PD-L1 immunostaining scoring for non-small cell lung cancer based on immunosurveillance parameters

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Abstract

Non-Small Cell Lung Cancer (NSCLC) is the leading cause of cancer death globally, and new immunotherapies developed and under development targeting PD-1/PD-L1 checkpoint inhibition require accurate patient selection to assure good clinical outcome. PD-L1 immunohistochemistry is the current biomarker assay used for patient selection, but still imprecise in predicting therapy response. Exploring this issue, we performed computational tissue analysis of PD-L1 immunostaining in procured NSCLC tissues (n = 50) using the Merck KGaA anti-PD-L1 clone MKP1A07310. Staining patterns and PD-L1 cut-off points were interrogated using relevant cancer immune-surveillance biomarkers. Groups with high PD-L1 expression levels (above 25/50% staining cut-off points) were enriched for a biomarker profile in the tumor-nest and microenvironment indicating escape from host-immunity, as represented by increased numbers of cells positive for CD8 and Granzyme B (immune-effectors), FOXP3 (immune-suppressive), and CD68 (P < 0.05). Manual analysis of PD-L1 staining patterns identified tumors with an immune-induced reactive pattern relevant for immunotherapy that would ordinarily be excluded by the arbitrary 25% staining threshold (P < 0.05). Conversely, some cases with completely or predominantly immune-independent constitutive PD-L1 staining patterns that indicate insensitivity to immunotherapy may have been incorrectly selected using this staining cut-off point criterion. Therefore, we propose differentiation of reactive vs constitutive PD-L1 staining patterns to improve the accuracy of this biomarker assay in selecting NSCLC patients for PD-1/PD-L1 immunotherapy.

Introduction

Lung cancer is the most lethal cancer type worldwide, with non-small cell lung cancer (NSCLC) accounting for the majority (~85%) of deaths [1]. Late diagnoses of NSCLC contribute to the high mortality and poor prognoses for those with advanced disease, treated with platinum-based chemotherapy [2]. Novel immunotherapies that target the Programed Cell
Death (PD-1)/ligand (PD-L1) check-point have been approved for NSCLC based on improved overall response rates in clinical trials using therapeutic monoclonal antibodies (mAb) directed against PD-1 (Nivolumab and Pembrolizumab) and PD-L1 (Atezolizumab), while other anti-PD-L1 mAbs (Avelumab and Durvalumab) are at advanced stages of development [3–5]. With the current range of options, it is important to understand which patients will benefit from each new immunotherapy drug. PD-L1 immunohistochemistry (IHC) assays are the most advanced biomarker strategies in-place for patient screening, as elevated tissue expression of PD-L1 has been correlated with a higher chance of prolonged post-therapy survival [3, 4, 6]. However, this approach has attracted some skepticism given that significant number of patients, even in highly selected cohorts, have treatment outcomes that contradict predictions based on PD-L1 status [6]. The use of multiple antibodies and staining thresholds (1–50%) complicates PD-L1 IHC data comparisons [4, 6, 7]. Furthermore, we lack a standard scoring algorithm for PD-L1 expression patterns in tumor-nests and the tumor microenvironment (TME) [3, 8]. Finally, the biological relevance of PD-L1 staining cut-off points, including their relationship to other immune biomarkers, remains unknown.

The PD-L1 staining profile targeted by PD-L1/PD-1 immunotherapies should be a reactive pattern induced by interferon (IFN)-γ secreted by tumor infiltrating lymphocytes (TIL) at tumor-nest-TME interfaces [6, 8–10]. This reactive expression, from immune and tumor cells responsive to IFN-γ, differs from constitutive expression arising from cell-intrinsic genetic/epigenetic events. A strong and homogeneous constitutive PD-L1 tumor cell staining pattern in the absence of any appreciable TIL component would predict a futile PD-L1/PD-1 therapeutic intervention [6, 8]. Conversely, a reactive patchy PD-L1 pattern surrounded by T-cell rich areas implies restraint of a potentially anti-tumoral cytotoxic adaptive response, and a greater likelihood of anti-PD-L1/PD-1 therapeutic success [11, 12]. PD-L1 negative tumors without TIL will not respond to PD-1/L1 therapy unless cancer antigens are exposed for a T-cell response, while genetic events leading to inability to express PD-L1 in the presence of TIL should identify acquired resistance to PD-1/L1 therapy. Currently, none of the applied PD-L1 scoring algorithms account for patterns that indicate cancer escape from immune surveillance [6, 9].

