Clinical utility of tumor mutational burden in patients with non-small cell lung cancer treated with immunotherapy

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Abstract: Anti-programmed death (ligand)-1 [anti-PD-(L)1] therapies such as pembrolizumab, nivolumab or atezolizumab have become standard of care for non-small cell lung cancer (NSCLC) patients either in first line or beyond. PD-L1 expression level allows enriching the treated population with responders, but it is still not an optimal biomarker. Even in patients with PD-L1 tumor proportion score (TPS) levels of ≥50% treated with first line pembrolizumab overall response rate (ORR) is only 44.8% and overall survival at one year is 70%. Moreover, in combination trials with chemotherapy and anti-PD-(L)1 therapy, a significant proportion of patients does not respond (ORR ranges from 45.3% to 64.0%), regardless of PD-L1 expression. Furthermore, PD-L1 expression level is not associated with improved benefit in patients treated with combinations of anti-PD-(L)1 and anti-cytotoxic T-lymphocyte-associated antigen (anti-CTLA4) therapy. One of the new promising biomarkers is tumor mutational burden (TMB). It has been discovered that especially tumor types with a known high mutation rate such as NSCLC and melanoma respond best to immune checkpoint inhibitors (ICIs). An explanation is that this high mutation rate makes it more likely that neoantigens arise that are targeted by activated immune cells, but it is not feasible to determine neoantigen load in daily practice. However, TMB of a certain tumor type is associated with neoantigen load and outcome on ICIs. In this comprehensive review, we discuss the TMB analysis methods, the rationale to use TMB as a predictive biomarker and the clinical utility of TMB in NSCLC patients treated with ICIs.

Keywords: Immunotherapy; mutation/immunology; non-small cell lung cancer (NSCLC); predictive value of tests; programmed death-ligand 1 (PD-L1)

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Introduction

In less than a decade, immune checkpoint inhibitors (ICIs) have become standard of care in the treatment of advanced non-small cell lung cancer (NSCLC), either in first line (alone in selected patients or in the near future in combination in others) or beyond. Pembrolizumab, a programmed death-1 (PD-1) inhibitor is registered in first line for advanced NSCLC patients, without an activating epidermal growth factor receptor (EGFR) mutation or anaplastic lymphoma kinase (ALK) aberration, but with a programmed death-ligand 1 (PD-L1) tumor proportion score (TPS) of ≥50%. It is also registered in second line for those with a PD-L1 TPS of ≥1%. This
outcome of ICI treatment. Until now, besides PD-L1, only mismatch repair (MMR) deficiency has been Food and Drug Administration (FDA) approved as a predictive biomarker (12). Moreover, there are several issues on PD-L1 testing such as the use of different antibodies, different platforms and different cut-offs for positivity that are extensively reviewed elsewhere (13).

One of the new promising biomarkers is tumor mutational burden (TMB). In this comprehensive review, we discuss the rationale to use TMB as a predictive biomarker, the analysis methods and the clinical utility of TMB in NSCLC patients treated with ICI.

**TMB definition and TMB assessment methods**

TMB has first been determined with whole exome sequencing (WES). Currently, WES seems to be less feasible in daily practice as it is costly, time consuming, labour intensive and needs large sequencing capacity (14,15). TMB can also be determined through hybrid-capture based targeted next generation sequencing [large next-generation sequencing (NGS) panel—NGS laboratory developed test (LDT)], for example using the FoundationOne® assay (first version of F1 LDT test) or the FoundationOne CDx™ assay (F1 CDx™, current FoundationOne assay) (16-18). In contrast to WES, large panel NGS is increasingly used in daily practice, usually with a limited number of genes. TMB is defined as the total number of somatic mutations in the tumor exome (17). There are different ways to estimate the total number of somatic mutations. The parameters are the type of mutations, the minimal allelic frequency and the strategy of excluding germline variants. In WES, the germline variants are oftenly excluded with a WES platform and different cut-offs for positivity that are extensively reviewed elsewhere (13).

TMB definition and TMB assessment methods
towards WES results increases significantly (16). Depending on the study in which a FoundationOne® assay (F1 LDT or F1 CDx™) is used, high TMB is defined as ≥9.9 to 20 mut/Mb (7,9,21-24). A good correlation for TMB determined by WES and large panel NGS was also found for other assays. For example, another study compared WES to a 500-gene targeted NGS panel including various cancer-relevant genes. Both approaches correlated linearly (regression equation =0.96), and the NGS panel achieved a sensitivity of >90% and a specificity of >85% compared to WES when classifying tumors as TMB high. Cutoff for TMB high values with NGS was 15 mut/Mb in this assay (25). The Memorial Sloan Kettering-Integrated Mutation Profiling of Actionable Cancer Targets (MSK-IMPACT) is another example with a panel of 341 (version 1) to 468 genes (version 3). TMB determined by the MSK-IMPACT assays (versions 1–3 used in this study) was highly correlated to WES (Spearman ρ=0.86; P<0.001), patients were considered TMB high when they had >7.4 mut/Mb (14). There are no published studies comparing the performance between each of the different large panel NGS but harmonization efforts are ongoing (26,27).

