Efficacy and safety of avelumab treatment in patients with metastatic Merkel cell carcinoma: experience from a global expanded access program

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ABSTRACT

Background Avelumab, a human anti–programmed death-ligand 1 immunoglobulin G1 monoclonal antibody, showed favorable efficacy and safety in patients with metastatic Merkel cell carcinoma (mMCC) in the phase II JAVELIN Merkel 200 trial, leading to approval in multiple countries. We describe real-world experience with avelumab in patients with mMCC from an expanded access program.

Methods Eligible patients had mMCC and progressive disease during or after chemotherapy or were ineligible for chemotherapy or clinical trial participation. Patients received an initial 3-month supply of avelumab (administered as 10 mg/kg intravenously every 2 weeks until progressive disease or unacceptable toxicity); resupply was allowed following complete response, partial response, stable disease, or clinical benefit per physician assessment.

Results Between December 15, 2015, and March 4, 2019, 558 of 620 requests from 38 countries were medically approved, and 494 patients received avelumab. Among 240 evaluable patients, the objective response rate was 46.7% (complete response in 22.9%, including 3 of 16 potentially immunocompromised patients), and the disease control rate was 71.2%. The median duration of treatment in evaluable patients with response was 7.9 months (range, 1.0–41.7) overall and 5.2 months (range, 3.0–13.9) in immunocompromised patients. No new safety signals were identified. The expanded access program closed for new requests on December 31, 2018, as required after regulatory approval; benefitting patients continued to receive avelumab.

Conclusions The avelumab expanded access program for patients with mMCC demonstrated efficacy and safety in a real-world setting, consistent with the results from JAVELIN Merkel 200, and provided a treatment for patients with limited options.

BACKGROUND

Merkel cell carcinoma (MCC) is an aggressive skin cancer associated with a high risk of local and distant metastasis and poor survival outcomes in patients with metastatic disease (5-year survival rate with distant metastases of ≤20%).1-3 MCC is associated with clonal integration of the Merkel cell polyoma-virus (MCPyV), ultraviolet radiation exposure, immunosuppression, and advanced age.4,5 MCC is considered a chemosensitive tumor, but responses to chemotherapy in patients with metastatic McC (mMCC) are seldom durable.6-10 In addition, commonly used platinum-based chemotherapy regimens incur significant toxicity, particularly in a population that largely comprise older patients. There is evidence that impairment of the immune system can contribute to the pathogenesis of MCC, providing a rationale for therapies that promote antitumor immune responses.9

Avelumab is a human anti–programmed cell death protein-ligand 1 (PD-L1) immunoglobulin G1 monoclonal antibody that inhibits the interaction between PD-L1 and programmed cell death protein 1 (PD-1),11,12 which may induce T-cell reinvigoration and effector cell functions against tumor cells, as shown in preclinical models.13,14 Avelumab has shown antitumor activity and a tolerable safety profile in patients with various tumors,12,15-17 including patients who received first-line (1L) or second-line or later (2L+) therapy for mMCC.18-20 In mMCC, avelumab was initially assessed in part A of the JAVELIN Merkel 200 trial (NCT02155647), which enrolled 88 patients with mMCC and progressive disease (PD) while receiving or after receiving chemotherapy.18 After ≥2 years of follow-up in all patients, the objective response rate (ORR) was 33.0%, including a complete response (CR) in 11.4%, and the response duration was ≥2 years in 67.0% of responders based on Kaplan-Meier estimates.21
treatment-related adverse events (TRAEs) were observed in 11.4% of patients, and no treatment-related deaths occurred. Part B of JAVELIN Merkel 200 enrolled patients with mMCC who had not received prior systemic treatment for metastatic disease. In the primary analysis (n=116; ≥15 months of follow-up), the ORR was 39.7%, including CR in 16.4%, 30.2% had a durable response (lasting ≥6 months), and the median overall survival was 20.3 months. Grade ≥3 TRAEs were observed in 18.1% of patients, and no treatment-related deaths occurred. Based on the findings from JAVELIN Merkel 200, avelumab became the first treatment approved for mMCC in several locations, including the USA, European Union, Switzerland, Japan, Australia, Israel, Brazil, and Canada. An expanded access program (EAP; sometimes called a compassionate use program) provides an investigational treatment to patients who have a serious or terminal disease outside of a clinical trial, although definitions vary by country. After data became available from the avelumab registrational clinical trial in mMCC, the EAP was initiated to provide avelumab for compassionate use to patients with mMCC who had limited treatment options. The EAP was an ad hoc program that incorporated treatment guidelines but was not protocol-driven.

