |
| **Renal dysfunction**           | 83 (70)         | 182 (50)        |
| **ECOG 2 and renal dysfunction**| 8 (7)           | 34 (9)          |
| **Other reasons**               | 20 (17)         | 33 (9)          |
| **Primary urothelial cancer site** |             |                 |
| **Upper tract**                 | 33 (33)         | 69 (18)         |
| **Lower tract**                 | 85 (71)         | 300 (81)        |
| **Other/unknown**               | 1 (<1)          | 1 (<1)          |
| **Disease sites**               |                 |                 |
| **Lymph node only**             | 31 (26)         | 50 (14)         |
| **Visceral metastasis**         | 78 (66)         | 316 (85)        |
| **Liver**                       | 25 (21)         | 78 (21)         |
| **Number of Bajorin/MSKCC risk factors** |             |                 |
| **0**                           | 35 (29)         | 29 (8)          |
| **1**                           | 66 (56)         | 210 (57)        |
| **2**                           | 18 (15)         | 131 (35)        |
| **PD-L1 status in ICs**         |                 |                 |
| **PD-L1 expression of <5%**     | 87 (73)         | Not applicable  |
| **PD-L1 expression of ≥5%**     | 32 (27)         | Not applicable  |

Data are presented as n (%). aIncludes Karnofsky Performance Status <80% and visceral (lung, liver, or bone) metastases. bProportion of PD-L1-stained tumor-infiltrating immune cells within the tumor area. Abbreviations: ECOG, Eastern Cooperative Oncology Group; ICs, immune cells; MSKCC, Memorial Sloan Kettering Cancer Center; PD-L1, programmed death-ligand 1.
Table 2. Efficacy results of Cohort 1 of IMvigor210 and KEYNOTE-052

| Variable                        | Cohort 1 IMvigor210 (atezolizumab) | KEYNOTE-052 (pembrolizumab) |
|--------------------------------|----------------------------------|--------------------------|
| No. of patients enrolled       | 119                              | 370                      |
| No. of BICR-assessed confirmed responders | 28                              | 106                      |
| ORR, % (95% CI)                | 23.5 (16.2–32.2)                 | 28.6 (24.1–33.5)         |
| Complete response, %           | 6.7                              | 6.8                      |
| Partial response, %            | 16.8                             | 21.9                     |
| Median DoR, months (range)     | NR (3.7 to 16.6+)                | NR (1.4+ to 17.8+)       |

Abbreviations: +, denotes a censored value; BICR, blinded independent central review; CI, confidence interval; DoR, duration of response; NR, not reached; ORR, objective response rate.

Responses (PRs) were observed. Median DoR in responders was not reached as of the data cutoff date of March 14, 2016, and the observed response durations ranged from 3.7 to 16.6+ months. At the time of data cutoff, responses were ongoing for at least 6 months in 64.2% of responding patients and for at least 12 months in 21.4% of responding patients.

Responses were observed in both PD-L1 expression subgroups. The confirmed ORR was 21.8% (95% CI: 13.7%–32.0%) in patients with PD-L1 expression of <5% and 28.1% (95% CI: 13.8%–46.8%) in those with PD-L1 expression of ≥5% in ICs. There were six CRs (6.9%) in patients with PD-L1 expression of <5% and two CRs (6.3%) in those with PD-L1 expression of ≥5%. Median duration of response was not reached in either subgroup. Responses were ongoing for at least 6 months in 82% of responding patients and for at least 12 months in 29% of responding patients.

In exploratory subgroup analyses, responses were observed in patients with nonbladder urothelial carcinoma, visceral metastases, and prior Bacillus Calmette-Guerin (BCG) treatment. Patients with a Bajorin risk score of 2 [11] had a lower response rate (11.1% [95% CI: 1.4%–34.7%]) compared with those with scores of 0–1 (25.7% [95% CI: 17.6%–35.4%]). Durable responses were observed in all the subgroups.