As an alternative to tissue, a recent method of TMB evaluation is exploring TMB directly in circulating tumoral DNA (ctDNA) from blood/plasma (bTMB). In one study, bTMB was determined using a 394-gene panel and was compared to tissue TMB (FoundationOne® assay, not mentioned whether this was F1 LDT or F1 CDx™) and to the FoundationACT (FACT) dedicated to ctDNA assay (including only 62 genes). A total of 259 patients were evaluable for both bTMB and tissue TMB. Overall agreement and positive percent agreement (PPA) were 81.5% and 63.6% respectively when using the 394-gene panel for bTMB. However, when the FACT assay was compared to tissue TMB, PPA dropped to 17%, probably due to the low number of genes included in the panel. However, the performance on variant detection was similar when overlapping allele regions were compared: 93% of variants were detected in both assays (28).

Causes of high TMB

There are multiple causes for somatic mutations in cancer such as defective DNA repair, intrinsic small errors in the DNA replication machinery, enzymatic modification of DNA and exogenous exposures (29). When a cell divides, multiple errors occur, mainly at the sites of microsatellites: most of these errors are corrected during “proofreading” (30,31). The MMR genes correct the remaining errors (32). Mutations in these damage repair genes can occur as germline mutation and as somatic mutations, both result in hypermutated tumors with a microsatellite instability-high (MSI-high) phenotype (33). Some colorectal cancers (e.g., the ones occurring in relation to the Lynch syndrome) are associated with a defect DNA MMR (34). Examples of exogenous exposures causing somatic mutations are tobacco smoking in lung cancer and ultraviolet light in skin cancers (35).

Variance of TMB number is high between and within tumor types (36-39). Tumors that are often found to have a high TMB are melanoma, NSCLC, bladder, oesophagus, colon, head and neck, gastric and endometrial cancers (38,40,41). A high TMB has been associated with smoking or a molecular smoking signature in lung cancer and head and neck cancer (38,40,41). Another dominant signature associated with a high TMB in lung cancer is the apolipoprotein B mRNA editing enzyme, catalytic polypeptide-like (APOBEC) signature (38,40). Stage of disease (I/II versus III/IV) is not correlated to TMB (41).

Several other mutations such as POLD1, POLE, MSH2, POLED2A, KEAP1, PAPPA2, PXDNL, RYR1, SCN8A and SLIT3 have been associated with a high TMB, however with the exception of KEAP1 mutations they seem rare in NSCLC (16,42,43).

Tobacco smoke carcinogens cause direct DNA damage and increase cancer risk by increasing the somatic mutation load as compared to non-smokers for whom the presence of a driver oncogene will be sufficient for the tumorigenesis (44,45).

Most driver mutations are found in non-smoking NSCLC patients and these patients have a lower TMB. In a large dataset including 1,775 EGFR mutated, 489 ALK/ROS1 rearranged, 286 MET exon 14 mutated, 493 v-Raf murine sarcoma viral oncogene homolog B (BRAF) mutated and 3,155 kirsten rat sarcoma viral oncogen (KRAS) mutated NSCLC patients, mean TMB and proportion of patients with a TMB of >10 or 20 mut/Mb was significantly lower for EGFR, ALK/ROS1 and MET exon 14 mutated patients compared to KRAS mutated patients. Mean TMB ranged from 3.1 to 6.2 mut/Mb for these patients compared to 10.3 mut/Mb for KRAS mutated patients. Mean TMB was similar for BRAF mutated compared to KRAS mutated patients (9.7 versus 10.3 mut/Mb), and TMB of these patient groups was comparable to all adenocarcinoma patients (mean TMB 9.1 mut/Mb). Squamous cell carcinoma patients had a slightly higher mean TMB (11.3).
TMB was evaluated through a large NGS panel (24). Updated results (using instead of mean values the median values, median being the correct measure to use in a skewed population such as the TMB samples) including over 15,000 analyzed lung cancer cases confirmed these results, except for the lower median TMB in BRAF mutated patients (3.8 mut/Mb) (46).

