Tumor mutational burden predicts the efficacy of pembrolizumab monotherapy: a pan-tumor retrospective analysis of participants with advanced solid tumors

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ABSTRACT

Background Several studies have evaluated the relationship between tumor mutational burden (TMB) and outcomes of immune checkpoint inhibitors. In the phase II KEYNOTE-158 study of pembrolizumab monotherapy for previously treated recurrent or metastatic cancer, high TMB as assessed by the FoundationOne CDx was associated with an improved objective response rate (ORR).

Methods We retrospectively assessed the relationship between TMB and efficacy in participants with previously treated advanced solid tumors enrolled in 12 trials that evaluated pembrolizumab monotherapy, including 3 randomized trials that compared pembrolizumab with chemotherapy. TMB was assessed in formalin-fixed, paraffin-embedded pretreatment tumor samples by whole-exome sequencing. High TMB was defined as ≥175 mutations/exome. Microsatellite instability (MSI) phenotype was based on whole-exome sequencing results. Programmed death ligand 1 (PD-L1) expression was assessed by immunohistochemistry. The primary end point was ORR assessed per RECIST V.1.1 by independent central review. Other end points included progression-free survival (PFS) assessed per RECIST V.1.1 by independent central review and overall survival (OS).

Results Of the 2234 participants in the analysis, 1772 received pembrolizumab monotherapy and 462 received chemotherapy. Among the pembrolizumab-treated participants. ORR was 31.4% (95% CI 27.1 to 36.0) in the 433 participants with TMB ≥175 mutations/exome and 9.5% (95% CI 8.0 to 11.2) in the 1339 participants with TMB <175 mutations/exome. The association of TMB with ORR was observed regardless of PD-L1 expression and not driven by specific tumor types or participants with very high TMB or high MSI. In the 3 randomized controlled trials, TMB was associated with ORR (p=0.016), PFS (p=0.005), and OS (p=0.029) of pembrolizumab but not of chemotherapy (p≥0.340, p≥0.643, and p=0.174, respectively), and pembrolizumab improved efficacy versus chemotherapy in participants with TMB ≥175 mutations/exome.

Conclusions TMB ≥175 mutations/exome is associated with clinically meaningful improvement in the efficacy of pembrolizumab monotherapy and improved outcomes for pembrolizumab versus chemotherapy across a wide range of previously treated advanced solid tumor types. These data suggest TMB has broad clinical utility irrespective of tumor type, PD-L1 expression, or MSI status and support its use as a predictive biomarker for pembrolizumab monotherapy in participants with previously treated advanced solid tumors.

BACKGROUND

The accumulation of neoantigens, defined as abnormal peptides arising from mutations in genes that encode proteins presented by the major histocompatibility complex, increases the likelihood that tumor cells will be recognized by infiltrating immune cells. Somatic tumor mutational burden (TMB) may result in neoantigen formation with the potential for immune system recognition.1 Therefore, agents that promote T-cell activation such as monoclonal antibodies against programmed death 1 (PD-1) or its ligand, PD-L1, may be particularly effective anticancer therapies in patients with high TMB. Several studies have shown positive correlations between TMB levels and response to immune checkpoint inhibitors across diverse tumor types.2-4

The anti-PD-1 monoclonal antibody pembrolizumab received accelerated approval from the US Food and Drug Administration (US FDA) for the treatment of adult and pediatric patients with unresectable or metastatic solid tumors with tumor mutational burden-high (TMB-H) (≥10 mutations/megabase as determined by an FDA-approved test) that have progressed following prior treatment and who have no satisfactory alternative...
treatment options. The FoundationOne CDx (Foundation Medicine, Cambridge, Massachusetts, USA), a targeted cancer gene panel, was contemporaneously approved as the TMB companion diagnostic for pembrolizumab. Approval was primarily based on a prespecified analysis of patients with select previously treated advanced solid tumors treated with pembrolizumab monotherapy in the phase II KEYNOTE-158 study. The objective response rate (ORR) was 29% in the 102 participants with TMB-H tumors per FoundationOne CDx and 6% in the 688 participants with non-TMB-H tumors. TMB was predictive of pembrolizumab outcomes irrespective of microsatellite instability (MSI) status, tumor PD-L1 expression, and tumor type.