This is the first publication to report IHC data for tumor-nest and TME compartments in NSCLC samples using the Merck KGaA anti-PD-L1 clone MKP1A07310 for Avelumab development together with an immune panel of effector and regulatory TIL [13], as well as to show the effects of PD-L1 staining cut-off points, and the stratification of biologically relevant PD-L1 staining patterns in tumor-nest and TME compartments. Using Computational Tissue Analysis (cTA™) capable of scoring entire sections, complementary to another NSCLC digital study [13], we expanded PD-L1 analyses to include all positively-stained TME cell types as all may support an immuno-suppressive environment. We demonstrated that simplistic thresholds based on PD-L1 staining amount over a whole section can exclude patients with lower amounts but potentially therapy-permissive patterns, while including cases with strong labeling that is not necessarily related to a restrained adaptive immune response.

**Materials and methods**

**Tissues and immunohistochemistry**

Formalin-fixed paraffin-embedded tissue blocks prepared using surgical resections from 50 treatment-naive NSCLC patients (Table 1) were procured from Indivumed GmbH (Hamburg, Germany) and processed in our Merck KGaA histopathology laboratory for IHC. Four-micrometer-thick serial sections were generated per tissue block and immunostained for CD8 (cytotoxic T-cells), FOXP3 (regulatory T-cells), CD56 (natural killer (NK) cells), Granzyme B
(functionally active cytotoxic NK/T-cells), or CD68 (pan macrophage marker), with corresponding matching isotype controls. Staining was performed in a BenchMark XT immunostainer (Roche Diagnostics, Germany). Antigen retrieval pretreatments, primary antibody (Ab) working conditions, as well as detection and hematoxylin counterstaining methods are listed in Table 2. Tris-EDTA buffer pH 8 for mild and standard CC1 (Cat. 950–124), UltraView Universal DAB (Cat. 760–500), the OptiView DAB IHC Detection kit (Cat. 760–700), and Hematoxylin II (Cat. 790–2208) were purchased from Ventana Medical Systems (Roche Diagnostics, Germany) and used according to the manufacturer’s instructions. Additional serial sections were immunostained under the conditions presented in Table 2 for PD-L1 (clone MKP1A07310), with respective matching isotype controls. Staining was performed in a Dako AutostainerLink 48 instrument (Hamburg, Germany). EnVision FLEX/HRP DAB detection kit (Cat. SM802), TRS high pH (Cat. K8004), and Hematoxylin EnVision FLEX (Cat. SM806) were purchased from Dako (Hamburg, Germany). All primary Ab and isotype controls were diluted in PBS pH 7.

Whole-slide scanning and manual annotation

Slides were digitized using Aperio’s AT Turbo and CS bright-field slide scanning systems (Leica Biosystems Inc., Buffalo Grove, IL) at Flagship Biosciences’ tissue analysis laboratory. A minimum of 4 focal points were set for each slide to allow for optimal scanning quality, with

Table 1. Patient’s clinical information.

| Gender | Male (n = 26) | Female (n = 24) |
|--------|--------------|----------------|
| Age (Average ± SD; years) | 67 ± 8 | 66 ± 10 |
| Disease Stage | IIA (n = 37) | IIB (n = 1) | IV (n = 12) |
| NSCLC histological subtype | Adenocarcinoma (n = 24) | Squamous (n = 16) | Large Cell (n = 5) | Pleomorphic (n = 5) |

Table 2.