A smaller series of 83 patients evaluated for TMB through the FoundationOne® assay (not mentioned whether F1 LDT or F1 CDx™) confirmed the lower TMB in EGFR and ALK positive patients but found also a lower TMB in KRAS mutated patients when EGFR, ALK and KRAS mutated patients were compared to driver mutation negative patients (47). The lower TMB of EGFR mutated tumors compared to wildtype tumors was confirmed in another series of 108 resected early stage lung adenocarcinomas (48). In lung adenocarcinoma, it was found that number of mutations or neoantigens was higher in tumors with high PD-L1 and high CD8A/cytolytic activity (CYT) compared to tumors with low PD-L1 and low CD8A/CYT, the latter being an immune ignorant TME (49). In contrast to driver mutated NSCLCs with mostly a low TMB, pulmonary sarcomatoid tumors were found to have a high TMB in one retrospective series: 41% of the 122 included sarcomatoid tumors had a TMB of >10 mut/Mb (in a large panel NGS) and 18% had a TMB of >20 mut/Mb in one study (50). Large-cell neuroendocrine carcinoma of the lung (LCNEC) has a TMB that seems similar to that reported of adenocarcinoma patients when compared to the adenocarcinoma TMB in the series of Spigel et al. although in the latter mean was provided instead of median (median 9.9 versus mean 9.1 mut/Mb) (24,51).

**Relevance of TMB in immunotherapy treated patients**

The goal of ICI treatment is that the patient’s immune system is stimulated to reject the tumor and by doing this induces a long-lasting tumor response. There are several steps necessary to achieve this. Among them, the tumor should express neoantigens (tumour-specific, mutated peptides presented on the surface of cancer cells) that should be adequately processed and presented by the major histocompatibility complex (MHC) and afterwards these neoantigens should be recognized by T-cells as non-self to achieve an adequate anti-tumor T-cell response (52-55). The higher the number of neoantigens in a certain tumor, the higher the chance that a T-cell response is initiated to one or more of these neoantigens. It has been shown that the number of candidates neoantigens per tumor correlates well with TMB (56,57).

**TMB and ICI outcomes in advanced NSCLC trials**

Several studies using samples of patients included in clinical trials as well as retrospective series reported ICI outcome for NSCLC patients in relation to TMB. These are described below and summarized in *Table 1*.

**Trial data**

**Pembrolizumab**

One of the first series demonstrating a superior clinical efficacy of ICI in NSCLC patients with high TMB was published in 2016. TMB was evaluated with WES, 2 independent cohorts of patients treated with pembrolizumab were sequenced (N=16 and N=18). A median of 209 nonsynonymous mutations was identified (range, 11–1,192). High TMB was defined as TMB above the median burden of the cohort (209 in discovery, 200 in validation cohort). In the discovery cohort 73% of patients with high TMB had durable clinical benefit compared to only 13% of those with low TMB (P=0.04). ORR was 63% versus 0% in those with high versus low TMB, respectively (P=0.03), progression free survival (PFS) was 14.5 versus 3.7 months, respectively [hazard ratio (HR) 0.19; 95% CI: 0.05–0.70; P=0.01]. Similar results were obtained in the validation cohort (56).

**Atezolizumab**

In a retrospective pooled analysis of the FIR and BIRCH trials (both single arm first line and beyond atezolizumab trials, including patients selected for PD-L1 positivity) a PFS benefit and a trend to an OS benefit was found in TMB high patients from the FIR and BIRCH trial. In total, 417 out of 797 patients were included in the biomarker evaluable population (BEP). TMB was determined in tissue with the F1 LDT panel. Patients with a TMB above the median (9.9 mut/Mb) had a PFS HR of 0.49 compared to those with a TMB below the median in first line (total of 87 patients in BEP) and a PFS HR of 0.64 in second line and beyond (total of 330 patients in BEP). HR for OS was 0.71 and 0.86, respectively [both not significant (NS)]. With a TMB high cutoff at the 75% quantile (16.2 mut/Mb), HR for PFS in first and second line/beyond was 0.7 (NS).
Table 1 Outcome of NSCLC patients treated with immune checkpoint inhibitors in relation to tumor mutational burden