To explore the value of TMB for predicting response to pembrolizumab monotherapy in cancer types not included in KEYNOTE-158, we assessed the relationship between TMB measured by whole-exome sequencing (WES) and efficacy using individual patient data from 12 additional trials of pembrolizumab monotherapy. Results of this supportive analysis, which used a WES cutpoint that maximized agreement to the 10-mutations/megabase FoundationOne CDx cutpoint and were submitted to and reviewed by the FDA as part of the application seeking regulatory approval of pembrolizumab for previously treated TMB-H advanced solid tumors, are presented herein.

METHODS
Trial design, participants, and treatment
Key eligibility criteria and full protocols for the trials included in this retrospective analysis of participants with previously treated advanced solid tumors have been published. Pembrolizumab was administered intravenously at a dose of 2 mg/kg, 10 mg/kg, or 200 mg once every 3 weeks or 10 mg/kg once every 2 weeks (online supplemental table S1). The chemotherapy regimens in the randomized trials varied by tumor type. Where permitted by local laws and regulations, broad genetic/genomic testing, including WES, was included as a mandatory element of 10 of 12 trials included in this analysis; in KEYNOTE-009 and KEYNOTE-010, genetic/genomic testing was optional. Only samples provided by properly consented participants were sent for testing.

Genomic profiling
WES was performed on formalin-fixed, paraffin-embedded pretreatment tumor samples using ImmunoSELECT-RUO (Personal Genome Diagnostics, Baltimore, Maryland, USA) or ACE Cancer Exome (Personalis, Menlo Park, California, USA). After pathology assessment and using a fresh scalpel, the tissue was scraped from the entire section and transferred to a 1.5 mL tube containing 200 µL of 100% ethanol; if the section contained <20% tumor, the tissue was macrodissected from the marked tumor area. DNA was isolated using the QIAamp DNA FFPE Tissue Kit (Qiagen, Valencia, California, USA). Tumor DNA was quantitated using the Qubit assay (Invitrogen, Carlsbad, California, USA); quality was assessed using the QuantideX qPCR DNA QC Assay (Assuragen, Austin, Texas, USA). WES was performed on matched normal DNA from whole blood collected in a PAXgene DNA Tube (Qiagen) at clinical sites and stored at ~20°C or ~70°C/80°C until processed in an approved central laboratory identified by the sponsor. The Chemagic STAR DNA Blood Kit (PerkinElmer, Waltham, Massachusetts, USA) run on either a Hamilton Chemagic STAR or PerkinElmer Chemagic 360 automated instrument was used to extract DNA in a final volume of 500 µL or 1.0 mL. Extracted DNA was subjected to volume and concentration determination and ultraviolet and visible spectral analysis to assess purity.

The WES bioinformatics pipeline for analyses across the pembrolizumab translational program has been previously described. WES reads were aligned to the Genome Reference Consortium Human Build 37 using Burrows-Wheeler Aligner MEM (V.0.7.12) followed by preprocessing steps including duplicate marking, indel realignment, and base recalibration with Picard (V.1.114) and Genome Analysis Toolkit (V.2) to generate analysis-ready binary alignment map (BAM) files. For the 89% of tumor samples for which matched normal DNA was available, MuTect (V.1.1.7) was used to generate somatic single nucleotide variant (SNV) calls using default parameters by comparing BAM files from tumor and matched normal samples. For the 11% of tumor samples without matched normal DNA, MuTect (V.1.1.7) was used to call SNVs, which were then analyzed by PureCN (V.1.6.3) to classify germline and somatic status; WES data from 50 platform-matched normal samples were used to build a pool of normal samples to do sequencing error and alignment artifact filtering and copy number normalization. MuTect-called SNVs present in the Single Nucleotide Polymorphism database (V.141) but not in the Catalogue of Somatic Mutations in Cancer (V.68) were filtered out. SNVs with mutant reads <4 in tumor samples were also eliminated. The mean coverage was 147 reads per base for tumor and 107 reads per base for normal samples; median coverage was 134 and 105 reads per base, respectively. Overall, 2199 of 2234 tumor samples (99.4%) had coverage >20 reads per base. Somatic mutations were annotated with Variant Effect Predictor (V.78), and non-synonymous mutations in protein coding regions were counted for TMB. TMB for individual participants was defined as the number of somatic non-synonymous SNVs and indels that passed all described filters. TMB of participants with tumor DNA only was highly concordant with TMB of participants with both tumor and matched normal DNA for a set of 505 tumors analyzed with both pipelines (Spearman’s correlation 0.96, with no substantial offset in intercept or slope). There was little association between computational tumor purity and TMB (Spearman’s correlation 0.22).

