Nivolumab monotherapy in recurrent metastatic urothelial carcinoma (CheckMate 032): a multicentre, open-label, two-stage, multi-arm, phase 1/2 trial

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Summary

Background Few effective treatments exist for patients with advanced urothelial carcinoma that has progressed after platinum-based chemotherapy. We assessed the activity and safety of nivolumab in patients with locally advanced or metastatic urothelial carcinoma whose disease progressed after previous platinum-based chemotherapy.

Methods In this phase 1/2, multicentre, open-label study, we enrolled patients (age ≥18 years) with urothelial carcinoma of the renal pelvis, ureter, bladder, or urethra at 16 sites in Finland, Germany, Spain, the UK, and the USA. Patients were not selected by PD-L1 expression, but tumour PD-L1 membrane expression was assessed retrospectively. Patients received nivolumab 3 mg/kg intravenously every 2 weeks until disease progression or treatment discontinuation because of unacceptable toxicity or other protocol-defined reasons, whichever occurred later. The primary endpoint was objective response by investigator assessment. All patients who received at least one dose of the study drug were included in the analyses. We report an interim analysis of this ongoing trial. CheckMate 032 is registered with ClinicalTrials.gov, NCT01928394.

Findings Between June 5, 2014, and April 24, 2015, 86 patients with metastatic urothelial carcinoma were enrolled in the nivolumab monotherapy group and 78 received at least one dose of treatment. At data cutoff (March 24, 2016), the minimum follow-up was 9 months (median 15·2 months, IQR 12·9–16·8). A confirmed investigator-assessed objective response was achieved in 19 (24·4%, 95% CI 15·3–35·4) of 78 patients. Grade 3–4 treatment-related adverse events occurred in 17 (22%) of 78 patients; the most common were elevated lipase (four [5%]), elevated amylase (three [4%]), and fatigue, maculopapular rash, dyspnoea, decreased lymphocyte count, and decreased neutrophil count (two [3%] each). Serious adverse events were reported in 36 (46%) of 78 patients and eight (10%) had a serious adverse event judged to be treatment related. Two (3%) of 78 patients discontinued because of treatment-related adverse events (grade 4 pneumonitis and grade 4 thrombocytopenia) and subsequently died.

Interpretation Nivolumab monotherapy was associated with a substantial and durable clinical response and a manageable safety profile in previously treated patients with locally advanced or metastatic urothelial carcinoma. These data support further investigation of nivolumab monotherapy in advanced urothelial carcinoma.

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Introduction

Nearly three decades have passed since the first ground-breaking treatments were developed for patients with metastatic urothelial carcinoma. The combination chemotherapy regimen methotrexate, vinblastine, doxorubicin, and cisplatin has not been surpassed in terms of response and survival.1 In 2000, gemcitabine plus cisplatin was tested as a less toxic alternative, but 37% of patients could not tolerate the treatment regimen.2 Decades of research exploring cytotoxic frontline chemotherapies followed,3 but no treatments were able to exceed the therapeutic outcomes achieved with the combination of methotrexate, vinblastine, doxorubicin, and cisplatin. About 25–50% of patients with metastatic urothelial carcinoma are unable to receive cisplatin-based chemotherapy because of renal impairment.4 Nonetheless, platinum-based combination chemotherapy remains the standard first-line treatment for patients with metastatic urothelial carcinoma.5 In the second-line setting, many drugs have been tested, but none have become established as a standard of care because of a low frequency of response (10% of patients or less). The most intensively studied regimen in the second-line setting—vinflunine plus best supportive care—did not significantly improve overall survival compared with best supportive care in a phase 3 trial (hazard ratio 0·9, 95% CI 0·7–1·1; intention-to-treat population).6 Although an increase in median overall survival of 2–6 months was noted with vinflunine in a subsequent analysis of the eligible population that excluded patients with protocol violations at baseline (hazard ratio 0·8, 95% CI 0·6–1·0; p=0·023). Immune checkpoint treatment, consisting of blockade of immune inhibitory pathways, has led to substantial advances in the treatment of cancer. The potential for...
nivolumab improved overall survival in melanoma,treating several solid tumours. Compared with comparator chemotherapy.

disease progression during or after platinum-containing chemotherapy responded to treatment,urothelial carcinoma who have received combination treatment with ipilimumab. The outcomes of patients who received combination treatment with ipilimumab will be reported separately.