<table>
<thead>
<tr>
<th>Author, year (retro- or prospective)</th>
<th>NR of TMB evaluable NSCLC patients</th>
<th>Treatment</th>
<th>TMB test</th>
<th>Definition TMB high</th>
<th>TMB associated with PD-L1 status</th>
<th>ORR high vs. low TMB</th>
<th>DCB high vs. low TMB</th>
<th>PFS high vs. low TMB</th>
<th>OS high vs. low TMB</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Samples used from trial patients</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rizvi, 2015 (retrospective; unclear which trials)</td>
<td>16 discovery, 18 validation</td>
<td>Pembro</td>
<td>WES</td>
<td>&gt;200 (median) mutations</td>
<td>NA</td>
<td>63% vs. 0%</td>
<td>73% vs. 13%</td>
<td>mPFS: 14.5 vs. 3.7 mo (HR 0.19)</td>
<td>NA</td>
</tr>
<tr>
<td>Kowanetz, 2016 (retrospective; FIR, BIRCH, POPLAR)</td>
<td>509</td>
<td>Atezo (± vs. doc)</td>
<td>F1 LDT</td>
<td>≥9.9 mut/Mb weak</td>
<td>NA</td>
<td>NA</td>
<td>F1 LDT: 1st line, PFS HR 0.49; 2nd line, PFS HR 0.64; POPLAR*: mPFS 7.3 vs. 2.8 mo (HR 0.49)</td>
<td>F1 LDT: 1st line, OS HR 0.71; 2nd line, OS HR 0.86; POPLAR*: mOS 16.2 vs. 8.3 mo (HR 0.48)</td>
<td></td>
</tr>
<tr>
<td>Gandara, 2017 (retrospective; POPLAR, OAK)</td>
<td>794</td>
<td>Atezo vs. doc</td>
<td>bTMB 394-gene panel</td>
<td>≥10 mut/Mb</td>
<td>No</td>
<td>NA</td>
<td>NA</td>
<td>PFS HR*; OAK 0.73; POPLAR 0.68</td>
<td>OS HR*; OAK 0.69; POPLAR 0.59</td>
</tr>
<tr>
<td>Carbone, 2017 (retrospective; CheckMate 026)</td>
<td>312</td>
<td>Nivo vs. platinum-doublet chemo</td>
<td>WES</td>
<td>≥243 mutations</td>
<td>No</td>
<td>47% vs. 28%*</td>
<td>NA</td>
<td>mPFS* 9.7 vs. 5.8 mo (HR 0.62)</td>
<td>NA</td>
</tr>
<tr>
<td>Hellmann, 2017 (retrospective; CheckMate 012)</td>
<td>75</td>
<td>Nivo + ipi</td>
<td>WES</td>
<td>&gt;158 (median) mutations</td>
<td>No</td>
<td>51% vs. 13%**</td>
<td>65% vs. 34%**</td>
<td>mPFS** 17.1 vs. 3.7 mo (HR 0.41)</td>
<td>NA</td>
</tr>
<tr>
<td>Ramalingam, 2018 (prospective; CheckMate 568)</td>
<td>98</td>
<td>Nivo + ipi</td>
<td>F1 CDxTM</td>
<td>≥10 mut/Mb</td>
<td>No</td>
<td>44% vs. 40%**</td>
<td>NA</td>
<td>mPFS 7.1 vs. 2.6 mo (HR NA)</td>
<td>NA</td>
</tr>
<tr>
<td>Hellmann, 2018 (prospective; CheckMate 227)</td>
<td>299</td>
<td>Nivo + ipi vs. platinum-doublet chemo or nivo</td>
<td>F1 CDxTM</td>
<td>≥10 mut/Mb</td>
<td>No</td>
<td>45.3% vs. 26.9%*</td>
<td>NA</td>
<td>mPFS* 7.2 vs. 5.5 mo (HR 0.58)</td>
<td>NA</td>
</tr>
<tr>
<td>Samples used from patients treated in daily practice or unknown</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Spigel, 2016 (retrospective; unknown)</td>
<td>64</td>
<td>Nivo, atezo or pembro</td>
<td>FoundationOne®</td>
<td>≥15 mut/Mb</td>
<td>NA</td>
<td>NA</td>
<td>DoT 64 vs. 17 weeks (HR 0.40)</td>
<td>NA</td>
<td>NA</td>
</tr>
</tbody>
</table>

Table 1 (continued)
<table>
<thead>
<tr>
<th>Author, year (retro- or prospective)</th>
<th>NR of TMB evaluable NSCLC patients</th>
<th>Treatment</th>
<th>TMB test</th>
<th>Definition TMB high</th>
<th>TMB associated with PD-L1 status</th>
<th>ORR high vs. low TMB</th>
<th>DCB high vs. low TMB</th>
<th>PFS high vs. low TMB</th>
<th>OS high vs. low TMB</th>
</tr>
</thead>
<tbody>
<tr>
<td>Park, 2017 (retrosp.; daily practice)</td>
<td>36</td>
<td>Nivo</td>
<td>NA</td>
<td>≥20 mut/Mb</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>mOS** NR vs. 10.3/12.4 mo (HR NA)</td>
</tr>
<tr>
<td>Singal, 2017 (retrosp.; daily practice)</td>
<td>444</td>
<td>Nivo</td>
<td>FoundationOne®</td>
<td>≥20 mut/Mb</td>
<td>NA</td>
<td>NA</td>
<td>DoT 7.5 vs. 4.6 mo (HR NA)</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Ross, 2017 (retrosp.; daily practice)</td>
<td>1,619</td>
<td>Not specified</td>
<td>FoundationOne®</td>
<td>≥20 mut/Mb</td>
<td>Weak</td>
<td>NA</td>
<td>High TMB predictive for longer DoT** (HR NA)</td>
<td>NA</td>
<td>BA</td>
</tr>
<tr>
<td>Rizvi, 2018 (retrosp.; 78% treated in daily practice)</td>
<td>240</td>
<td>Anti-PD(L)1 + anti-CTLA4</td>
<td>MSK-IMPACT assay</td>
<td>&gt;7.4 mut/Hb</td>
<td>No</td>
<td>NA</td>
<td>38.6% vs. 25.1% PFS HR 1.38 (low vs. high TMB)</td>
<td>NA</td>
<td>NA</td>
</tr>
</tbody>
</table>