The FoundationOne CDx was performed at the Clinical Laboratory Improvement Amendments-certified laboratory of Foundation Medicine using the Dx1 baitset. TMB was calculated by counting the number of synonymous and non-synonymous mutations across an 0.8 Mb
region spanning 324 genes, with computational germline status and oncogenic driver filtering.

Participants were evaluable for TMB if their samples provided valid TMB scores on sequencing and analysis. Participants were non-evaluable for TMB if they did not provide any tumor tissue, if they did not consent to exploratory biomarker research, if the amount of tumor tissue was inadequate for analysis, if the samples did not yield sufficient DNA from either tumor tissue or matched normal DNA for WES, or if the samples failed sequencing and subsequent quality control steps.

Expression of an 18-gene T-cell inflamed gene expression profile (GEP) was assessed using the NanoString nCounter gene expression platform (NanoString Technologies, Seattle, Washington, USA) as described by Ayers et al. in RNA isolated from formalin-fixed, paraffin-embedded tumor samples (processed as outlined above for WES). MSI genotype was detected by applying mSINGs on WES data from tumor samples. The stability of each mononucleotide microsatellite locus was evaluated, and the proportion of unstable microsatellite loci of each mononucleotide microsatellite locus was evaluated, and the proportion of unstable microsatellite loci of each mononucleotide microsatellite locus was determined as the MSI score. MSI was confirmed by PCR using the Promega MSI Analysis System, V.1.2. in the 304 pembrolizumab-treated participants from KEYNOTE-059 and KEYNOTE-061 included in the analysis.

**Histology**

PD-L1 expression was assessed by immunohistochemistry in formalin-fixed tumor samples using either the PD-L1 IHC 22C3 pharmDx assay (Agilent, Carpinteria, California, USA) or a laboratory-developed prototype assay based on the 22C3 antibody (Qualtex, Goleta, California, USA). The definition of positivity using the PD-L1 IHC 22C3 pharmDx assay for all tumors except melanoma and non-small-cell lung cancer was combined positive score (CPS) ≥ 1, where CPS was defined as the number of PD-L1-expressing tumor cells, lymphocytes, and macrophages divided by the total number of viable tumor cells, multiplied by 100. For melanoma, positivity was defined as Allred proportion score ≥ 2, which is equivalent to membranous PD-L1 expression in ≥ 1% of tumor and tumor-associated immune cells; for non-small-cell lung cancer, positivity was defined as tumor proportion score (TPS) ≥ 1%, where TPS was defined as the percentage of PD-L1-expressing tumor cells out of the total number of viable tumor cells. The definition of positivity using the prototype assay was membranous PD-L1 expression on ≥ 1% of tumor and associated inflammatory cells or positive staining in the stroma.

**Statistical analysis**

Individual participant data from all pembrolizumab monotherapy trials that included patients with previously treated advanced solid tumors for which WES data were available at the time the TMB regulatory submission package was prepared in 2020 were assembled as supportive evidence and reviewed by the FDA. Data from the KEYNOTE-002 chemotherapy group were excluded because there were too few participants with available TMB for reliable between-arm treatment comparisons. Data from KEYNOTE-199 cohort were excluded because participants had bone-predominant metastatic castration-resistant prostate cancer, precluding them from being assessed for the primary end point of this analysis. The analysis population included participants with available TMB scores who received ≥ 1 dose of study treatment and received prior treatment for their cancer.

Among a range of cutoffs showing clear enrichment for objective response, the data for pembrolizumab monotherapy by evaluating the relationship between TMB and inflammation in the tumor microenvironment as assessed using an 18-gene T-cell inflamed GEP using a method similar to that of Panda et al. The WES cutoff that maximized agreement to the 10-mutations/megabase FoundationOne CDx cutoff was assessed in participants in the training set who had TMB assessed by both methods.