Patients aged at least 18 years were eligible if they had progressive disease after at least one previous study who had locally advanced or metastatic urothelial carcinoma (unselected for PD-L1 expression) and were previously treated with platinum-based chemotherapy.

Methods

Study design and participants

CheckMate 032 is a multicentre, open-label, two-stage, multi-arm, phase 1/2 study. Patients with histologically or cytologically confirmed carcinoma of the renal pelvis, ureter, bladder, or urethra were enrolled at 16 sites in five countries (Finland, Germany, Spain, the UK, and the USA; appendix p 3). The protocol is available in the appendix.

The urothelial carcinoma part of this trial consisted of two treatment regimens: the first with nivolumab monotherapy and the second with nivolumab in combination with ipilimumab. The outcomes of patients who received combination treatment with ipilimumab will be reported separately.

Evidence before this study

We searched PubMed from Dec 1, 1994, to May 31, 2014, using the search terms “metastatic urothelial carcinoma”, “relapsed urothelial carcinoma”, “clinical trials”, “immune response”, “immune checkpoint blockade”, and “immunotherapy”. Studies identified from the search revealed poor outcomes for patients with recurrent or relapsed urothelial carcinoma, with few treatment options to improve survival. A few papers reported findings from studies investigating immunotherapy in advanced urothelial carcinoma, including BCG treatment. The immune system is a target for treatment in urothelial carcinoma; immunotherapy with BCG is standard treatment for superficial urothelial carcinoma, reducing the risk of local recurrence by about 60% and leading to 5-year survival of about 90% in patients with unifocal disease. Additionally, CD8+ tumour-infiltrating lymphocytes are predictive of survival in muscle-invasive urothelial carcinoma; patients with advanced urothelial cancer and higher numbers of CD8+ tumour-infiltrating lymphocytes within the tumour (≥8 cells) seem to have better disease-free and overall survival than those with similar-staged urothelial carcinoma and fewer intra-tumoural CD8+ tumour-infiltrating lymphocytes. Together with the promising anti-PD-L1 data reported in advanced urothelial carcinoma, these findings provided the rationale for further investigation of immune checkpoint blockade with the PD-1 inhibitor nivolumab for recurrent metastatic urothelial carcinoma.

Added value of all the available evidence

Patients with recurrent metastatic urothelial carcinoma have poor clinical prospects because of the scarcity of effective treatments. A growing body of evidence suggests that immune checkpoint inhibition can offer effective treatment for patients with urothelial carcinoma, as in other tumour types. The results presented here support further development of PD-1 checkpoint inhibition in larger trials of this disease.

A promising target for immunotherapy is the PD-1/PD-L1 immune checkpoint. PD-1 is expressed on T cells and can inhibit T-cell responses on interaction with its ligands, PD-L1 and PD-L2; high PD-L1 expression has been found in bladder tumour cells. In a clinical trial with atezolizumab, an antibody that blocks PD-L1, 15% of patients with metastatic or surgically unresectable urothelial carcinoma who were previously treated with platinum-based chemotherapy responded to treatment, leading to US Food and Drug Administration approval of this drug for the treatment of patients with locally advanced or metastatic urothelial carcinoma who have disease progression during or after platinum-containing chemotherapy or disease progression within 12 months of neoadjuvant or adjuvant treatment with platinum-containing chemotherapy.

Treatment with nivolumab, a fully human monoclonal IgG4 antibody that blocks PD-1, has proven effective in several solid tumours. Compared with comparator treatments (eg, dacarbazine, docetaxel, or everolimus), nivolumab improved overall survival in melanoma, non-small-cell lung cancer, renal-cell carcinoma, and head and neck cancer, and studies have shown promising clinical activity in several additional malignancies including Hodgkin’s lymphoma and microsatellite-unstable colorectal cancer.