*high TMB checkpoint inhibitor vs. high TMB chemotherapy instead of high versus low TMB when treated with checkpoint inhibitor; **, high TMB vs. low/intermediate TMB when treated with checkpoint inhibitor instead of high versus low TMB when treated with checkpoint inhibitor; *, not mentioned in abstract whether it was F1 LDT or F1 CDx™ assay. NSCLC, non-small cell lung cancer; TMB, tumor mutational burden; PD-L1, programmed death-ligand 1; pembro, pembrolizumab; WES, whole exome sequencing; NA, not available; ORR, objective response rate; mPFS, median progression free survival; mo, months; HR, hazard ratio; DCB, durable clinical benefit; atezo, atezolizumab; doc, docetaxel; bTMB, blood tumor mutational burden; mut, mutations; Mb, megabase; OS, overall survival; nivo, nivolumab; chemo, chemotherapy; ipi, ipilimumab; DoT, duration of therapy; NR, not reached; CTLA4, cytotoxic T-lymphocyte-associated antigen; LDT, laboratory developed test.
and 0.55, respectively. HR for OS was 0.35 and 0.77 (NS) respectively. Comparable results were found for the patients in the POPLAR trial (second line and beyond atezolizumab versus docetaxel, no PD-L1 selection). In the BEP for this trial, a total of 92 out of 287 patients were included. Median PFS was 7.29 months for atezolizumab and 2.83 months for docetaxel for patients with a TMB above the median (HR 0.49; 95% CI: 0.25–0.93). Median PFS was 8.48 and 2.83 months respectively when a TMB high cutoff at the 75% quantile was used (HR 0.49; 95% CI: 0.19–1.3). Median OS for atezolizumab versus docetaxel was 16.23 versus 8.31 months in the TMB > median population (HR 0.48; 95% CI: 0.23–1.04) and not reached versus 12.16 months in the TMB >75% quantile population (HR 0.50; 95% CI: 0.15–1.67). In the pooled analysis, TMB was weakly correlated with PD-L1 expression but not with tumor infiltrating lymphocytes (TIL). The highest ORR was found in the patient population with high TMB and PD-L1 expression (ORR just below 40%) (21).

As is clear from the limited number of BEP patients in the atezolizumab trials, obtaining enough tissue for large panel NGS or WES is challenging. As described above, TMB can also be evaluated in blood (bTMB) and this was evaluated in an exploratory analysis of the OAK and POPLAR trials (794 out of 1,137 patients were evaluable). To determine the bTMB, 394 genes from plasma cell-free DNA were interrogated for single nucleotide variants (SNVs) and a score was based on the number of high-confidence SNVs identified. bTMB did not correlate with PD-L1, but improved PFS and OS were seen with atezolizumab compared to docetaxel for patients with a bTMB of ≥10 in both the OAK and POPLAR trial [PFS HR 0.73 and 0.68, respectively; overall survival (OS) HR 0.69 and 0.59, respectively] (58).

bTMB is currently prospectively tested in the first line single-arm phase II B-FI1RST of atezolizumab monotherapy for NSCLC patients (NCT 02848651). Co-primary endpoints are ORR and PFS in the bTMB negative and positive subgroups. Arm C of the phase II/III B-FAST trial (NCT03178552) evaluates atezolizumab monotherapy versus platinum-doublet chemotherapy in advanced NSCLC patients. Primary endpoint is PFS.

**Nivolumab ± ipilimumab**

The first TMB data for nivolumab came from the CheckMate 026 trial (6). In this first line randomised phase III trial, stage IV/recurring NSCLC patients with a PD-L1 expression of ≥1% were randomised to nivolumab monotherapy or platinum-doublet chemotherapy. The trial was negative for its primary endpoint (PFS in the group of patients with a PD-L1 expression of ≥5%). However, in an exploratory analysis, TMB determined with WES correlated with outcomes. Low TMB was defined as 0–100 mutations, median as 100–242 mutations and high as 243 or more. A total of 312 out of 541 (58%) randomized patients were evaluable for TMB and a slightly higher percentage of patients with a high TMB was found in the chemotherapy group compared to the nivolumab group (39% versus 30%). ORR as well as median PFS were higher in high TMB patients treated with nivolumab compared to high TMB patients treated with chemotherapy. ORR was 47% and 28% respectively, median PFS was 9.7 versus 5.8 months, respectively (HR 0.62; 95% CI: 0.38–1.00). Median OS was not significantly different but cross-over rate was high (68%). No correlation was found between TMB and PD-L1 status, but those with high TMB as well as high PD-L1 (≥50%) had, when treated with nivolumab, a superior ORR (75%) compared to those with only one of the markers defined as high (ORR 32–34%) or when both markers were classified as low (ORR 16%) (6).