Statistical analyses were performed using SAS V.9.4 (SAS Institute, Cary, North Carolina, USA). The primary end point was ORR (ie, proportion of participants with confirmed complete or partial response) assessed per RECIST V.1.1 by independent central review. Other end points were duration of response (DOR, ie, time from initial response to disease progression or death in participants with complete or partial response) and progression-free survival (PFS, ie, time from randomization (first dose of pembrolizumab in non-randomized trials) to disease progression or death) assessed per RECIST V.1.1 by independent central review and overall survival (OS, ie, time from randomization (first dose of pembrolizumab in non-randomized trials) to death). The Kaplan-Meier method was used to estimate DOR, PFS, and OS. ORR, DOR, PFS, and OS were summarized descriptively for participants with TMB ≥ 175 and < 175 mutations/exome. For analyses by tumor type, tumor types represented by ≤ 10 participants were pooled into a single category termed ‘other’. Although our focus was to provide clinical findings for the 175 mutations/exome WES TMB cutoffs that were aligned with the 10-mutations/megabase FoundationOne CDx cutoff, we have also provided a profile of ORR in patients with TMB above a range of progressively higher cutoffs, beginning with 0 and increasing in increments of 25 mutations/exome, for the pooled dataset along with pointwise 95% CIs; at least 50 participants must have had TMB above the cut-off for estimating ORR and the CI. The impact of key baseline characteristics on the robustness of the relationship between TMB status and response to pembrolizumab was assessed using logistic regression analysis that included the following covariates: TMB status (≥ 175 vs < 175 mutations/exome), PD-L1 expression status (positive vs negative), MSI status (high vs non-high), age (< 65 vs ≥ 65 years), sex (female vs male), Eastern Cooperative Oncology Group (ECOG) performance status (0 vs 1 or 2), number of lines of prior therapy for advanced disease (≤ 1 vs ≥ 2), and tumor...
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type (model parameterized as difference relative to gastric cancer).

The treatment effect of pembrolizumab versus chemotherapy was assessed in the 3 randomized studies9 15 18 by comparing ORR, DOR, PFS, and OS in participants with TMB ≥175 and <175 mutations/exome. To evaluate associations between TMB and efficacy by treatment, logistic regression analyses adjusted for ECOG performance status were performed separately in participants treated with pembrolizumab and in participants treated with chemotherapy for each randomized study; two-sided p values (not adjusted for multiplicity) are presented as a measure of the strength of association between each efficacy end point and TMB assessed as a continuous, log_{10}-transformed variable.

All genomic and clinical data were provided to and reviewed by the FDA as part of the application seeking regulatory approval of pembrolizumab for previously treated TMB-H advanced solid tumors.

Reanalysis of data from Samstein et al

Data from Samstein et al26 were downloaded from http://www.cbioportal.org/study?id=tmb_mskcc_2018 in September 2019. The only publicly available data were for the 1662 participants treated with an immune checkpoint inhibitor as monotherapy or in combination. Cox regression analyses associating high versus low TMB with OS across tumor types, within tumor types, and at various TMB cutpoints were executed in MATLAB (MathWorks, Natick, Massachusetts, USA) and HRs and 95% CIs were reported. Because data for estrogen receptor status were not provided in the downloaded dataset, we could not perform analyses for estrogen receptor-positive and estrogen receptor-negative breast cancer. Using the published tumor-specific cutoffs, we observed HRs and 95% CIs that qualitatively appeared identical to those reported by Samstein et al.26 After this verification, we analyzed the data using a single, pan-tumor cutpoint of 10 mutations/megabase and plotted the resultant HRs and 95% CIs against those we obtained using the tumor-specific cutoffs (online supplemental figure S1).26

RESULTS
Cutpoint verification

We verified WES TMB 175 mutations/exome as the optimal cutpoint in distinct analyses. One analysis used a training set that included the first 976 participants with previously treated cancer for whom TMB was obtained, of whom 810 also had GEP data. We found that WES TMB 175 mutations/exome enriched for response to pembrolizumab and was associated with peak statistical significance for differences in inflammation in the tumor microenvironment as measured by the 18-gene T-cell inflamed GEP (figure 1A). In the second analysis, which included the 338 participants in the training set who had TMB assessed by both WES and FoundationOne CDx, there was high concordance between TMB assessed by both methods (Spearman’s correlation, 0.71), and 175 mutations/exome was the WES cutpoint that maximized average positive and negative agreement with the 10-mutations/megabase FoundationOne CDx cutpoint (area under the receiver-operating characteristics curve, 0.92) (figure 1B).