Nivolumab is being investigated in an ongoing multicentre, open-label, phase 1/2 clinical study of several advanced or metastatic solid tumour types (CheckMate 032). We report the activity and safety of nivolumab monotherapy in a cohort of patients from this study who had locally advanced or metastatic urothelial carcinoma (unselected for PD-L1 expression) and were previously treated with platinum-based chemotherapy.

this approach in the treatment of urothelial carcinoma is suggested by the effectiveness of immunotherapy with BCG: given intravesically, BCG induces an immune response against tumour cells and is indicated as adjuvant treatment after surgical resection in patients with high-grade non-muscle-invasive urothelial carcinoma. The immune checkpoint inhibitor ipilimumab, which blocks CTLA-4, enhanced immune responses and tumour regression in studies of patients with localised urothelial carcinoma. A promising target for immunotherapy is the PD-1/PD-L1 immune checkpoint. PD-1 is expressed on T cells and can inhibit T-cell responses on interaction with its ligands, PD-L1 and PD-L2; high PD-L1 expression has been found in bladder tumour cells. In a clinical trial with atezolizumab, an antibody that blocks PD-L1, 15% of patients with metastatic or surgically unresectable urothelial carcinoma who were previously treated with platinum-based chemotherapy responded to treatment, leading to US Food and Drug Administration approval of this drug for the treatment of patients with locally advanced or metastatic urothelial carcinoma who have disease progression during or after platinum-containing chemotherapy or disease progression within 12 months of neoadjuvant or adjuvant treatment with platinum-containing chemotherapy.

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The urothelial carcinoma part of this trial consisted of two treatment regimens: the first with nivolumab monotherapy and the second with nivolumab in combination with ipilimumab. The outcomes of patients who received combination treatment with ipilimumab will be reported separately.

Patients aged at least 18 years were eligible if they had progressive disease after at least one previous
platinum-based chemotherapy treatment for metastatic disease or locally advanced unresectable disease, recurrence within 1 year of completing previous platinum-based neoadjuvant or adjuvant treatment, or had previously refused standard treatment with chemotherapy for the treatment of metastatic (ie, stage IV) or locally advanced unresectable disease. Inclusion criteria were locally advanced or metastatic urothelial carcinoma, an Eastern Cooperative Oncology Group (ECOG) performance status of 0 or 1, and measurable disease by CT or MRI (per Response Evaluation Criteria in Solid Tumors [RECIST] version 1.1).23 Patients were not selected based on tumour PD-L1 expression. Baseline laboratory tests used to assess eligibility were white blood cell counts and measurement of neutrophil, platelet, haemoglobin, serum creatinine, alanine aminotransferase, aspartate aminotransferase, total bilirubin, albumin, lipase, and amylase concentrations. Key exclusion criteria included active brain or leptomeningeal metastases; any serious or uncontrolled medical disorder; history of or active, known, or suspected autoimmune disease (vitaligo, type 1 diabetes mellitus, residual hypothyroidism caused by autoimmune thyroiditis, and disorders not expected to recur in the absence of an external trigger were permitted); need for immunosuppressive doses of systemic corticosteroids (>10 mg per day prednisolone equivalents) for at least 2 weeks before study drug treatment; and previous treatment with experimental antitumour vaccines or any modulator of T-cell function or checkpoint pathway. Median survival for patients with relapsed advanced transitional cell carcinoma of the urothelium has been reported as about 4-6-6-9 months.3

The study was approved by the institutional review board or independent ethics committee for each centre and was done in accordance with Good Clinical Practice guidelines defined by the International Conference on Harmonisation. All patients provided written informed consent to participate before study participation based on the principles of the Declaration of Helsinki.

Procedures

Patients with metastatic urothelial carcinoma were enrolled by an interactive voice response system to receive nivolumab 3 mg/kg intravenously every 2 weeks until progression or treatment discontinuation because of unacceptable toxicity or other protocol-defined reasons, whichever occurred later. Patients in the nivolumab monotherapy group could switch to nivolumab plus ipilimumab (nivolumab 1 mg/kg and ipilimumab 3 mg/kg or nivolumab 3 mg/kg and ipilimumab 1 mg/kg intravenously, every 3 weeks for four cycles) after progression if they met prespecified criteria, as part of this clinical trial (appendix p 4). Treatment beyond RECIST-defined progression was permitted if nivolumab was tolerated and clinical benefit was noted, on the basis of investigator assessment (appendix p 4). No dose reductions or modifications were permitted. Criteria for dose delay until resolution of a treatment-related adverse event to grade 1 or lower have been described previously.23 Adverse events that could lead to a dose delay were grade 2 or worse non-skin events (except for grade 2 fatigue), grade 3 skin events, and grade 3 laboratory abnormalities (except for asymptomatic amylase and lipase increases).