| Primary Ab (Clone) | Manufacture | Catalogue | Isotype | Concentration (μg/ml) | Incubation | Isotype control | Antigen retrieval | Detection | Counterstain |
|--------------------|-------------|-----------|---------|-----------------------|------------|----------------|------------------|-----------|-------------|
| CD56 (MRQ-42)      | Ventana     | 760–4596  | Rabbit IgG | 0.5–1                 | 32 min, 37°C | Cell Signaling, Cat. 3900 | Mild CC1 | UltraView DAB | Hematoxylin II |
| CD68 (PG-M1)       | Dako        | M0876     | Mouse IgG3, k | 0.06                | 32 min, 37°C | Biologic, Cat. 401302 | Standard CC1 | OptiView DAB | Hematoxylin II |
| CD8 (Gr/144B)      | Dako        | M7103     | Mouse IgG1, k | 8                   | 32 min, 37°C | Dako, Cat. X093 | Standard CC1 | UltraView DAB | Hematoxylin II |
| FoxP3 (236A/E7)    | Abcam       | ab20034   | Mouse IgG1 | 10                   | 32 min, 37°C | Dako, Cat. X093 | Standard CC1 | OptiView DAB | Hematoxylin II |
| Granzyme B (GrB-7) | Dako        | M7235     | Mouse IgG2a, k | 5.5                | 60 min, RT | Biologic, Cat. 400224 | Standard CC1 | UltraView DAB | Hematoxylin II |
| PD-L1 (MKP1A07310) | Merck KGaA | Proprietary | Rabbit IgG | 0.25                | 30 min, RT | Cell Signaling, Cat. 3900 | TRS high pH | EnVision FLEX/HRP | Hematoxylin EnVision FLEX |

Mild CC1: 28 min at 96 °C
Standard CC1: 56 min at 100 °C
TRS high pH: 20 min at 95 °C
Hematoxylin II: 8 min
Hematoxylin EnVision FLEX: 5 min

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scanning performed at x20. All whole-slide digital images were assessed for scanning artifacts including blurring due to incomplete cleaning, and 'stitching' artifacts. Scanning was repeated until all slides reached Flagship’s quality criteria for digital scans. Digital images were viewed using Aperio ImageScope (Leica Biosystems Inc., Buffalo Grove, IL), with manual annotation to include region(s) of interest (ROI). Inclusion annotations were made just outside the invasive margin of the tumor mass when present (Fig 1A and 1B), excluding adjacent pre-existing normal lung and/or pleural tissue and in-situ carcinomatous changes. Exclusion annotations were made around staining and tissue artifacts including folds and tears, mounting media artifacts, stain smudges, and other artifacts deemed irreversible in the scanning process. Additional exclusions were areas of necrosis (Fig 1B), entrapped airway tissue (including hyperplastic or metaplastic epithelium), ectopic lymphoid structures, and large intraluminal clusters of macrophages and/or cellular debris. One tissue sample was dropped from the study due to widespread anthracosis.

**Computational tissue analysis (cTA)**

Full scan analysis was performed using Flagship Bioscience’s proprietary cTA image analysis software system (Westminster, CO, USA) available for research. The platform identified nuclei in each sample based upon hematoxylin staining, then quantified the intensity of DAB staining...
for each nucleus-identified cell. To identify positive cells, thresholds for DAB staining were set using biomarker-specific algorithms for CD8, FOXP3, CD56, Granzyme B, CD68, and PD-L1. For PD-L1, multiple thresholds were identified according to a negative, +1, +2, and +3 scoring paradigm contiguous with manual scoring approaches (Fig 1C and 1D). The biomarker-specific algorithms were then further developed on a sample-by-sample basis to assign cells to either the tumor-nest or TME compartments based on morphometric and/or staining qualities (Fig 1D). Instances where compartmental separation was impossible due to similar morphologies of tumor nest and TME cell nuclei, or an insufficient quantity of TME, cell information was studied from both compartments. Data from these cases were included in the “All Cells” category and excluded from the solo tumor-nest and TME compartment-specific analyses. All annotations and image analysis markups were assessed by a pathologist (JPK) to verify performance and accuracy. Stained cell counts and the percentage of positive cells for each tissue sample in every biomarker on a per-compartment basis was performed. Heterogeneity was further analyzed for PD-L1 stained samples using an algorithm-calculated H-Score within the tumor-nest compartment, the TME compartment, and the entire tissue sample respectively. Digital H-scores ranging from 0 to 300 were calculated using the following standard formula: [3 × % cells +3 intensity] + [2 × % cells +2 intensity] + [1 × % cells +1 intensity].