Figure 1 Validation of the optimal cutpoint for TMB assessed by WES. (A) Comparison of inflammation in the tumor microenvironment assessed using an 18-gene T-cell inflamed GEP in participants with TMB above versus below potential WES cutpoints for mutational burden. P values are two-sided and assessed using the Wilcoxon rank-sum test. The vertical dotted line represents TMB 175 mut/exome. (B) The optimal TMB cut-off assessed by WES that corresponds with the FoundationOne CDx TMB cutpoint of 10 mut/megabase. The Youden index of 175 mut/exome assessed per WES is the point of maximal average positive and negative agreement rates with 10 mut/megabase by the FoundationOne CDx. The area under the receiver-operating characteristic curve is 0.92. GEP, gene expression profile; mut, mutations; TMB, tumor mutational burden; WES, whole-exome sequencing.
Clinical utility of the cutpoint

To assess the clinical utility of the 175-mutations/exome cutpoint, we explored efficacy in a pooled population of participants with previously treated cancer enrolled in 12 clinical trials—9 single-arm trials of pembrolizumab monotherapy and 3 randomized trials that compared pembrolizumab monotherapy with chemotherapy and enrolled participants between 2011 and 2017 (online supplemental table S1). Including the 876 eligible participants from the training set, 2234 participants representing 24 tumor types—1772 who received pembrolizumab monotherapy and 462 who received chemotherapy—were included in the WES TMB analysis population (online supplemental figure S2). Median (IQR) follow-up duration, defined as time from the first dose of study treatment to data cut-off, was 26.1 months (20.1–32.9). Among 1772 pembrolizumab-treated participants, 433 (24.4%) had TMB $\geq$ 175 mutations/exome and 1339 (75.6%) had TMB $<$ 175 mutations/exome (online supplemental table S2). Among 462 chemotherapy-treated participants, 140 (30.3%) had TMB $\geq$ 175 mutations/exome and 322 (69.7%) had TMB $<$ 175 mutations/exome (online supplemental table S2). The prevalence of TMB $\geq$ 175 mutations/exome among tumor types represented by $>10$ participants ranged from 4.1% (ovarian cancer) to 56.6% (melanoma) (online supplemental table S2). The most prevalent tumor types in the TMB $\geq$ 175 and $<$ 175 mutations/exome populations were non-small-cell lung cancer and ovarian cancer, respectively (figure 2). Aside from imbalances in sex, baseline characteristics were generally similar in the TMB $\geq$ 175 and $<$ 175 mutations/exome populations (online supplemental table S3). Twenty-one (1.2%) participants, all of whom had TMB $\geq$ 175 mutations/exome, had MSI-high (MSI-H) tumors per WES. More participants in the $\geq$ 175 vs $<$ 175 mutations/exome population completed (7.4% vs 1.6%) or remained on (8.5% vs 4.1%) pembrolizumab at the time of data cut-off (online supplemental table S4).

In the pooled pembrolizumab population, we found that participants with TMB $\geq$ 175 mutations/exome had better outcomes than those with TMB $<$ 175 mutations/exome. ORR was 31.4% (95% CI 27.1 to 36.0) in participants with TMB $\geq$ 175 mutations/exome and 9.5% (95% CI 8.0 to 11.2) in those with TMB $<$ 175 mutations/exome (online supplemental table S5). Reductions from baseline in target lesion size were observed in 67.5% and 41.5% of participants, respectively (online supplemental figure S3). Excluding participants from the training set, ORR was 26.3% (95% CI 20.3 to 33.0) and 10.2% (95% CI 8.0 to 12.7) in the $\geq$ 175 mutations/exome (n=198) and $<$ 175 mutations/exome (n=698) populations, respectively. In the pooled population of 1772 participants, ORR increased as the WES TMB cutpoint was increased (online supplemental figure S4).