Tumour assessments (CT or MRI, or both) were done by the investigator at baseline (within 28 days before the first dose of study drug), every 6 weeks (within 1 week earlier or later) until week 24, and every 12 weeks (within 1 week earlier or later) thereafter until disease progression per RECIST. If study treatment was discontinued for reasons other than disease progression, tumour assessments were continued. Laboratory assessments were done within 72 h before dose until week 24, and within 72 h after every alternate dose thereafter. Safety assessments were done continuously in all treated patients, and adverse events and laboratory values were graded according to the National Cancer Institute’s Common Terminology Criteria for Adverse Events, version 4.0. As described previously,23 assessment of tumour PD-L1 protein expression was done retrospectively with a Dako PD-L1 immunohistochemical 28-8 pharmDx kit (Dako North America, Carpinteria, CA, USA) in pretreatment tumour biopsy specimens that were fresh or archived within 3 months before treatment start. Several pretreatment specimens could be tested for PD-L1 expression and used to define PD-L1 status. The most recently collected specimen before the start of study treatment was used to define PD-L1 status, although results were not needed before the start of study treatment. Tumour PD-L1 expression was categorised as positive when staining of tumour-cell membrane, at any intensity, was noted in at least 1% or at least 5% of tumour cells in a section that included at least 100 assessable tumour cells in any sample (ie, not necessarily the sample collected closest to the time of study drug treatment). PD-L1 expression thresholds were chosen based on findings from previous studies in other tumour types.12-15

Outcomes

The primary endpoint was the proportion of patients with a confirmed investigator-assessed objective response, defined as the number of patients with a best overall response of complete or partial response per RECIST divided by the number of treated patients. For a complete or partial response to be judged to be a best overall response, the assessment needed to be confirmed by a second scan no less than 4 weeks after the criteria for response was first met. Patients who did not meet response-evaluable criteria (ie, at least one target lesion at baseline and at least one on-study assessment) were judged to be not assessable.

Secondary endpoints were safety, defined as the incidence of treatment-related adverse events leading to drug discontinuation within the first 12 weeks of treatment in patients who had at least one dose of study drug;
duration of response, defined as the time from partial or complete response until progressive disease or death from any cause; progression-free survival, defined as the time from treatment assignment to the date of the first documented tumour progression or death from any cause, whichever occurred first; and overall survival, defined as the time between the date of treatment assignment and the date of death from any cause.

Objective response, overall survival, and progression-free survival by PD-L1 expression were exploratory endpoints. In a post-hoc analysis, we also identified patients who had a non-conventional benefit; that is, patients who did not have a best overall response of partial response or complete response before initial RECIST-defined progression, and met at least one of the following criteria: appearance of a new lesion followed by a decrease from baseline of at least 10% in the sum of the target lesions; initial increase from nadir at least 20% in the sum of the target lesions followed by reduction from baseline of at least 30%; and initial increase from nadir at least 20% in the sum of the target lesions followed by at least two tumour assessments showing no further progression, defined as a 10% additional increase in the sum of target lesions and new lesions.

Statistical analysis
We enrolled patients according to a one-stage design, with a total sample size of 60–100 patients needed to assess whether nivolumab resulted in an objective

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**Figure 1: Study profile**
*One pneumonitis and one thrombocytopenia. 1One each of increased blood creatinine concentration, hepatitis C infection, anaemia, and urosepsis. 1Not including adverse events in patients who received combination treatments after switching.

| Nivolumab (n=78) |
|------------------|
| Age (years) 65.5 (31-85) |
| <65 years 37 (47%) |
| ≥65 and <75 years 31 (40%) |
| ≥75 years 10 (13%) |
| Sex Male 54 (69%) |
| Female 24 (31%) |
| Race White 72 (92%) |
| Black or African-American 4 (5%) |
| Asian 1 (1%) |
| Other 1 (1%) |
| Smoking status Present or former 48 (62%) |
| Never 29 (37%) |
| Unknown 1 (1%) |
| Number of previous regimens 1 26 (33%) |
| 2-3 42 (54%) |
| >3 10 (13%) |
| ECOG performance status 0 42 (54%) |
| 1 36 (46%) |
| Baseline metastatic disease Visceral 61 (78%) |
| Liver 20 (26%) |
| Lymph node only 13 (17%) |
| Baseline haemoglobin concentration (g/dL) <10 11 (14%) |
| ≥10 67 (86%) |
| Number of Bellmunt risk factors* 0 27 (35%) |
| 1 39 (50%) |
| 2 8 (10%) |
| 3 4 (5%) |
| Tumour PD-L1 expression Assessable 67 (86%) |
| <1% 42 (54%) |
| ≥1% 25 (32%) |
| <5% 53 (68%) |
| ≥5% 14 (18%) |
| Indeterminate, not assessable, or missing 11 (14%) |