Pathologist scoring of PD-L1 staining patterns

In addition to the image analysis detailed above, all slide images were reviewed and categorized according to the PD-L1 staining patterns (I-V) noted in tumor-nest and TME compartments, including the tumor-nest-TME interfaces (Table 3). The system used was an adaptation/extension of those previously suggested for cancer classification based on PD-L1 expression by tumor cells and co-localization with TIL [6, 8]. The aim of this categorization was an attempt to differentiate between constitutive versus reactive PD-L1 expression (or mixtures of the two) in order to predict potential responders to PD-1/PD-L1-based immunotherapy [6, 8]. In this study, we assessed PD-L1 expression by all cell types in the TME compartment rather than using TIL counts.

Statistical analysis

Hierarchical clustering of the data for all biomarkers was performed using the MATLAB bioinformatics toolbox version 2015a (MathWorks, Natick, MA) with Euclidean distance metric.

Table 3. Classification of NSCLC according to PD-L1 expression patterns in tumor-nest and TME compartments.

| Pattern | PD-L1 expression† | Biological relevance |
|---------|-------------------|----------------------|
| I       | TUM+ (diffuse)    | Mixed of genetic/epigenetic (constitutive) and adaptive immunity effect (induced) |
|         | TME+              |                      |
| II      | TUM- (patchy)     | Only driven by adaptive immunity (induced) |
|         | TME+              |                      |
| III     | TUM- TME-         | Immune ignorance     |
| IV      | TUM+ TME-         | Constitutive only    |
| V       | TUM- TME+         | Tumor cells might be genetically unable to express PD-L1 and/or functional interferon-γ receptors |

* Based on system proposed by Teng et al. [8], but with subdivision of category I by Ribas and Hu-Lieskov et al. [6], and assessment of TME rather than only tumor-infiltrating lymphocytes.

†TUM (tumor-nest compartment) and TME (tumor microenvironment compartment). (+) and (-): positive and negative PD-L1 IHC, respectively.

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and average linkage to generate the hierarchical tree. Data were normalized such that the mean was 0 and the standard deviation was 1 across samples for each marker score. Pearson correlation coefficients were also calculated between marker measurements and a heatmap was generated to visualize the strength of the correlations.

Percentages of cells positive for each biomarker were compared for tumors above and below 5%, 25%, 50%, and 80% cut-off points for PD-L1 immunostaining. These comparisons were made for the tumor-nest, TME, and “All Cells” compartments. Additionally, positive PD-L1 percentages, immune cell percentages, and digital H-scores were compared between the PD-L1 staining pattern categories in the tumor-nest and TME compartments, respectively. Data were plotted as box-and-whisker plots. Boxes show the median and interquartile range (IQR) and whiskers show the range of the data excluding outliers. Outliers are defined as data more than 1.5*IQR beneath the lower quartile or more than 1.5*IQR above the upper quartile. No data points were removed from analysis. P-values were calculated using a two-tailed Student’s t-test, assuming unequal sample variances. The Holm method was used to correct for multiple comparisons, with a significance threshold of $P<0.05$ for each subset of analyses.

Results

Correlations between immune biomarkers in tumor-nests and TME

Multiple biomarkers were analyzed in the tumor-nest, TME, and All Cells compartments. This allowed us to accurately assess the relative proportions of effector and regulatory cells in the typically heterogeneous NSCLC tissue microenvironment, and their possible relationships. In a small number of cases ($n = 7$), it was not possible to separate tumor-nest and TME compartments digitally as the latter was minimal; All Cells scores only were analyzed for those slides. In the group of cases where a separation between tumor-nest and microenvironment was feasible ($n = 43$), scores derived from a single marker showed a greater correlation across compartments than scores derived from different markers in the same compartment (Fig 2A). For example, percentages of CD68-positive cells in the tumor-nest and TME compartments had a Pearson correlation coefficient ($\rho$) of 0.73, which fell to 0.42 when comparing CD68 and CD8 in the TME alone. However, PD-L1 data showed a lower correlation between the tumor-nest and TME compartments ($\rho = 0.43$) than the other markers (Fig 2B). CD56 showed the least correlation with other markers, while CD8 and Granzyme B (both expressed by cytotoxic T-cells) showed the greatest association with each other (All Cells $\rho = 0.93$, $P < 0.0001$; Fig 2B). Of the biomarkers beside PD-L1, CD8- and CD68-positive cells had the highest mean percentages in both tumor-nest (10.7%; 8.3%) and TME compartments (11.1%; 13.2%) (Table 4). Finally, we could not identify significant clustering based on the four NSCLC subtypes represented in our sample set (adenocarcinoma, squamous cell carcinoma, large cell carcinoma, and pleomorphic carcinoma).

