Relationships between apparent diffusion coefficient (ADC) histogram analysis parameters and PD-L1 expression in head and neck squamous cell carcinomas: a preliminary study

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Background. Immunotherapy has become a cornerstone of the modern cancer treatment. It might be crucial to predict its expression non-invasively by imaging. The present study used diffusion-weighted imaging (DWI) quantified by whole lesion apparent diffusion coefficient (ADC) values to elucidate possible associations with programmed cell death ligand 1 (PD-L1) expression in head and neck squamous cell cancer (HNSCC).

Patients and methods. Overall, 29 patients with primary HNSCC of different localizations were involved in the study. DWI was obtained by using a sequence with b-values of 0 and 800 s/mm² on a 3 T MRI. ADC values were evaluated with a whole lesion measurement and a histogram approach. PD-L1 expression was estimated on bioptic samples before any form of treatment using 3 scores, tumor positive score (TPS), immune cell score (ICS), and combined positive score (CPS).

Results. An inverse correlation between skewness derived from ADC values and ICS was identified (r = -0.38, p = 0.04). ADCmax tended to correlate with ICS (r = -0.35, p = 0.06). Other ADC parameters did not show any association with the calculated scores.

Conclusions. There is a weak association between skewness derived from ADC values and PD-L1 expression in HNSCC, which might not be strong enough to predict PD-L1 expression in clinical routine. Presumably, ADC values are more influenced by complex histopathology compartments, comprising cellular and extracellular aspects of tumors than only of a single subset of tumor associated cells.

Key words: head and neck squamous cell cancer; apparent diffusion coefficient; diffusion weighted imaging; programmed cell death ligand 1

Introduction

Head and neck squamous cell carcinoma (HNSCC) is one of the most frequent malignancies.¹ Modern functional imaging modalities, comprising diffusion-weighted imaging (DWI) and Dynamic-contrast enhanced MRI provide further insight into tumor microstructure.² With these approaches, several diagnostic and prediction aspects were made possible, including the prediction of several histological features in tumors, treatment success and new prognostic.²,₅
DWI assesses random-water motion in tissues, namely Brownian molecular movement and can be quantified by apparent diffusion coefficient (ADC).\textsuperscript{3,5} This water movement is hindered predominantly by cellularity and thus it is a well-known fact that ADC values are inversely correlated with cell count with yet different results across several tumor entities.\textsuperscript{5} However, ADC values are not only associated with cellularity, but also with other histopathologic features, such as proliferation potential, neoangiogenesis, tumor invading lymphocytes or extracellular matrix. This was also shown for HNSCC.\textsuperscript{6-9}

Recently, various studies elucidated further associations of ADC values and tumor characteristics in oncology. ADC values aid in diagnostic purposes, as benign tumors have significantly higher ADC values than malignant tumors. This was also reported for head and neck lesions.\textsuperscript{10-12} Another important aspect is treatment prediction. A higher pretreatment ADC value and, therefore, presumably a tumor with a lower cell density and a less malignant tumor biology, is an independent favorable prognostic factor to chemotherapy and radiochemotherapy.\textsuperscript{13-15} Moreover, ADC values are independently associated with overall prognosis.\textsuperscript{13-15}

There is a growing interest in targeting immune modulatory checkpoints, especially the programmed cell death protein 1 (PD-1) around oncology.\textsuperscript{16} It is a transmembrane protein that is frequently expressed on the surface of T-lymphocytes. Programmed cell death ligand 1 (PD-L1) is a transmembrane protein, which is physiologically expressed at low levels in several cells, such as vascular, endothelial and immune cells. Interestingly, it can be highly expressed in neoplastic cells.\textsuperscript{16} The binding of PD-L1 to PD-1 forms an immunological situation that can impair the proliferation and the function of the immune cells against the tumor. That is why high expression of PD-L1 enables cancer cells to escape from immunologic response.\textsuperscript{16}

Immunotherapy was stated as a new paradigm shift in HNSCC treatment.\textsuperscript{17} Moreover PD-L1 expression on immunohistochemical specimen was shown to predict progression free survival in in advanced HNSCC stages and can predict metastases.\textsuperscript{18-20}

However, it is unclear, whether ADC values derived from MRI are associated with the amount of PD-L1 stained tumor cells and the associated PD-L1 stained immune cells and not only with the whole tumor cellularity of the lesion, which might not be as clinically relevant as to predict several subtypes of cells.

Therefore, the present study sought to elucidate possible associations between ADC values derived from MRI and the amount of PD-L1 stained tumor parts including tumor cells and immune cells within the tumor.

### Patients and methods

This study is a retrospective analysis of a prospectively acquired patient sample of consecutive HNSCC patients undergoing MRI evaluation for staging purposes. It was approved by the institutional review board (Ethic committee of the University of Leipzig, study codes 180-2007, 201-10-12072010, and 341-15-05102015) and every patient gave their written consent.

#### Patients

Overall, 29 patients with primary HNSCC of different localizations were involved in the study. There were 8 (27.6\%) women and 21 (72.4\%) men with a mean age of 55.2 ± 10.8 years, range 33 - 77 years. The patients received no specific treatment before the MRI.

Table 1 gives an overview of the included patients. In 12 patients (37.5\%) moderately differentiated tumors (G2), and in 17 cases (62.5\%) poorly differentiated carcinomas (G3) were diagnosed. The identified tumors were staged as T2 in 6 pa-
tients (20.7%), T3 in 10 patients (34.5%), and as T4 in 13 cases (44.8%). Most patients (n = 25, 86.2%) had nodal metastases.