Toxicity

All 119 patients enrolled in Cohort 1 received at least one dose of atezolizumab and were included in the safety analysis. Immune-mediated adverse events (imAEs) were defined as events requiring the use of systemic steroids with no alternate etiology, endocrine events, and other events thought to be immune-related. Immune-mediated adverse events were generally managed with administration of high-dose (1–2 mg/kg daily of prednisone or equivalent) corticosteroids followed by a taper and interruption of atezolizumab therapy. In total, 19.3% of patients experienced an imAE, including 12.6% of patients who required systemic corticosteroids and 6.7% who required only hormone replacement therapy. Five percent of patients received an oral prednisone dose equivalent to ≥40 mg daily for an immune-mediated adverse event. The reported imAEs in this cohort included hypothyroidism, rash, hepatic injury, colitis, arthritis, adrenal insufficiency, and rhabdomyolysis. The pattern of imAEs was generally consistent with that observed with other approved PD-1/PD-L1-targeted products. Thyroid function tests were routinely collected only at baseline and end of study, such that the reported incidence of hypothyroidism may underestimate the true incidence.

Five patients (4.2%) who were treated with atezolizumab experienced one of the following events, which led to death: sepsis, cardiac arrest, myocardial infarction, respiratory failure, or respiratory distress, within 30 days of last drug administration. One additional patient with herpetic meningoencephalitis was reported to have died as a result of disease progression. Four percent of patients treated with atezolizumab discontinued treatment because of an adverse event. The causes of discontinuation were diarrhea/colitis, fatigue, hypersensitivity, and dyspnea.

The most common grade 1–4 adverse events occurring in at least 20% of patients treated with atezolizumab were fatigue (52%), decreased appetite (26%), nausea (25%), urinary tract infections (22%), pyrexia (21%), and constipation (21%). The most common grade 3–4 adverse events occurring in at least 2% of patients treated with atezolizumab were fatigue (8%), urinary tract infection (5%), diarrhea (4%), intestinal obstruction (3%), decreased appetite (3%), sepsis (3%), back/neck pain (3%), renal failure (3%), and hypotension (3%). The laboratory abnormalities worsening from baseline to grade 3–4 in at least 2% of patients treated with atezolizumab included hyponatremia (15%), hyperglycemia (10%), lymphopenia (9%), anemia (7%), increased alkaline phosphatase (7%), increased creatinine (5%), hypophosphatemia (4%), increased alanine aminotransferase (4%), increased aspartate aminotransferase (4%), hyperkalemia (3%), hypermagnesemia (3%), and hyperbilirubinemia (3%).

Pembrolizumab

KEYNOTE-052 enrolled 370 patients. Key demographic and disease characteristics of these patients are summarized in Table 1. The median age was 74 years. Approximately 14% of patients had lymph node-only metastatic disease, whereas 21% had liver involvement. The most common reason for cisplatin ineligibility was impaired renal function. As of the data cutoff, 50% of patients were on study treatment and 50% of patients discontinued study treatment. Of the discontinued patients, 80% were due to disease progression and 10% due to adverse events. The median treatment duration was 3.4 months (range: 0.03–19.94 months).

Efficacy

Efficacy results are shown in Table 2. With a median follow-up time of 7.8 months, ICR-confirmed ORR was 28.6% (95% CI: 24.1%–33.5%) in all treated patients. Both CRs and PRs were observed. Median DoR in responders was not reached as of the data cutoff date of December 19, 2016, and the observed response durations ranged from 1.4+ to 17.8+ months.
Responses were observed in both PD-L1 expression subgroups. The confirmed ORR was 21% (95% CI: 16%–26%) in patients with CPS <10 or unknown and 47% (95% CI: 38, 57%) in those with CPS ≥10. There were 8 CRs (3%) in patients with CPS <10 or unknown and 17 CRs (15%) in those with CPS ≥10. Median duration of response was not reached in either subgroup. At the time of data cutoff, responses were ongoing for at least 6 months in 52% of responding patients and for at least 12 months in 7% of responding patients.