Similar results were found in an exploratory analysis of arm G the CheckMate-012 study (nivolumab + ipilimumab). Totally, 75 of the 197 included patients had sufficient tumor tissue and paired blood for WES. ORR, durable clinical benefit (DCB) and PFS were significantly higher in those with a TMB above the median [158] compared to those with a TMB below the median. ORR was 51% versus 13% (P=0.0005), DCB was 65% versus 34% (P=0.011), median PFS was 17.1 months for those with a high TMB compared to 3.7 months for those with a low TMB (HR 0.41; 95% CI: 0.23–0.73; P=0.0024). PD-L1 expression was not correlated to TMB (P=0.804). Patients with high TMB as well as PD-L1 expression ≥1% had the best ORR and PFS (57).

In the CheckMate 568 trial, a single-arm phase II study of first line nivolumab plus ipilimumab in advanced NSCLC, the investigators sought to identify an appropriate TMB cut-off to select patients for this first line ICI combination. Primary endpoint was ORR, one of the secondary endpoints was ORR by TMB. In contrast to the Checkmate 026 trial, TMB was determined through large panel NGS with the validated F1 CDx™ assay. 288 patients were included, and overall ORR was 27%. 98 patients (34%) were evaluable for TMB. ORR plateaued with the threshold of ≥10 mutations/Mb. ORR was 4% with TMB <5, 10% with TMB <10, 44% with TMB ≥10 and 39% with TMB ≥15 mut/Mb. With receiver operating characteristic (ROC)
Median PFS was not significantly different and was 3.2 (95% CI: 2.7–4.3) and 5.5 (95% CI: 4.3–5.6) months, respectively (HR 1.07; 95% CI: 0.84–1.33). Interestingly, and in contrast to the CheckMate 026 trial, PFS was not superior for nivolumab versus chemotherapy in the TMB ≥13 mut/Mb and PD-L1 level of ≥1% (HR 0.95; 95% CI: 0.61–1.48; P=0.78) (7).

**Retrospective data from daily practice**

**Anti-PD(L)1 therapy**

In an analysis of 64 NSCLC patients treated with nivolumab, pembrolizumab or atezolizumab, a high TMB (FoundationOne® assay (not mentioned whether F1 LDT or F1 CDx™), TMB high: ≥15 mut/Mb) was associated with a longer duration of therapy: median was 64 versus 17 weeks (HR 0.396; 95% CI: 0.190–0.825; P=0.010). Moreover MSI-high status correlated strongly with TMB (24).

**Nivolumab**

A numerically higher median OS for high TMB NSCLC patients treated with nivolumab was found in a small study (N=36): median OS (95% CI) was not reached for TMB high (≥20 mut/Mb), 10.3 (4.8–15.7) for intermediate TMB (6–19 mut/Mb) and 12.4 (7.1–17.1) months (P=0.211). Interestingly, a clinical score (iSEND) based on gender, performance status, neutrophil-to-lymphocyte ratio (NLR) and deltaNLR correlated significantly with OS. Median OS (95% CI) was 15.7 (10.8–20.6), 10.3 (4.8–15.7) and 3.7 (0–7.8) months for iSEND good, intermediate and poor groups, respectively (P<0.001). TMB and iSEND were not combined to one predictive score (59).

In a series of 444 NSCLC patients treated with nivolumab, high TMB (determined through the FoundationOne® (not mentioned whether F1 LDT or F1 CDx™) assay, TMB high defined as ≥20 mut/Mb), patients with a high TMB had a longer duration of nivolumab therapy (7.5 versus 4.6 months, P<0.001) as well as a longer median OS (not reached versus 10 months, P<0.01). In this series, PD-L1 status (available for 282 patients) was not associated with outcome (23).

In another large series (1,619 samples from the Flatiron Health Analytic Database) of NSCLC patients treated with IC with available TMB [FoundationOne® assay (not mentioned whether F1 LDT or F1 CDx™)], OS was superior for nivolumab compared to chemotherapy in the TMB ≥13 mut/Mb and PD-L1 level of ≥1% (HR 0.95; 95% CI: 0.61–1.48; P=0.78) (7).
with duration of therapy but high TBM predicted for longer duration of therapy (P=0.001). Interestingly, STK11 mutations correlated with a high TMB/low-PD-L1 status and in a preliminary analysis an inferior outcome on ICI. BRAF mutations were correlated with a longer duration of therapy irrespective of TMB (22).