We confirmed that the response enrichment provided by TMB $\geq$ 175 mutations/exome was robust and not driven by associations with other confounding variables using subgroup analyses, participant-level data, and logistic regression. TMB $\geq$ 175 mutations/exome enriched for response in each of the 8 tumor types that included $>10$ participants and in the pooled set of 16 tumor types that included $\leq$ 10 participants each (figure 3A). Participant-level data showed that the overall association of TMB with ORR was not driven by outliers with very high TMB or by specific tumor types (figure 3B). The ORR was higher among participants with TMB $\geq$ 175 mutations/exome regardless of PD-L1 status (figure 3C) or MSI genotype (figure 3D). In a multivariate logistic regression analysis, the OR for ORR in participants with TMB $\geq$ 175 vs $<$ 175 mutations/exome was 3.076 (95% CI 2.243 to 4.220).

Figure 2  Proportion of participants in the pooled pembrolizumab population with whole-exome sequencing TMB $\geq$ 175 mut/exome and TMB $<$ 175 mut/exome by tumor type. The tumor types are presented in order of descending median tumor mutational burden. For a listing of tumor types included in the ‘other’ category, see online supplemental table S2. HNSCC, head and neck squamous cell carcinoma; mut, mutations; NSCLC, non-small-cell lung cancer; TMB, tumor mutational burden; TNBC, triple-negative breast cancer.
after adjusting for the effects of other prognostic variables (table 1).

Among responders, median DOR was not reached in the TMB ≥175 mutations/exome population and was 12.5 months in the TMB <175 mutations/exome population; 63.8% and 32.2% of responses, respectively, were estimated to last for ≥24 months (figure 4A). Median PFS was 4.2 months (95% CI 3.6 to 5.8) in participants with TMB ≥175 mutations/exome and 2.1 months (95% CI 2.0 to 2.1) in participants with TMB <175 mutations/exome (figure 4B). Median OS was 15.5 months (95% CI 13.1 to 17.7) and 10.1 months (95% CI 9.4 to 10.9), respectively; the estimated percentage of participants alive at 24 months was 40.0% and 23.2% (figure 4C).

To further assess the association between TMB and outcomes, we used exploratory regression models and data from the 3 randomized clinical trials, which enrolled patients with non-small-cell lung cancer (KEYNOTE-010),9 urothelial cancer (KEYNOTE-045),15 and gastric cancer (KEYNOTE-061).18 In these trials, TMB as a continuous log_{10}-transformed variable was significantly associated with ORR (p≤0.016), PFS (p≤0.005), and OS (p≤0.029) in the pembrolizumab groups but not the chemotherapy groups (p≥0.340, 0.643, and 0.174, respectively) (online supplemental table S6). The benefit of pembrolizumab versus chemotherapy was enhanced in participants with TMB ≥175 mutations/exome in all 3 randomized trials (table 2; figure 5, online supplemental figure S5, S6).

**DISCUSSION**

In this retrospective analysis of individual participant data from 12 clinical trials across 24 tumor types that contributed to the US FDA approval of pembrolizumab monotherapy for the treatment of patients with unresectable or metastatic TMB-H solid tumors that progressed following prior treatment and who have no satisfactory treatment options, we found that TMB ≥175 mutations/exome assessed by WES was predictive of outcomes to pembrolizumab. TMB ≥175 mutations/exome enriched for response to pembrolizumab across tumor types, including those with generally high TMB like melanoma and those with generally low TMB like ovarian cancer.27 28 TMB ≥175 mutations/exome was also associated with longer DOR, PFS, and OS, although pooled OS estimates should be interpreted with caution given the different representation of tumor
types with differing prognosis in the TMB ≥175 and <175 mutations/exome populations. Our finding that TMB was consistently and significantly associated with efficacy of pembrolizumab but not of chemotherapy and that the relative benefit of pembrolizumab versus chemotherapy was enhanced in participants with TMB ≥175 mutations/exome across the 3 randomized trials included in our analysis supports the clinical utility of TMB as a predictive biomarker for pembrolizumab monotherapy.