Data are median (range) or number (%). Some percentages do not add up to 100 because of rounding. ECOG=Eastern Cooperative Oncology Group. *ECOG performance status >0, liver metastases, visceral involvement (defined as liver, lung, bone, or any non-lymph node), lymph-node-only involvement, and haemoglobin concentration <10 g/dL.

**Table 1: Baseline patient demographics and clinical characteristics**
change in size of target lesion from baseline. at baseline and at least one on-treatment tumour assessment. Dotted lines show 30% decrease, no change, and 20% increase in tumour burden over time in 74 response-evaluable patients.

Responders were response-evaluable patients with a complete or partial response as best overall response per Response Evaluation Criteria in Solid Tumors version 1.1. Response-evaluable patients were those with a target lesion at baseline and at least one on-treatment tumour assessment. Dotted lines show 30% decrease, no change, and 20% increase in size of target lesion from baseline.

Table 2: Antitumour activity

|                     | Nivolumab (n=78) | PD-L1 <5% (n=42) | PD-L1 ≥5% (n=25) |
|---------------------|------------------|------------------|------------------|
| Confirmed objective response | 19 (24.4%, 15:3–35:4) | 11 (26.2%, 13:9–42:0) | 6 (24.0%, 9:4–45:1) |
| Best overall response |                  |                  |                  |
| Complete response    | 5 (6%)           | 1 (2%)           | 4 (16%)          |
| Partial response     | 14 (18%)         | 10 (24%)         | 2 (8%)           |
| Stable disease       | 22 (28%)         | 11 (26%)         | 8 (32%)          |
| Progressive disease  | 30 (38%)         | 18 (43%)         | 8 (32%)          |
| Unable to establish  | 7 (9%)           | 2 (5%)           | 3 (12%)          |

Data are number (%), 95% CI or number (%). Some percentages do not add up to 100 because of rounding.

Table 2: Antitumour activity

Role of the funding source
The funder of the study provided the study drug and worked with the investigators to design the study and to collect, analyse, and interpret the data. All drafts of the report were prepared by the corresponding author with input from all co-authors and editorial assistance from professional medical writers, funded by the sponsor. All authors and professional medical writers had full access to all the data in the study and all authors had final responsibility for the decision to submit for publication.

Results
Between June 5, 2014, and April 24, 2015, 86 patients with metastatic urothelial carcinoma were enrolled in the nivolumab monotherapy group and 78 were treated with nivolumab monotherapy. At the time of data cutoff (March 24, 2016), 60 (77%) of 78 patients had discontinued treatment, 50 (64%) because of disease progression (figure 1), and 53 (68%) were continuing to be followed up. 18 (23%) of 78 patients switched to combination treatment with ipilimumab upon disease progression. At data cutoff, minimum follow-up was 9 months (median 15–2 months, IQR 12·9–16·8) and patients had received a median of 8·5 doses (range 1–46) of nivolumab.

Table 1 shows baseline demographics and clinical characteristics of all treated patients. Three patients who
previously refused standard treatment had not received previous platinum-containing chemotherapy in any setting. 71 (91%) of 78 patients had tumour tissue samples collected at baseline and PD-L1 expression could be measured in samples from 67 patients; 45 biopsies were obtained from a primary tumour site and 22 from a metastatic site. At least one Bellmunt risk factor (ECOG performance status >0, liver metastases, or haemoglobin concentration <10 g/dL, visceral involvement, or lymph-node-only involvement) was present in 51 (65%) of 78 patients.

A confirmed investigator-assessed objective response was achieved in 19 (24·4%, 95% CI 15·3–35·4) of 78 treated patients, with five patients (6%) achieving a complete response and 14 (18%) a partial response (table 2). Four patients were not assessable because they did not have at least one target lesion at baseline and at least one on-study assessment to meet response-evaluable criteria. Of 74 response-evaluable patients, 19 (25·7%) had a complete or partial response, resulting in an objective response rate of 25·7% (95% CI 16·2–37·2). Change in tumour burden over time for response-evaluable patients is shown in figure 2. 31 (40%) of 78 patients continued nivolumab monotherapy beyond progression, of whom nine were judged to have a non-conventional benefit (appendix p 4).