Responses were observed in patients with lymph node-only metastases, visceral metastases, and prior BCG treatment, and in patients who received prior adjuvant/neoadjuvant therapy.

Toxicity
All the 370 patients in this study received at least one dose of pembrolizumab and were included in the safety analysis. In total, 17% of patients experienced an immune-mediated adverse event, including 8% of patients who required systemic corticosteroids and 8% who required hormone replacement therapy. Five percent of patients received an oral prednisone dose equivalent to ≥40 mg daily for an immune-mediated adverse event. One imAE was myositis that led to multiple organ failure and death despite use of corticosteroids. The pattern of immune-related adverse events was generally consistent with other approved agents targeting the PD-1/PD-1 pathway.

Eighteen patients (5%) treated with pembrolizumab died as a result of an adverse event. Thirteen of these patients (3.5%) died within 30 days of the last dose of sepsis (5), renal failure (2), stroke, pneumonia, aspiration, respiratory failure, colonic perforation, and unknown cause. Eleven percent of patients discontinued pembrolizumab because of an adverse event. The most common cause of discontinuation were infections, including pneumonia, urinary tract infections, and sepsis.

The most common grade 1–4 adverse events occurring in at least 20% of patients treated with pembrolizumab were fatigue (38%), musculoskeletal pain (24%), decreased appetite (22%), constipation (21%), rash (21%), and diarrhea (20%). The most common grade 3–4 adverse events occurring in at least 2% of patients treated with pembrolizumab were urinary tract infection (9%), anemia (7%), fatigue (6%), musculoskeletal pain (4.9%), hyponatremia (4.1%), elevated liver function tests (3.5%), hematuria (3%), abdominal pain (2.7%), and diarrhea (2.4%). The grade 3–4 laboratory abnormalities occurring in at least 2% of patients treated with pembrolizumab included anemia (7%), elevated blood creatinine (4.3%), hyponatremia (4.1%), elevated alkaline phosphatase (2.2%), and hyperkalemia (2.2%).

IMvigor130 and KEYNOTE-361
In the two ongoing clinical trials of atezolizumab (IMvigor130) and pembrolizumab (KEYNOTE-361), the Data Monitoring Committee (DMC) for each study performed an early unplanned review and found that patients in the monotherapy arms of both trials with PD-L1-low status had decreased survival compared with patients who received cisplatin- or carboplatin-based chemotherapy. There was no change in the adverse event profile of either drug. Both Merck, manufacturer of pembrolizumab, and Genentech, manufacturer of atezolizumab, stopped enrolling patients whose tumors have PD-L1-low status to the atezolizumab or pembrolizumab monotherapy arms per the DMCs’ recommendations. For patients already recruited in the mono-therapy arms, the DMC recommended continuation in the trial without treatment modification but that they be reconsented. The monotherapy arms remain open for new enrollment only to patients whose tumors have PD-L1-high status; the combination arms and the chemotherapy arms of both studies also remain open to all patients regardless of PD-L1 status.

Discussion
FDA review of the initial supplemental applications found that treatment with atezolizumab or pembrolizumab had a favorable benefit-risk profile (Table 3) in patients with locally advanced or metastatic urothelial carcinoma who are not eligible for cisplatin-based chemotherapy. Treatment with either agent elicited confirmed objective antitumor responses in approximately 24%–29% of patients in the single-arm trials. Objective response rates in these trials were similar to or greater than those seen in the treatment of patients who had received prior platinum-based therapy. This may be due to the extent of prior therapy and to the higher proportion of patients with lymph node-only disease in the trials of cisplatin-ineligible patients. Although the median DoR were not reached at the data cutoff for the response analyses in either study, the duration of response was substantially longer than that seen historically with cytotoxic chemotherapy. The majority of responding patients maintained their response for ≥6 months and some for ≥12 months, indicating durable responses with either product. There was a slight decrease in ORR in patients treated with atezolizumab or pembrolizumab who had tumors considered PD-L1 negative by the Ventana PD-L1 (SP142) Assay or the Agilent PD-L1 IHC 22C3 pharmDx assay, respectively, compared with those with tumors considered PD-L1 positive. However, in both trials, responses were observed with long durations of response in both the PD-L1-high and PD-L1-low subgroups. Responses were also observed in all demographic and disease subgroups in both trials, including patients with various primary tumor sites, prior BCG treatment, and visceral metastases. Duration of response did not appear to differ by subgroup.