Distribution of immune biomarkers relative to frequently used PD-L1 cut-off points

We compared the distributions of our immune markers relative to PD-L1 positive staining cut-offs (5, 25, 50, and 80%) currently used in Merck KGaA PD-L1 clinical development programs. We studied cut-off points in the tumor nest, TME, and All Cells compartments, with a full analysis of the immune markers in each compartment. A 50% PD-L1 TME cut-off put only 14% of samples in the high PD-L1 group, increasing to 100% at a 5% PD-L1 TME cut-off. Whereas, data from tumor-nest cut-offs permitted a more balanced sample size distribution. For example, using the 25% and 50% PD-L1 tumor-nest cut-offs, approximately half of the
Fig 2. Relationships between biomarkers in our NSCLC cohort. (A) Hierarchical clustering heat map of percentages of positive cells for FOXP3, PD-L1, CD8, Granzyme B, and CD68. Each biomarker (row) was normalized before clustering to give a mean of 0 and a standard deviation of 1 across columns (samples). The color scale indicates the relative size of the biomarker score compared to the other samples in the cohort with blue = low, white = neutral, and brown = high. Bracket height indicates the degree of correlation between measurements with shorter brackets representing higher correlations and taller brackets representing smaller correlations. (B) Heatmap of Pearson correlation coefficients between percent positive cells measurements across samples for FOXP3, PD-L1, CD8, Granzyme B, and CD68. The color scale ranges from the minimum (blue) and maximum (red) correlation coefficient observed in this analysis. Therefore red represents strong positive correlations, white represents moderate correlations, and blue represents weak correlations. No strong negative correlations were observed.

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samples were in the high PD-L1 group (53% and 44%, respectively). A plot of 50% PD-L1 tumor-nest cut-off (most frequently scored by pathologists) for overall immune cell expression (All Cells) provides a summary of the global changes in cellular distribution for the PD-L1 high/low groups (Fig 3A). Higher percentages ($P < 0.05$) of stained cells for CD68, CD8, FOXP3, and Granzyme B were detected in the high PD-L1 group when compared vs the low PD-L1 group (Fig 3A). Results were similar when comparing immune marker distribution in the tumor-nest, TME, and All Cells compartments using 25% and 50% PD-L1 tumor-nest cut-off points (non-shown data). Additionally, we found no significant differences in Granzyme B/CD8 ratio for the 50% cut-off ($P = 0.14$), but a higher FOXP3/CD8 ratio beneath the 50% threshold ($P = 0.03$). When applying a 5% PD-L1 tumor-nest cut-off, 70% of samples segregated to the high PD-L1 group, with CD8, Granzyme B, and CD68 significantly higher in those samples ($P < 0.05$). Statistical comparisons using an 80% PD-L1 tumor-nest cut-off value were impossible given that only one sample segregated to the high PD-L1 group.

Classification of PD-L1 staining patterns and relation with cut-Off points

The staining patterns for PD-L1 (Table 3) were classified for cases with tumor-nest and TME compartment data ($n = 43$) to determine whether any NSCLC case with evidence of reactive membrane expression would be falsely excluded at certain cut-off points, while some NSCLC with only constitutive tumor expression might be inappropriately included. Most of the tumors in this sample set were classified as having pattern I ($n = 16$) or II ($n = 16$); both showing PD-L1 expression in tumor cells at the tumor-nest-TME interface that suggested contribution of a reactive process (Fig 4A and 4B).