At data cutoff, the median duration of response was 9·4 months (IQR 5·7–12·5) and the median time to response was 1·5 months (1·2–4·1). Of 19 responders, 12 had an ongoing response (11 of whom were still on nivolumab monotherapy), three switched to nivolumab plus ipilimumab combination therapy (one of whom subsequently died), two continued on nivolumab monotherapy after progression, and two died (figure 3).

Median overall survival was 9·7 months (95% CI 7·3–16·2) and 46 (59%) of 78 patients had died at the time of data cutoff (figure 4A). 1-year overall survival was 46% (95% CI 34–56). In a post-hoc sensitivity analysis of overall survival excluding patients who switched to combination treatment, 1-year overall survival was 43% (95% CI 31–55). Median progression-free survival in the overall treated population was 2·8 months (95% CI 1·5–5·9) and 60 (77%) of 78 patients had disease progression or died by data cutoff (figure 4B). 1-year progression-free survival was 21% (95% CI 12–31).

An objective response was achieved in six (24%, 95% CI 9–45) of 25 patients with PD-L1 expression of at least 1%, and in 11 (26%, 14–42) of 42 patients with PD-L1 expression of less than 1% (table 2). Tumour reduction from baseline in target lesions is shown by PD-L1 expression in the appendix (p 6–7).

Median overall survival was 16·2 months (95% CI 7·6–not estimable) in patients with PD-L1 expression of at least 1% and 9·9 months (7·0–not estimable) in those with PD-L1 expression of less than 1% (table 2). Median progression-free survival was 5·5 months (95% CI 1·5–5·9) in patients with PD-L1 expression of at least 1% and 2·8 months (1·4–6·5) in those with PD-L1 expression of less than 1% (appendix p 9). Similar results were noted in patients with at least 5% PD-L1 expression (appendix p II).

The post-hoc analysis of objective response by Bellmunt prognostic risk factors is shown in table 3. Grade 3 or 4 treatment-related adverse events occurred in 17 (22%) of 78 patients (table 4). The most commonly reported grade 3 or 4 treatment-related adverse events were elevated lipase (four [5%]), elevated amylase
(three [4%]), and fatigue, maculopapular rash, dyspnoea, decreased lymphocyte count, and decreased neutrophil count (two [3%] each). Treatment-related adverse events of special interest potentially associated with the use of nivolumab of any grade were skin (33 [42%]), gastrointestinal (eight [10%]), renal (seven [9%]), hepatic (four [5%]), and pulmonary adverse events (two [3%]).

A post-hoc analysis of immune-mediated adverse events, regardless of causality, that occurred within 100 days of the last dose of nivolumab is shown in the appendix (p 10).

Serious adverse events were reported in 36 (46%) of 78 patients. Ten serious adverse events judged to be treatment related occurred in eight (10%) of 78 patients: colitis (grade 3–4), diarrhoea (grade 1–2), mouth ulceration (grade 1–2), nausea (grade 3–4), oral pain (grade 1–2), thrombocytopenia (patient died), fatigue (grade 3–4), hyponatraemia (grade 3–4), acute kidney injury (grade 3–4), and pneumonitis (patient died).

Two (3%) of 78 patients discontinued treatment because of treatment-related adverse events (grade 4 pneumonitis and grade 4 thrombocytopenia). Both events were fatal and are described in the appendix (p 5). 31 patients died of disease progression. Other than disease progression and treatment-related adverse events as causes of death, the following additional deaths were reported: three (4%) from unknown causes, and one (1%) due to sepsis that was not deemed related to study drug. Adverse events leading to treatment discontinuation but not deemed related to study drug were reported in four patients: one patient each had increased blood creatinine (grade 2), hepatitis C infection (grade 3), anaemia (grade 3), and urosepsis (grade 3). 28 (36%) of 78 patients had at least one dose delay; 52 doses were delayed from a total of 982 doses received (5%), 34 (65%) of which were delayed because of adverse events.

Discussion

In this study, nivolumab monotherapy was associated with a substantial and durable tumour response, promising survival, and acceptable safety in patients with locally advanced or metastatic urothelial carcinoma who were previously treated with platinum-based chemotherapy. Objective response was not dependent on tumour PD-L1 expression and was consistent across patient subgroups on the basis of key prognostic factors.