Given the toxicities associated with cisplatin, carboplatin is often substituted for cisplatin in combination with other agents, including gemcitabine, as first-line therapy. In a randomized phase III trial comparing two carboplatin-based regimens in this population, the median survival was only 8–9 months [7]. For patients who received carboplatin and gemcitabine, the confirmed objective response rate was 36%, whereas the median response duration was only 5.4 months. The most common severe toxicities associated with chemotherapy regimens in this patient population were neutropenia (46%), thrombocytopenia (19%), and anemia (18%) [6].
Table 3. Benefit-risk assessments of atezolizumab and pembrolizumab for first-line use in cisplatin-ineligible patients with advanced urothelial carcinoma at the time of initial accelerated approval

| Dimension              | Evidence and uncertainties                                                                 | Conclusions and reasons                                                                 |
|------------------------|----------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------|
| Analysis of condition  | • Advanced urothelial carcinoma in patients ineligible for cisplatin has a poor prognosis, with a median survival of 7–10 months. Approximately 15,000 deaths from advanced urothelial carcinoma each year | This disease is serious and life-threatening. This represents a significant unmet medical need. |
| Current treatment options | • No approved products in the U.S. for first-line therapy for this patient population<br>• Carboplatin-based combination chemotherapy regimens are generally administered, but are associated with high toxicity, short duration of response, and poor outcome. | All the products are palliative and have significant adverse reactions and/or intolerance. Responding patients generally have short response durations. |
| Benefit                | Atezolizumab:<br>• Of the unenriched population of 119 patients, 23.5% had confirmed responses, including 8 (6.7%) complete responders. The ORR was 28.1% in the PD-L1 IC 2/3 group and 21.8% in the PD-L1 IC 0/1 group.<br>• Median response duration was not reached (range 3.7 to 16.6+ months). Of the responders, 75% (21/28) had ongoing responses with a median follow-up time of 14.4 months.<br>Pembrolizumab:<br>• Of the 370 patients, 28.6% had confirmed responses, including 25 (6.8%) complete responders.<br>• Median response duration was not reached (range 1.4 to 17.8+ months). Of the responders, 75% (80/106) had ongoing responses with a median follow-up time of 7.8 months. | Substantial evidence of effectiveness for first-line use of atezolizumab or pembrolizumab in this patient population was durable responses. The durable responses are reasonably likely to provide an advantage over conventional chemotherapy, where duration of response has generally been short. |
| Risk                   | • Tolerated in most study patients<br>• Important risks include hepatitis, pneumonitis, endocrine disorders, colitis, infection, and neurological disorders. | The safety profiles of atezolizumab and pembrolizumab are similar to those observed in PD-1/PD-L1-targeted products. The adverse event profile of either product appears less toxic and better tolerated than that of conventional chemotherapy. |
| Risk management        | • Nonendocrine immune-mediated adverse events were largely reversible with the use of corticosteroids. A medication guide is distributed to describe these risks. | The safe use of atezolizumab and pembrolizumab can be managed through accurate labeling and routine pharmacovigilance. |

Abbreviations: ORR, objective response rate; PD-1, programmed death receptor-1; PD-L1, programmed death-ligand 1.