In NSCLC classified as pattern I, the tumor cells showed variable percentages of PD-L1 membrane labeling (1+, 2+, and 3+ scores) and positive staining in TME cells (Fig 4A). Heavier PD-L1 staining of tumor cells adjacent to positive TME cells were observed by the pathologist in some fields, which varied within and between sections. In tumors classified as pattern II, the variable staining intensity was largely limited to patches that were often located at the tumor-nest-TME interface (Fig 4B). PD-L1 positive cells in the TME for pattern I and II included morphological cellular components compatible with lymphocytes, macrophages, fibroblasts, and endothelial cells. Looking at the other immune biomarkers, CD8 and to lesser extent CD68, most often showed pronounced infiltration on both sides of the tumor-nest-TME interface, as opposed to the other immune markers that tended to be more diffusely arranged in both compartments. Only one tumor showed predominantly negative staining for PD-L1 in both compartments (tumor-nest and TME), consistent with pattern III (Fig 4C). Pattern IV (diffuse and intense staining of tumor cell membranes with minimal to no TME expression) was also only noted in one case (Fig 4D). Finally, there were a number of NSCLC

| Biomarker | All Cells* | Tumor-nests (n = 43) | Tumor microenvironment (TME) (n = 43) |
|-----------|------------|---------------------|--------------------------------------|
| PD-L1     | 32.90      | 35.60               | 31.40                                |
| CD8       | 8.36       | 6.02                | 11.11                                |
| Granzyme B| 3.17       | 2.05                | 4.41                                 |
| FOXP3     | 4.42       | 1.81                | 7.16                                 |
| CD68      | 8.61       | 4.14                | 13.43                                |
| CD56      | 0.63       | 0.22                | 1.07                                 |

*“All Cells” is the combined data for tumor-nest and TME

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Fig 3. Immune markers in NSCLC relative to a 50% cut-off point for PD-L1 staining. A: Percentage of positive cells for each biomarker in "All Cells" (tumor nest and TME compartments combined) calculated by using Computational Tissue Analysis (cTA™). NSCLC with >50% PD-L1-positive tumor cells were compared with those with lower expression. Statistical differences were calculated using two-tailed Student’s t-tests, assuming unequal sample variances (*P ≤ 0.05, **P ≤ 0.01, ***P ≤ 0.001 and ****P ≤ 0.0001). Percentages of cells positive for CD68, CD8, FoxP3, and Granzyme B were significantly higher in the >50% PD-L1 group. B: Representative IHC images of PD-L1-, CD8-, and CD68-positive cells (the most numerous biomarkers) in an NSCLC (solid carcinoma) case with >50% PD-L1 expression in the tumor nest compartment, and an NSCLC (adenocarcinoma with mixed subtypes) case with lower PD-L1 expression. Scale bars = 100 μm. NSCLC = non-small cell lung carcinoma; IHC = immunohistochemistry; TME = tumor microenvironment.

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cases (n = 9) with PD-L1 staining in the TME of variable intensity and minimal to no tumor cell staining consistent with pattern V (Fig 4E).

Comparison of tumor groups with patterns I and II revealed a lower percentage of PD-L1 positive cells in the tumor-nests from the group showing pattern II staining (P < 0.0001). Importantly, most of those pattern II samples fell beneath the cut-off points when applying 50% and 25% PD-L1 tumor-nest thresholds (Fig 4F). However, comparable percentages of PD-L1-positive TME cells were noted for each of these patterns (Fig 4G). A similar separation of these two groups could be achieved by using a tumor-nest digital H-score, with pattern I
and II groups having significantly different averages of 160 and 41, respectively (Fig 4H). In terms of other immune biomarker cellular percentages (and their digital H-scores), there were no significant differences between pattern I and II groups for TME or All Cells analyses. However, the tumor nest compartment for pattern I tumors showed significantly higher percent of FOXP3-positive cells ($P = 0.002$; data not shown).