The use of TMB determined by a large NGS panel (MSK-IMPACT assay) has also been evaluated in 240 patients treated with anti-PD-(L)1 or anti-PD-(L)1 plus anti-CTLA4 combination therapy. 186 (78%) of the treated patients were treated outside of a clinical trial. Totally, 29% of the patients had DCB and TMB was significantly higher in those with DCB compared to those without (median 8.5 versus 6.6 SNVs/Mb, P=0.0062). DCB was 38.6% vs. 25.1% in those with a TMB above versus below the 50th percentile (P<0.001). PFS HR was 1.38 (P=0.024). TMB was also evaluated in a non-ICI, chemotherapy treated cohort of NSCLC patients: an inverse relationship between TMB and survival was identified (14). PD-L1 expression level was also significantly correlated with PFS (HR 0.526; P=0.011 for PD-L1 ≥1% versus 0%) but not with TMB. Patients with high TMB and PD-L1 positivity (≥1%) had the highest rate of DCB (50%) (14).

**Drawbacks of TMB**

Determining TMB becomes increasingly feasible with the availability of NGS technology. However, the determination of TMB on tissue raises several limitations in term of sample consumption and timing of the results. For example, for the commercially available F1 CDx™ test, a tumor block or 10 unstained tumor slides cut at 4–5 microns thickness are preferred with an optimum of 25 mm² surface area and at least 20% (preferred 30%) of tumor nuclei. In 2018, turnaround time is approximately two weeks, and cost is between 2,500 and 5,000 US dollars although it is expected that price will decrease with time (18,19,60).

Secondly, the attrition rate due to sample quality and quantity could be higher than current molecular testing. The percentage of TMB evaluable patients enrolled in clinical trials is lower than that for PD-L1. For example, in the CheckMate 568 trial, 88% of patients were evaluable for PD-L1 but only 34% were evaluable for TMB (F1 CDx™) (9).

Furthermore, TMB testing is not standardized for the different assays used and the definition of high TMB varies across studies from ≥7.4 to 20 mut/Mb when measured with a large NGS panel (7,9,14,21-24,58,59). As already mentioned, harmonization efforts are ongoing (26,27), but our opinion is that more prospective clinical trials will be needed to correctly determine the threshold and validate the clinical utilization of the TMB result.

Besides quantity and quality of the tissue, time, costs and defining the optimal cutoff there are drawbacks of measuring TMB as not only total TMB influences outcome on ICI. For example, when measuring TMB from a single sample, both clonal and subclonal neoantigens are counted in the TMB. Clonal neoantigens are present in all tumor cells while subclonal neoantigens are only present in a subset of tumor cells. In a series of 31 tumors from NSCLC patients treated with pembrolizumab it was shown that especially those with a high TMB as well as a low neoantigen subclonal fraction (defined as <5% subclonal) had the most durable clinical benefit with pembrolizumab. The authors also demonstrated that T-cells only recognized clonal neoantigens and not subclonal neoantigens. This is important as chemotherapy seems to enhance TMB mainly through the increase of subclonal neoantigens. Even when T-cells are generated against these subclonal antigens, these T-cells will not be able to control all tumor cells (61). Similar results were found in an exploratory analysis of the CheckMate 012 trial: clonal predicted neoantigen burden was more predictive of improved PFS compared with total predicted neoantigen burden (57).

Not only the load of neoantigens but also the neoantigen fitness should also be taken into account (62), This can be modelled by not only using information on the frequency of the clone but also using amplitude of MHC presentation (using relative MHC affinity of wild-type and mutant peptide) and T-cell recognition probability (62).

**Other factors influencing immunotherapy outcome**

Besides PD-L1 status and TMB, outcome on ICI can be influenced by several factors. An exhaustive coverage of biomarkers and other factors that influence outcome on ICI is outside of the scope of this review. Some examples are summarized here. It has been demonstrated that some cancer types have a better outcome with ICI treatment than would be expected from the TMB and for other tumors outcome is worse (17). Other factors such as viral antigen presentation on tumors (e.g., Merkel-cell carcinoma) may also result in a favourable ORR despite a lower TMB (63). Several oncogenic pathways can also have an impact on evasion of antitumor response, as was reviewed in Spranger...
et al. (64). In summary, defective critical steps in antitumour T-cell priming might result in a poor outcome despite high TMB. For example, functional dendritic cells are necessary for cross-presenting tumour neoantigens through the MHC class I pathway. Moreover, there are a certain number of gain-of-function alterations such as WNT-beta-catenin signalling or gain of MYC function that result in reduced T-cell recruitment and reduced T-cell activation and infiltration, respectively. Loss-of-function alterations such as LKB1 (STK11) mutations, PTEN loss, and TP53 loss can result in decreased T-cell infiltration, reduced T-cell priming and decreased T-cell infiltration, respectively.