Although the response rates for atezolizumab and pembrolizumab were lower relative to those for chemotherapy in the same disease setting, the responses were more durable, with durable responses of more than 12 months observed in a small number of patients. The safety profiles were largely consistent with those seen in the second-line setting. Approximately 10%–20% of patients experienced immune-mediated adverse events that required either use of systemic corticosteroids or hormone replacement. Apart from endocrine events, immune-mediated toxicity was typically reversible. Although the nature of toxicity differed from that of cytotoxic chemotherapy, the incidence of high-grade toxicity related to atezolizumab and pembrolizumab compared favorably, and only 5% of patients required high-dose corticosteroids in either trial. These durable responses combined with different and potentially improved safety profile represent an improvement over available chemotherapies and are reasonably likely to predict clinical benefit in a patient population with a high unmet medical need.

Taken together, the evidence, as summarized in our Benefit-Risk Assessment in Table 3, was considered sufficient for the respective accelerated approvals of atezolizumab and pembrolizumab for the intended clinical use. This provides the first nonchemotherapeutic treatments in this disease setting and addresses an unmet medical need for this unique patient population. The results from these trials should not be extrapolated to patients who may be eligible for cisplatin-based chemotherapy.

Based on the regulatory requirements for accelerated approval, the continued approval of atezolizumab and pembrolizumab for this indication may be contingent upon verification and description of clinical benefit in a confirmatory trial(s). Preliminary, early analysis reports generated by the respective DMCs from IMvigor130 (17% of OS events) and KEYNOTE-361 (19% of OS events) demonstrated decreased survival in patients with PD-L1-low status on the monotherapy arms. Although the study stopped accrual of patients with low PD-L1 status to the monotherapy arms, given the
previous accelerated approvals for pembrolizumab and atezolozubab monotherapy for the cisplatin-ineligible populations, FDA no longer considered the benefit-risk profile favorable for all cisplatin-ineligible patients. Therefore, on June 18, 2018, the indication for both agents was modified to include only patients who are not eligible for cisplatin-containing chemotherapy and who have high expression of PD-L1 or are not eligible for any platinum-containing chemotherapy regardless of the level of PD-L1 expression. As there are some patients for whom any platinum-containing chemotherapy (cisplatin or carboplatin) is not indicated, FDA worded part of the indication for use of these drugs in patients not eligible for any platinum-containing regimen regardless of tumor PD-L1 status. There were substantial differences between IMVigor210 and KEYNOTE-052 and the ongoing randomized trials. The key difference is that both IMVigor210 and KEYNOTE-052 required patients to meet strict criteria concerning platinum eligibility, whereas the choice of cisplatin- or carboplatin-based therapy was made by the Investigator on the randomized phase III trials.

This change in indication based on PD-L1 expression was made prior to companion diagnostic approval, given the urgency in refining the product labeling for concerns over decreased survival. The companion diagnostic approvals were agreed upon as postmarketing commitments and were subsequently approved on July 2, 2018 (atezolizumab), and August 16, 2018 (pembrolizumab).

**CONCLUSION**

The efficacy and safety results described above demonstrated an acceptable benefit-risk profile to support the initial accelerated approval of atezolizumab and pembrolizumab for the treatment of patients with locally advanced or metastatic urothelial carcinoma who are not eligible for cisplatin-based chemotherapy. These approvals were based on confirmed ORRs comparable to available therapy in combination with improved DoRs and favorable safety profiles as well as the unmet need in this patient population. Per accelerated approval regulations, continued approval in patients with urothelial carcinoma who are not eligible for cisplatin-containing chemotherapy may be contingent upon verification and description of clinical benefit in confirmatory trials. Based on emerging results from these ongoing, large, randomized trials, with coprimary endpoints of PFS and OS, the indication statement for both agents was revised to exclude patients eligible for platinum-based chemotherapy with low PD-L1 expression. This revision reflects the balance between the benefit of early availability of promising drugs to patients and the risk of accelerated approval without verification of clinical benefit [12]. The early reports of decreased survival are preliminary and the confirmatory studies are still ongoing; thus, further revisions to the indication may be warranted.

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