### Table 1 Baseline characteristics of approved patients in avelumab MCC EAP

| Characteristics | All approved patients (n=558) | Approved immunocompromised patients (n=37) | Approved 1L patients (n=36) |
|----------------|-------------------------------|------------------------------------------|----------------------------|
| **Age (years), n (%)** |                               |                                          |                            |
| <65             | 116 (20.8)                    | 6 (16.2)                                 | 4 (11.1)                   |
| ≥65             | 442 (79.2)                    | 31 (83.8)                                | 32 (88.9)                  |
| **Sex, n (%)**  |                               |                                          |                            |
| Male            | 372 (66.8)                    | 21 (56.8)                                | 24 (66.7)                  |
| Female          | 185 (33.2)                    | 16 (43.2)                                | 12 (33.3)                  |
| **Weight (kg), n (%)** |                               |                                          |                            |
| <80             | 229 (50.9)                    | 24 (64.9)                                | 16 (44.4)                  |
| ≥80             | 221 (49.1)                    | 13 (35.1)                                | 20 (55.6)                  |
| **Median (range)** | 78.8 (40–150)                | 75 (41–124)                              | 80 (49–138)                |
| **ECOG PS, n (%)** |                               |                                          |                            |
| 0               | 175 (39.0)                    | 14 (37.8)                                | 15 (41.7)                  |
| 1               | 233 (51.9)                    | 18 (48.7)                                | 18 (50.0)                  |
| 2               | 33 (7.4)                      | 3 (8.1)                                  | 3 (8.3)                    |
| 3               | 8 (1.8)                       | 2 (5.4)                                  | 0                          |

**Line of therapy, n (%)**

| Line of therapy | 1 | ≥2 |
|-----------------|---|----|
| 36 (6.5)        | 4 (10.8) | 36 (100) |
| 522 (93.6)     | 33 (89.2) | 0 |

*Weight and ECOG PS at baseline were available only for the 450 patients enrolled via the EAP portal. EAP, expanded access program; ECOG PS, Eastern Cooperative Oncology Group performance status; 1L, first-line; MCC, Merkel cell carcinoma.
A 3-component intervention to prevent serious complications of PD.avelumab, stable ECOG PS, and no delay to any immuno new or worsening of existing symptoms, tolerance to
including age, renal disorder
were ineligible for chemotherapy for various reasons,
patients received 1L avelumab after individual evaluation
received avelumab as 2L+ treatment, but a minority of
patients for several reasons, which included patients
was 73 years (range, 23–95) and 66.8% of patients were
Countries providing the most requests were Italy (n=109),
France (n=96), and Australia (n=83). The median age
was 73 years (range, 23–95) and 66.8% of patients were
male. Most patients (90.9%) had an ECOG PS of 0 or
1. Other available baseline characteristics are shown in
table 1. Among approved patients, 522 (93.6%) were
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ment on December 31, 2018, after regulatory approval of
avelumab in multiple countries.

RESULTS
Between December 15, 2015, and March 4, 2019 (data
cut-off date), 620 requests for avelumab were received
from 38 countries (online supplementary additional file
1, additional file 2). Of these requests, 558 were approved,
11 were withdrawn, and 51 were medically rejected for
various reasons, including incorrect diagnosis, lack of
appropriate prior therapy, and incomplete information.

Table 2

| Response parameter* | All patients (n=240) | Immunocompromised patients (n=16) | 1L (n=15) | 2L+ (n=225) |
|---------------------|---------------------|----------------------------------|-----------|-------------|
| ORR, %              | 46.7                | 37.5                             | 46.7      | 46.7        |
| DCR, %†             | 71.2                | 66.8                             | 66.7      | 71.6        |
| Confirmed BOR, n (%)|                      |                                  |           |             |
| CR                  | 55 (22.9)           | 3 (18.8)                         | 2 (13.3)  | 53 (23.6)   |
| PR                  | 57 (23.8)           | 3 (18.8)                         | 5 (33.3)  | 52 (23.1)   |
| SD                  | 59 (24.6)           | 5 (31.3)                         | 3 (20.0)  | 56 (24.9)   |
| PD‡                 | 69 (28.8)           | 5 (31.3)                         | 5 (33.3)  | 64 (28.4)   |
| Duration of treatment in patients with response§ | | | | |
| Median (range), months | 7.9 (1.0–41.7) | 5.2 (3.0–13.9) | 4.5 (3.0–19.8) | 7.9 (1.0–41.7) |

Immunocompromised, 1L, and 2L+ are subsets of the total number of evaluable patients.
*Response was reported according to treating physician assessment of follow-up scans at the time of resupply.
†Among patients treated for a minimum of 3 months with available data.
‡Patients with PD or AEs who required treatment discontinuation within the first 90 days were never resupplied and did not have a follow-up response evaluation; thus, these values may be under-reported.
§Duration of avelumab treatment/drug supply is reported as a surrogate for duration of response or clinical benefit.