Two (3%) of 78 patients had a treatment-related event that led to death (one case of thrombocytopenia and one case of pneumonitis); both cases are described in greater detail in the appendix (p 5). "This patient subsequently died from pneumonitis and was counted in the grade 5 rather than the grade 4 total as a treatment-related death."

### Table 3: Objective response by Bellmunt prognostic risk factor

| Number of patients (%) | Grade 1–2 | Grade 3 | Grade 4 |
|------------------------|-----------|---------|---------|
| ECOG performance status|           |         |         |
| 0                      | 42 (26.2%, 13–9–42.0) |         |         |
| 1                      | 36 (22.2%, 10–1–39.2)   |         |         |
| Liver metastasis       |           |         |         |
| Yes                    | 20 (15.0%, 3–2–37.9)    |         |         |
| No                     | 58 (27.6%, 16–7–40.9)   |         |         |
| Visceral metastasis    |           |         |         |
| Yes                    | 66 (21.2%, 12–1–33.0)   |         |         |
| No                     | 12 (41.7%, 15–2–72.3)   |         |         |
| Lymph node only        |           |         |         |
| Yes                    | 11 (36.4%, 10–9–69.2)   |         |         |
| No                     | 67 (22.4%, 13–1–34.2)   |         |         |
| Haemoglobin concentration <10 g/dL |           |         |         |
| Yes                    | 11 (18.2%, 2–3–51.8)    |         |         |
| No                     | 67 (25.4%, 15–5–37.5)   |         |         |

**ECOG=Eastern Cooperative Oncology Group.**

### Table 4: Treatment-related adverse events of any grade reported in at least 10% of patients and all grade 3–4 events in 78 patients who received nivolumab

| Grade 1–2 | Grade 3 | Grade 4 |
|-----------|---------|---------|
| Any event | 46 (59%)| 17 (22%)| 0 |
| Fatigue   | 26 (33%)| 2 (3%)  | 0 |
| Pruritus  | 23 (29%)| 0       | 0 |
| Rash, maculopapular | 12 (15%)| 2 (3%)  | 0 |
| Lipase elevated | 7 (9%)  | 4 (5%)  | 0 |
| Nausea    | 9 (12%) | 1 (1%)  | 0 |
| Arthralgia| 9 (12%) | 0       | 0 |
| Anaemia   | 8 (10%) | 0       | 0 |
| Amylase increased | 4 (5%)  | 3 (4%)  | 0 |
| Dyspnoea  | 4 (5%)  | 1 (1%)  | 1 (1%) |
| Lymphocyte count decreased | 3 (4%)  | 2 (3%)  | 0 |
| Hyperglycaemia | 4 (5%)  | 1 (1%)  | 0 |
| Neutrophil count decreased | 1 (1%)  | 2 (3%)  | 0 |
| White blood cell count decreased | 2 (3%)  | 1 (1%)  | 0 |
| Hyponatraemia | 1 (1%)  | 1 (1%)  | 0 |
| Dermatitis acniform | 1 (1%)  | 1 (1%)  | 0 |
| Wheezing  | 1 (1%)  | 1 (1%)  | 0 |
| Acute kidney injury | 0       | 1 (1%)  | 0 |
| Aspartate aminotransferase increased | 0       | 1 (1%)  | 0 |
| Back pain | 0       | 1 (1%)  | 0 |
| Colitis   | 0       | 1 (1%)  | 0 |

Data are number of patients (%). Adverse events that occurred after patients crossed over from nivolumab 3 mg/kg to combination treatment were excluded.

Some patients had more than one adverse event. Two patients had a treatment-related event that led to death (one case of thrombocytopenia and one case of pneumonitis); both cases are described in greater detail in the appendix (p 5). "This patient subsequently died from pneumonitis and was counted in the grade 5 rather than the grade 4 total as a treatment-related death."
compare favourably with outcomes reported in a recent phase 2 clinical trial in urothelial carcinoma with the anti-PD-L1 antibody atezolizumab. The proportion of patients who achieved an objective response in our study seemed consistent across different prognostic risk factor subgroups, although sample sizes were small in these analyses, and in the post-hoc sensitivity analysis of overall survival excluding patients who switched to combination treatment, there was no effect on 1-year survival.