Other aberrations in oncogenic pathways, such as mutations in isocitrate dehydrogenase 1 and 2 (IDH1 and IDH2) are associated with reduced T-cell infiltration while STAT3 signalling can decrease expression of pro-inflammatory cytokines (57,64). It has been shown that TILs represent both a predictive and a prognostic biomarker (65,66).

Furthermore, gene expression profiling performed in longitudinal tumor biopsies showed dynamic changes in multiple genes after the initiation of PD-1 therapy. These immune signatures could also be a complementary solution to guide treatment decisions. Targeted gene expression panels quantify the specific RNA expression profiles from formalin-fixed paraffin embedded (FFPE) biopsy. The quantification of the RNA expression will comprehensively delineate an inflamed tumor microenvironment. Immune expression profiling has the potential to accurately determine the inflammatory status of a tumor better than the PD-L1 status alone. The correlations between RNAseq and those signatures have been explored and seem very acceptable (67). Several approaches are proposed with RNA extracted from FFPE: a 25-gene interferon-gamma (IFN-γ) signature derived from crude extract (EdgeSeq, HTG Molecular Diagnostics Tucson, AZ, USA) and 700 genes from isolated RNA (NanoString Technologies, Seattle, WA, USA). For lung cancer, several trials have used this option: for example, the POPLAR trial with atezolumb and the T-effector/IFN-γ signature. An OS HR of 0.43 (95% CI: 0.24–0.77) was found for atezolizumab versus docetaxel in the high gene expression level group compared to an OS HR of 1.1 (95% CI: 0.68–1.76) for atezolizumab versus docetaxel in the low gene expression group (68).

It is also important to take into account exogenous factors such as use of antibiotics and steroids. It has been shown in a retrospective study (N=360, of which 239 NSCLC) that use of antibiotics within 30 days of ICI start is associated with a worse outcome compared to patients that did not use antibiotics. For NSCLC patients, median PFS was 1.9 versus 3.8 months, respectively (HR 1.5; 95% CI: 1.0–2.2; P=0.03). Median OS was 7.9 versus 24.6 months, respectively (HR 4.4; 95% CI: 2.6–7.7; P<0.01). Use of antibiotics retained its significance for OS in multivariate analysis (69). An imbalance in gut microbiota has also been associated with poor outcomes on ICI, and it is suggested that this might be caused by use of antibiotics (70).

Baseline steroid use (prednisolon equivalent of ≥10 mg/day) was also associated with worse outcome on ICI in a retrospective series of 640 NSCLC patients and steroid use retained its significance in multivariate analysis [HR for PFS 1.31 (P=0.03), HR for OS 1.66 (P<0.001)] (71).

**Future: TMB and other biomarkers to guide treatment decisions**

There are besides PD-L1 and TMB multiple parameters associated with ICI outcome (e.g., prognostic scores such as LIPI and dNLR, TILs, T-effector/IFN-γ signature, general immune fitness, soluble inhibitors, tumor metabolism, microbiome), and as such it would be interesting to combine several parameters (16,52,66,68,72).

Challenge when combining multiple biomarkers is the increasing number of subgroups that is created. For example, with two biomarkers there are already four subgroups and this doubles to eight when there are three biomarkers combined. It then becomes increasingly difficult to define cutoffs and interdependence. In reports of clinical trials, only PD-L1 and TMB were combined.

However, obtaining enough tissue to perform all analysis necessary to make a treatment decision (define histological subtype, test for molecular drivers, PD-L1 and TMB) can be difficult. Furthermore, most predictive biomarkers are expensive to perform in daily routine. Exceptions are the predictive scores using laboratory values already determined in routine clinical care as the LIPI score which uses only pretreatment neutrophils, lymphocytes and LDH (72). Now, the performance of all those scores should be evaluated and compared with PD-L1 or TMB testing.

**Conclusions**

Anti-PD(L)1 therapy has become standard of care for NSCLC, and some of the combination strategies with chemotherapy (or anti-CTLA4 therapy) are likely to become standard of care in the near future for certain subgroups of patients. With all of these possible treatment options,
Biomarkers are needed to identify which subgroup of patients is likely to benefit the most from a certain therapy. Although not perfect, a high PD-L1 expression level can aid in selecting the right patient population for monotherapy pembrolizumab. TMB is an interesting biomarker, however assays and definition for high TMB should be standardized and it is challenging to obtain an adequate amount of tissue. Importantly, high TMB predicted for improved PFS with nivolumab plus ipilimumab but OS data of the CheckMate 227 are still immature and data for nivolumab monotherapy versus chemotherapy are conflicting. When assays are standardized and an OS benefit is shown, TMB in combination with PD-L1 expression level could help in defining subgroups of patients that will likely benefit from monotherapy or combination strategies.

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Footnote


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