A series of requests were processed for avelumab in multiple countries.

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Of 494 patients supplied with avelumab, starting from
April 6, 2016, response data were not available in 254
patients for several reasons, which included patients
whose entry predated the implementation of the EAP
portal and who did not have outcome data, regard-
less of response or resupply, and patients who had no

**Procedures**
Patients received avelumab 10 mg/kg by 1-hour intrave-
nous infusion every 2 weeks until confirmed PD, unac-
ceptable toxicity, or occurrence of any other criterion for
withdrawal. Continuation of avelumab beyond radiolog-
ical PD was permitted in the absence of significant clinical
deterioration and based on physician assessment of poten-
tial benefit on a case-by-case basis, with criteria including
no new or worsening of existing symptoms, tolerance to
avelumab, stable ECOG PS, and no delay to any immi-
ent intervention to prevent serious complications of PD.
A 3-month supply of avelumab was provided to treating
physicians, and resupply was allowed for patients who had
CR, partial response (PR), stable disease, or other clinical
benefit according to radiography or treating physician
assessment. Data were provided by the treating physician
during the application for avelumab resupply. Antitumor
activity was assessed by the treating physician per treat-
ment guidelines and according to Response Evaluation
Criteria in Solid Tumors 1.1 where applicable to deter-
mine best overall response. Other assessments included
duration of treatment for patients with response, safety,
and tolerability. Adverse events (AEs) were reported to
Merck KGaA Global Patient Safety. An online portal to
process EAP requests and collate responses was imple-
ment in May 2017.
evaluable data beyond the 90-day period of initial drug supply (including those with rapid PD). Outcomes were therefore provided for 240 patients. Among patients who discontinued treatment, the most common reasons were radiographic or clinical progression. Of the 240 evaluable patients, 16 were classified as potentially immunocompromised based on available medical history (examples included patients who had chronic lymphocytic leukemia, were HIV-positive, had received a prior organ transplant, or were receiving high-dose steroids), and 15 received 1L avelumab (patients ineligible for chemotherapy).

Duration of avelumab treatment (or duration that the drug was supplied) was considered a surrogate for duration of response or clinical benefit because resupply was dependent on these criteria. At data cut-off, the median duration of avelumab treatment was 7.9 months (range, 1.0–41.7) among all evaluable patients (n=240), 5.2 months (range, 3.0–13.9) in immunocompromised patients (n=16), and 4.5 months (range, 3.0–19.8) in 1L (chemotherapy-ineligible) patients (n=15). In total, 55 patients had CR and 57 had PR, representing an ORR of 22.7% using the denominator of 494 patients in the total population. Considering only the 240 response-evaluable patients, the ORR was 46.7%, including CR in 22.9% and PR in 23.8%. In immunocompromised patients, the ORR was 37.5% (n=6), including CR in 18.8% (n=3; these patients had a history of acute myeloid leukemia, had undergone splenectomy, or had Sjögren’s syndrome). In the 1L subgroup, the ORR was 46.7% (n=7) (table 2). Images of responses in individual patients are shown in figure 1.

Physician-reported safety data from the EAP are summarized in table 3. Infusion-related reaction, fever, fatigue, and rash were the most frequently occurring TRAEs. Safety events were likely under-reported because data were reported at the treating physician’s discretion at the time of resupply, and many patients had no evaluable data beyond the 3-month supply.

**Discussion**

The global MCC EAP for avelumab answered an urgent, unmet medical need for patients with mMCC, a rare cancer with limited treatment options; to our knowledge, this is the largest EAP performed for this patient population. Before 2017, no treatment was approved for mMCC by the US Food and Drug Administration, European Commission, or any other regulatory authority. In this large EAP population, avelumab provided a clinical benefit in both immune-competent and immunocompromised patients (based on available medical history). The physician-reported ORR was 46.7% among all evaluable patients, including three immunocompromised patients who achieved CR and three who achieved PR, and the ORR was consistent in 1L and 2L+ subgroups. Durable responses were observed in both immune-competent and immunocompromised patients. Importantly, no new safety signals were identified in the EAP population. In both the EAP and JAVELIN Merkel 200 study, infusion-related reaction, fatigue, and rash were among the most frequently occurring TRAEs. Infusion-related reactions were managed per established guidance for avelumab.