Importantly, the association of TMB with response in our study was not driven by outliers with very high TMB, by the minority of participants with MSI-H tumors, or by specific tumor types. Although ORR was higher among participants with PD-L1-positive tumors, TMB ≥175 mutations/exome enriched for response regardless of PD-L1 expression. After adjusting for key baseline demographics and disease characteristics, including PD-L1 expression, the adjusted OR for response to pembrolizumab in participants with TMB ≥175 versus <175 mutations/exome was 3.076, confirming that the enrichment in response provided by high TMB is robust and not driven by associations of confounding variables. Together with a growing body of evidence suggesting that TMB and PD-L1 expression are not correlated but are jointly informative for clinical outcomes in patients treated with anti-PD-1 or anti-PD-L1 monotherapy,29-33 and the findings from KEYNOTE-158 showing that high TMB expression is predictive of response to pembrolizumab in a PD-L1-unselected population,34 our data support the use of TMB as a clinically meaningful biomarker regardless of PD-L1 expression.

We verified that TMB 175 mutations/exome assessed by WES is well aligned with the FoundationOne CDx TMB cutoff of 10 mutations/megabase that is known to enrich for response,34 is the cut-off agreed on by a multi-stakeholder group convened by the Friends of Cancer Research to harmonize on a standard cut-off for clinical trials,25 and is the cutpoint used in the KEYNOTE-158 TMB analysis.35 The high concordance we observed between TMB assessed by WES and the FoundationOne CDx suggests that molecular biomarker discoveries made using exploratory techniques such as WES may have clinical utility.

Table 1: Multivariate logistic regression analysis of response to pembrolizumab assessed per RECIST V.1.1 by independent central review in the pooled pembrolizumab population

<table>
<thead>
<tr>
<th>Covariate</th>
<th>OR for ORR (95% CI)</th>
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<tr>
<td>TMB status (≥175 vs &lt;175 mutations/exome)</td>
<td>3.076 (2.243 to 4.220)</td>
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<tr>
<td>PD-L1 expression status (positive vs negative)</td>
<td>2.187 (1.442 to 3.316)</td>
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<tr>
<td>MSI status (high vs non-high)</td>
<td>3.798 (1.453 to 9.925)</td>
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<td>Age (≤65 vs &gt;65 years)</td>
<td>0.707 (0.525 to 0.952)</td>
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<td>Sex (female vs male)</td>
<td>1.004 (0.713 to 1.413)</td>
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<td>ECOG performance status</td>
<td>1.471 (1.094 to 1.978)</td>
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<tr>
<td>No. of lines of prior therapy for advanced disease (≥1 vs ≤2)</td>
<td>1.237 (0.898 to 1.705)</td>
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<td>Tumor type (HNSCC vs gastric*)</td>
<td>1.841 (1.057 to 3.207)</td>
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<td>Tumor type (melanoma vs gastric*)</td>
<td>3.664 (1.877 to 7.153)</td>
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<td>Tumor type (NSCLC vs gastric*)</td>
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<td>Tumor type (ovarian vs gastric*)</td>
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<td>Tumor type (prostate vs gastric*)</td>
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<td>Tumor type (TNBC vs gastric*)</td>
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<tr>
<td>Tumor type (urothelial vs gastric*)</td>
<td>2.224 (1.291 to 3.832)</td>
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</table>

*Gastric cancer was chosen as the reference because, alphabetically, it was the first tumor type group included in the analysis.
ECOG, Eastern Cooperative Oncology Group; HNSCC, head and neck squamous cell carcinoma; MSI, microsatellite instability; NSCLC, non-small-cell lung cancer; ORR, objective response rate; PD-L1, programmed death ligand 1; TNBC, triple-negative breast cancer.
be successfully translated to smaller cancer gene panels like FoundationOne CDx, which are more easily implemented in routine clinical practice and may be more cost-effective. 35 36

While other studies also show that a single TMB cutpoint enriches for outcomes of checkpoint inhibitors regardless of tumor type, 9 24 37–40 there are reports that suggest tumor-specific cutpoints are needed. 26 40–42 When we reanalyzed the publicly available data from one such study, published by Samstein et al, 26 we found that a 10-mutations/megabase cutpoint was associated with improved OS across tumor types, with HRs for death generally consistent with those based on the originally reported tumor-specific cutpoints (online supplemental figure S1). The results of our study and our reanalysis of the Samstein et al data show that although the distribution of TMB scores varies across tumor types, a single cutpoint that provides clinically meaningful enrichment of response to PD-1 blockade across tumor types can be identified. This highlights the importance of distinguishing between the conditional distribution of response associated with a biomarker and the marginal distribution of the biomarker itself. Identification of a single cutpoint with broad clinical utility is valuable for fully realizing the initial diagnostic implementation of TMB as a predictive biomarker.