Nivolumab was also well tolerated in this population; the adverse-event profile compares favourably with previously tested chemotherapies and no new safety signals were noted compared with nivolumab studies across a wide range of tumour types. Additionally, in this cohort of patients who tend to have several comorbidities and impaired renal function, nivolumab was tolerated over several doses, suggesting a manageable tolerability profile in the longer term. Two patients discontinued treatment because of treatment-related adverse events (pneumonitis and thrombocytopenia); both events were subsequently fatal.

PD-L1 expression on tumour cells, as defined by the Dako immunohistochemical assay, was not associated with objective responses; patients whose tumours were defined as having at least 1% of tumour cells expressing PD-L1 achieved an objective response with a similar frequency as those whose tumours had less than 1% of tumour cells expressing PD-L1. Patients whose tumours had at least 1% of tumour cells expressing PD-L1 had a median overall survival of over 16 months, whereas those whose tumours had less than 1% of tumour cells expressing PD-L1 had a median overall survival of almost 10 months. Longer follow-up is needed to clarify whether this difference in median overall survival translates into long-term differences. Inter-tumour heterogeneity, both between primary and metastatic lesions and between different metastatic lesions, could have contributed to a false-negative PD-L1 result, although this is mitigated by the fact that patients were classified as having positive PD-L1 expression if they had any positive sample (ie, not necessarily the sample collected closest to the time of study drug treatment). Establishing whether these data are necessarily the sample collected closest to the time of tumour response with nivolumab according to immune cell PD-L1 expression is warranted.

Limitations of the study include the absence of a standard current practice comparator because of the scarcity of effective treatments in this setting, the small sample size, and the short follow-up period, which precluded further insights into the effect of nivolumab on long-term survival. Also, data on the efficacy and safety of treatment beyond progression, as well as protein expression scoring, were not available, but are being analysed and will be reported at a later date.

In summary, nivolumab monotherapy was active and safe in previously treated patients with locally advanced or metastatic urothelial carcinoma. These data suggest that this immune checkpoint therapy has activity in a patient population with limited treatment options.

Contributors
PSh and MKC conceived and designed the study. PSh, MKC, PB, JK, PSp, ECa, RNP, PAO, FdB, MM, DTL, DJ, ECh, and JER collected and assembled data. All authors analysed and interpreted data, and wrote and revised the manuscript.

Declaration of interests
PSh has received fees for advisory board participation for Jounce and Kite; fees for consultancy from Jounce, Kite, Bristol-Myers Squibb, AstraZeneca, and Amgen; and stock or stock options from Jounce and Kite. MKC has received grants from Bristol-Myers Squibb; consultancy fees from AstraZeneca and Moderna; and payment for lectures from Clinical Care Options. PB has received honoraria from Bristol-Myers Squibb, Pfizer, MSD, and Orion Pharma, and research funding from Novartis. RNP has received a travel grant from Bristol-Myers Squibb for an investigator meeting. PAO has received a grant from Bristol-Myers Squibb and consultancy fees from Bristol-Myers Squibb, Amgen, Celldex, Alexion, and Cytoora. FdB has received consultancy fees from Tizziana Life Sciences, Bristol-Myers Squibb, MSD, Servier, Eli Lilly, Merck Serono, GlaxoSmithKline, and Novartis, and speaker fees from Bristol-Myers Squibb, Eli Lilly, Roche, and ACCMED. MM has received consultancy fees from Eutibics and Boehringer Ingelheim, and speaker fees from Genentech, Novartis, Sanofi, Regeneron, Lexicon, Ipsen, Onyx, Bayer, Taiho, Merinimack, and Celgene. DJ and AA are employees of Bristol-Myers Squibb. ECh has received grants from Bristol-Myers Squibb and consultancy fees for advisory board participation from EMD Serono, Taiho, Bayer, Advaxis, Amgen, Lilly, and Castle Biosciences. CH, C-SL, and MT are employees of and hold stock options with Bristol-Myers Squibb. JER has received grants from Bristol-Myers Squibb, Novartis, and Roche/Genentech; has received consultancy fees from Roche/Genentech, AstraZeneca, Eli Lilly, Agensys, Sanofi US Services, Oncogenex, Onyx, Dendreon, Bristol-Myers Squibb, and Boehringer Ingelheim; and holds stock or stock options with Illumina and Merck. JK, PSp, ECa, and DTL declare no competing interests.

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