The data summarized in this manuscript have some limitations. Safety and efficacy data for the EAP were reported at the treating physician’s discretion and are therefore potentially under-reported in this program. Furthermore, because resupply of avelumab was only provided to patients following an assessment of clinical benefit by the treating physician, this may have affected reporting. Additionally, patients with PD or an AE who required treatment discontinuation within the first 90 days did not receive a resupply of avelumab and subsequently were not provided with a follow-up response evaluation. Limited information was provided by clinicians about each patient’s medical history, and...
for this rare aggressive cancer. has revealed the need for additional treatment options safety profile of avelumab in a real-setting, this EAP is the largest factors that excluded participation in clinical trials. Efficacy and safety data were consistent with pivotal trial data. the classification of patients as immunocompromised was made without specific assessments or knowledge of ongoing medications. Nevertheless, these findings highlight that in a real-world setting, avelumab showed efficacy and safety consistent with results from the JAVELIN Merkel 200 trial, despite including patients who would have been ineligible for the trial (e.g., those with ECOG PS of 2 or 3, treated brain metastases, or immunosuppressive conditions including HIV infection). Although the results presented are limited by the nature of data reporting within the MCC EAP, these results provide valuable insight into responses to anti-PD-1/PD-L1 antibodies in vulnerable or immunocompromised patients with this disease.

EAPs enable patients with limited treatment options to access therapies that they otherwise would not receive. To the best of our knowledge, this MCC EAP is the largest and only EAP for an immune checkpoint inhibitor in this rare disease. In addition to evaluating the efficacy and safety profile of avelumab in a real-world setting, this EAP has revealed the need for additional treatment options for this rare aggressive cancer.

In conclusion, avelumab provided benefit in a real-world setting, and avelumab showed efficacy and safety consistent with results from the JAVELIN trial. This EAP was conducted in accordance with the Declaration of Helsinki and the International Council for Harmonisation Guideline for Good Clinical Practice. The protocol was approved by the institutional review board or independent ethics committee of each center, and all patients provided written informed consent before enrollment.

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Correction notice Since the online publication of this article, the authors have noticed an error in Table 1 and 2. In Table 1, the header ‘Approved immunocompromised patients (n=33)’ should read ‘Approved immunocompromised patients (n=37)’ as the total number of patients is 37. Additionally in Table 2 row ‘CR’, column ‘1L (n=15)’, the data value is incorrectly stated as ‘5 (33.3%)’ (13.3%). The correct data value is ‘2 (13.3%)’.

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Patient consent for publication Not required.

Ethics approval The trial was conducted in accordance with the Declaration of Helsinki and the International Council for Harmonisation Guideline for Good Clinical Practice. The protocol was approved by the institutional review board or independent ethics committee of each center, and all patients provided written informed consent before enrollment.

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after January 1, 2014, Merck KGaA, Darmstadt, Germany will share patient-level and study-level data after deidentification, as well as redacted study protocols and clinical study reports from clinical trials in patients. These data will be shared with qualified scientific and medical researchers, upon researchers’ request, as necessary for conducting legitimate research. Such requests must be submitted in writing to the company’s data sharing portal. More information can be found at https://www.merckgroup.com/en/research/our-approach-to-research-and-development/healthcare-clinical-trials/commitment-responsible-data-sharing.html, Where Merck KGaA has a coresearch, coderevelopment or co-marketing/copromotion agreement or where the product has been out-licensed, it is recognized that the responsibility for disclosure may be dependent on the agreement between parties. Under these circumstances, Merck KGaA will endeavor to gain agreement to share data in response to requests.

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Correction: Efficacy and safety of avelumab treatment in patients with metastatic Merkel cell carcinoma: experience from a global expanded access program

Walker JW, Lebbé C, Grignani G, et al. Efficacy and safety of avelumab treatment in patients with metastatic Merkel cell carcinoma: experience from a global expanded access program. *J Immunother Cancer* 2020;8:e000313. doi: 10.1136/jitc-2019-000313.

Since the online publication of this article, the authors have noticed errors in Table 1 and 2. In Table 1, the header 'Approved immunocompromised patients (n=33)' should read 'Approved immunocompromised patients (n=37)' as the total number of patients is 37. Additionally in Table 2 row ‘CR’, column ‘1L (n=15)’, the data value is incorrectly stated as ‘5 (33.3) 2 (13.3)’. The correct data value is ‘2 (13.3)’.

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