**Discussion**

Using procured NSCLC tissues from naïve-treated patients with advanced disease, we adopted the cTA platform to identify and quantify multiple immune biomarkers in tumor-nest and TME compartments in whole sections. Pivotal to this analysis was the usage of the Merck KGaA anti-PD-L1 clone MKP1A07310, which is the analytical Ab for the evaluation of Avelumab [5], with same specificity and higher sensitivity than Dako anti-PD-L1 clone 22C3 as shown in a NSCLC PD-L1 IHC assay comparison study [14]. We demonstrated that all our immune biomarkers, except for PD-L1, had a strong correlation across tumor-nest and TME compartments, with CD8 and Granzyme B being the TIL markers with strongest association. PD-L1 tumor-nest cut-off points, but not TME-related ones, allowed a balanced sample size distribution that showed similar results at 25 and 50% in terms of TIL-related and non-related immune biomarkers, clustering higher levels of these immune biomarkers with higher levels of PD-L1 expression. This is consistent with the hypothesis that higher PD-L1 tumor-nest expression would be expected in the context of an adaptive immune response to tumor antigens. However, we propose that a simple quantification of PD-L1 immunostaining levels is not enough to select patients for PD-1/PD-L1 immunotherapy [6], because cut-off points do not allow discrimination between constitutive and reactive expression patterns. Our data show that most cases with a patchy PD-L1 reactive pattern II fell below both 25 and 50% cut-off points, indicating that they would be excluded from PD-1/PD-L1 immunotherapy if those cut-off criteria were used for patient selection. Also, the inclusion of cases with high PD-L1 tumor expression related to a constitutive expression either alone (pattern IV) or predominant (pattern I) may not necessarily favor therapeutic outcome. In conclusion, we propose that current scoring systems based simply on amount of staining should be complemented by algorithms that discriminate between constitutive and reactive PD-L1 staining patterns, being the last one the appropriate target for PD-1/PD-L1 checkpoint inhibition.

We have demonstrated enrichment of effector and regulatory immune cell types in groups with higher PD-L1 expression as defined by IHC thresholds of over 25 and 50%. This included CD8 and Granzyme B positive cells, with the latter being a marker that highlights functional effector T-cells [15]. The surprisingly low percentage of Granzyme B$^+$ cells versus total CD8$^+$ cells in our study might be explained by a large proportion of exhausted CD8 T cells [9, 16]. The Granzyme B$^+$/CD8$^+$ ratio did not differ significantly across the 25% and 50% thresholds, but might not be expected in our particular cohort of patients with advanced disease. Increased percentages of FOXP3$^+$ cells with higher PD-L1 expression also suggested a tumor cell modulation of the adaptive immune response given that PD-L1/PD-1 interactions promote T-cells differentiation into regulatory cells [17]. The balance between FOXP3$^+$ and CD8$^+$ T-cells is another clinically linked parameter, as low-FOXP3/ high-CD8$^+$ ratios have been linked with positive survival effects across multiple cancer types including NSCLC [18–19]. In the current study, the FOXP3$^+$/CD8$^+$ ratio was lower in NSCLC above the 50% PD-L1 cut-off point, but this result does not necessarily indicate a favorable TME for an adaptive anti-tumoral immune response due to the likelihood that most of those TIL are exhausted. Overall, these results support using PD-L1 immunostaining cut-off points to select patients with higher chances of responding to PD-1/PD-L1 checkpoint inhibition, as they are consistent with previous positive...
correlations of PD-L1 expression, numbers of TIL, and longer overall survival in NSCLC and other solid tumors [15, 20–21, 22–23]. However, our further analysis of PD-L1 staining patterns in tumor-nests and TME suggested the need to take into account this additional aspect.

The categorization of tumor PD-L1 expression patterns related to TIL distribution has been discussed previously, suggesting clear guidelines for checkpoint inhibition immunotherapy [6, 8]. This approach may simplify PD-L1 scoring paradigms, but detracts from complex immunologic read-outs reflecting the contribution from various combinations of immune biomarkers. In fact, tumor-associated macrophages, fibroblasts, dendritic cells, and even endothelial cells can modulate immune responses via PD-L1 expression [16, 24–30]. PD-L1 may also be expressed on T-cells themselves [31]. Cells with morphologies consistent with all these cell types were found to express variable PD-L1 labeling intensities in our tissue sections classified as pattern I, II and V. Moreover, these PD-L1 expressing cells have been previously shown to be therapeutically important, as positive responses to Atezolizumab were enriched in NSCLC when PD-L1 was expressed by infiltrating immune cells [31]. In fact, absence of PD-L1 expression in the immune infiltrate was categorized as a “non-functional immune response” [31].