One limitation of our analysis is the inclusion of 876 participants from the training set in the overall pooled pembrolizumab population (n=1772). However, the ORR in the TMB ≥175 and <175 mutations/exome populations was similar regardless of whether participants from the training set were included in the analysis. Another limitation is the inclusion of tumors without matched normal DNA in our analyses; the low frequency of these samples and the high concordance of TMB derived from the tumor-only and matched-normal pipelines suggests that their inclusion did not bias the analysis. Although the data set includes tumor types for which pembrolizumab is already approved and could be biased by the tumor types for which pembrolizumab trials are available and the participants in the trials for whom WES data were available, we believe that a population of almost 2500 participants that includes 24 tumor types with a mix of those associated with high TMB (eg, melanoma, non-small-cell lung cancer, and urothelial cancer) and those associated with generally low TMB (eg, triple-negative breast, ovarian, and prostate cancers) is robust and supports the ability of TMB to predict response to pembrolizumab regardless of tumor type. While formal statistical testing of the predictive value of TMB was not specified, logistic regression analysis that included key baseline demographics and disease characteristics was performed, and the resultant OR estimates confirm the robustness of our findings.

In conclusion, our data show that a single TMB cutpoint determined by WES that has broad clinical utility irrespective of tumor type, PD-L1 expression, or MSI genotype can be identified, mapped to a targeted cancer gene panel, and used to define a population that derives clinically meaningful benefit from pembrolizumab monotherapy. Our data support TMB as a predictive biomarker for pembrolizumab and the FDA approval of pembrolizumab monotherapy for the treatment of patients with unresectable or metastatic, TMB-H solid tumors that have progressed following prior treatment and who have no satisfactory alternative treatment options.

Acknowledgements The authors would like to thank the patients and their families and caregivers for participating in the clinical trials; the investigators...
Ethics approval
All clinical trial participants provided written informed consent before enrollment. All study protocols were consistent with the global standards of the International Conference on Harmonization Good Clinical Practices, the Council for International Organizations of Medical Public Policy Statement: Clinical Trial Ethics Sciences International Ethical Guidelines for Biomedical Research Involving Human Subjects, the Pharmaceutical Research and Manufacturers of America Principles on Conduct of Clinical Trials, applicable local regulatory requirements, and the ethical principles that have their origin in the Declaration of Helsinki.

Provenance and peer review
Not commissioned; externally peer reviewed.

Data availability statement
Data are available on reasonable request. Merck Sharp & Dohme Corp, a subsidiary of Merck & Co, Inc, Kenilworth, New Jersey, USA (MSD) is committed to providing qualified scientific researchers access to anonymized data and clinical study reports from the company’s clinical trials for the purpose of conducting legitimate scientific research. MSD is also obligated to protect the rights and privacy of trial participants and, as such, has a procedure in place for evaluating and fulfilling requests for sharing company clinical trial data with qualified external scientific researchers. The MSD data sharing website (available at: http://engagezone.msd.com/ds_documentation.php) outlines the process and requirements for submitting a data request. Applications will be promptly assessed for completeness and policy compliance. Feasible requests will be reviewed by a committee of MSD subject matter experts to assess the scientific validity of the request and the qualifications of the requestors. In line with data privacy legislation, submitters of approved requests must enter into a standard data-sharing agreement with MSD before data access is granted. Data will be made available for request after product approval in the USA and European Union or after product development is discontinued. There are circumstances that may prevent MSD from sharing requested data, including country-specific or region-specific regulations. If the request is declined, it will be communicated to the investigator. Access to genetic or exploratory biomarker data requires a detailed, hypothesis-driven statistical analysis plan that is collaboratively developed by the requester and MSD subject matter experts; after approval of the statistical analysis plan and execution of a data-sharing agreement, MSD will either perform the proposed analyses and share the results with the requester or will construct biomarker covariates and add them to a file with clinical data that is uploaded to an analysis portal so that the requester can perform the proposed analyses.

Supplemental material
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REFERENCES