PD-L1 expression in TME immune and non-immune cells could be an over-arching biomarker for the anti-tumor response, allowing selection of appropriate patients for checkpoint inhibitor immunotherapy. However, it is unknown which are the characteristics of the PD-L1 staining required in the TME (pattern I, II and V) for a successful therapeutic intervention. In our study, the TME expression of PD-L1 alone did not facilitate separation of patients using cut-off points, but it’s relationship to tumor cell staining in the context of our PD-L1 pattern classification system (I-V) suggested why simple cut-off points may exclude therapy-responsive patients, while including some who are unlikely to respond.

The analysis of PD-L1 staining patterns of tumor cells was performed to titrate the presence of morphological structures where PD-L1/PD-1 interactions should occur like an ‘immunologic synapse’. This is the classic TCR/antigen presentation interface where T-cell immune activation and inhibitory signals interplay [32]. In our tissue context, localization of TIL at the tumor-nest-TME interface and among other PD-L1-expressing cell types within the TME compartment (captured in our pattern I, II and V) provide ample opportunities for PD-L1/PD-1 interactions to attenuate or paralyze any adaptive immune response. This physical interaction was most clearly seen in our staining pattern II, which had a prominent and patchy tumor-nest-TME interface (Fig 4B). In contrast, where PD-L1 expression by the tumor cells is diffuse, often intense, and not only localized to the interface (patterns I or III), it is logical to assume that this expression reflects tumoral-intrinsic events related to genetic mutation or epigenetic disturbance [6]. Such tumors will tend to be given high scores that are out of proportion to the numbers of infiltrating immune cells [33], although many of such cases in our study did also contain interface components (pattern I). We hypothesized that NSCLC tissues with interface-only staining (pattern II) as well as the cases showing mixed interface and diffuse staining (pattern I) contain inducible PD-L1/PD-1 interactions potentially adequate for therapeutic blocking. Of note, current NSCLC PD-L1 scoring methods do not allow us to identify patterns I and II [3], as traditional cut-off point fail to recognize these essential interactions at the interface. Moreover, we have shown that most of the pattern II tumors fell below a 25% and 50% cut-off points, which are commonly used in current scoring methods [3]. Also, these methods will not assess if a sufficient extent and/or magnitude of adaptive immunity is present to select patients who could benefit from PD-1/PD-L1 therapeutic intervention. In fact, some pattern II tumors may contain a more extensive and targetable adaptive immune response than those included in the pattern I category. Setting a lower tumor cell cut-off point (e.g. 1% or 5%) risks including pattern II without sufficient immunological synapse interfaces, while still including potentially recalcitrant pattern I or III tumors. Then, current scoring
algorithms based on amount of staining should be complemented with qualitative algorithms that allow the discrimination between constitutive and relevant reactive PD-L1 staining patterns.

In conclusion, using our cTA analysis of PD-L1 immunostaining relative to immune bio-markers that might be involved in NSCLC immune surveillance during this late disease stage, we demonstrated that groups with high levels of PD-L1 defined by cut-off points of 25 and 50% enriched also for higher amounts of FOxP3, CD8, Granzyme B, and CD68 positive cells in tumor-nest and TME. This association between high levels of PD-L1 and increased effector and suppressor adaptive as well as augmented innate immune tumor infiltrates may reflects escape from a potential host anti-tumoral immune response. Furthermore, high amounts of PD-L1 in NSCLC defined by these or other cut-off points may not accurately predict if anti-PD-1/PD-L1 therapy will successfully unleash a host anti-tumoral immune response. Selection of a specifically reactive PD-L1 staining pattern within regions of the tumor architecture where immunological synapses are formed is likely to be required for more accurate determination of which patients will benefit from this therapeutic intervention. We anticipate that the current data may trigger a new way to analyze clinical trial specimens, because discrimination between constitutive and reactive PD-L1 staining patterns should improve the precision of immunotherapy patient selection.

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