**DWI**

DWI was obtained by using an axial EPI (echo planar imaging) sequence with b-values of 0 and 800 s/mm² (TR/TE: 8620/73 ms, slice thickness 4 mm, and voxel size 3.2 × 2.6 × 4.0 mm) on a 3T MRI scanner (Siemens Biograph mMR; Siemens Healthcare, Erlangen, Germany) within a routinely acquired MRI protocol. ADC maps were automatically generated by the implemented software. DWI data was processed with a custom-made Matlab-based application (The Mathworks, Natick, MA, USA). On the ADC maps a volume of interest was manually drawn at tumor boundary in accordance to the T1-weighted contrast enhancing tumor areas using all slices, as performed before. All measures were performed by one experienced author (AS, 16 years of general radiological experience and 6 years of experience on the field of head and neck radiology) blinded to the histopathology results. The following histogram parameters were calculated: mean, maximum, minimum, median, mode, 10th, 25th, 75th and 90th percentile as well as kurtosis, skewness and entropy.

**PD-L1 expression analysis**

The histopathology analysis was performed by an experienced board-certified pathologist blinded to the imaging results. PD-L1 expression was retrospectively evaluated by using an immunohistochemistry (IHC) assay, a routinely used version of PD-L1 (SP142) IHC assay (Roche Diagnostics, 68305 Mannheim, Germany). The staining protocol used in this study was as described in the instructions for the approved commercial assay. The assay was used to determine PD-L1 expression on tumor samples derived from clinical pretherapeutic biopsy specimen of the primary tumor. The biopsy was performed before the MRI within a short time interval of a few days. Three scoring methods were used to assess the immunohistochemical expression. Firstly, only the amount of positive tumor cells were scored, tumor proportion score (TPS). Of note, membrane as well as cytoplasmatic staining was assessed as positive. Secondly, the immune related cells comprising macrophages and lymphocytes were assessed, immune cell score (ICS). Finally, a combined positive score (CPS) of both was calculated. For all scores, the amount of positive cells was calculated in relation to tumor cells multiplied by 100. All calculations were made on 20-fold magnification.

**Statistical analysis**

Statistical analysis was performed using performed using GraphPad Prism 5 (GraphPad Software, La Jolla, CA, USA). Collected data were evaluated by means of descriptive statistics. Mean values were stated with standard deviation in all instances. Spearman’s correlation coefficient (r) was used to analyze associations between investigated parameters. A two-sided Mann-Whitney-Test was used to test between groups. P-values < 0.05 were taken to indicate statistical significance.

**Results**

Table 2 summarizes the investigated ADC histogram parameters.

For the PD-L1 expression of the TPSa mean value of 3.1 ± 14.8 was identified. Overall, 23 tumors (79.3% of all patients) showed no expression.

Furthermore, for the PD-L1 expression of the CPS a mean value of 4.7 ± 14.0 was identified and 13 tumors (45.0%) showed no expression.

Finally, for the PD-L1 expression of the ICS a mean value of 2.5 ± 3.8 was identified. 13 patients (45.0%) showed no expression.

| Parameter | Mean ± standard deviation, x10⁻³ mm²/s | Range, x10⁻³ mm²/s |
|-----------|---------------------------------------|--------------------|
| Mean      | 1.14 ± 0.22                           | 0.78-1.68          |
| Min       | 0.70 ± 0.25                           | 0.17-1.24          |
| Max       | 1.78 ± 0.32                           | 1.35-2.39          |
| P10       | 0.90 ± 0.21                           | 0.54-1.41          |
| P25       | 1.0 ± 0.21                            | 0.64-1.50          |
| P75       | 1.27 ± 0.24                           | 0.87-1.82          |
| P90       | 1.41 ± 0.26                           | 0.94-2.03          |
| Median    | 1.12 ± 0.22                           | 0.76-1.64          |
| Mode      | 0.97 ± 0.28                           | 0.78-1.55          |
| Kurtosis  | 3.74 ± 1.40                           | 2.23-7.93          |
| Skewness  | 0.49 ± 0.49                           | -0.54-1.49         |
| Entropy   | 2.44 ± 0.49                           | 1.67-3.75          |

P = percentile
There was a moderate correlation between TPS and CPS ($r = 0.57$, $p = 0.0015$) and ICS ($r = 0.57$, $p = 0.016$). There was a strong correlation between ICS and CPS ($r = 0.95$, $p < 0.001$).

Regarding associations between ADC histogram parameters and PD-L1-expression scores, there was only one weak correlation identified between skewness and ICS ($r = -0.38$, $p = 0.04$). \( \text{ADC}_{\text{max}} \) tended to correlate with ICS ($r = -0.35$, $p = 0.06$). For TPS, the best association was identified with \( \text{ADC}_{p25} \) ($r = -0.36$, $p = 0.052$).

In discrimination analysis there were no significant differences between TPS negative and positive tumors in regard of ADC histogram parameters.

In tumors with high expression of ICS (5–15 points) skewness was significantly lower than in lesions without ICS expression, $0.09 \pm 0.26$ vs. $0.56 \pm 0.51$ ($p = 0.025$), respectively. Other parameters did not differ significantly.
Discussion

This study sought to elucidate associations between DWI and PD-L1-expression in HNSCC. As shown, a moderate association between skewness derived from ADC values and the immune cell score was identified.

There is increasing body of evidence that different ADC values are capable to reflect tumor microstructure.\(^3\)\(^-\)\(^5\) Thus, results pooled out of many tumor entities suggests that there is an overall moderate inverse correlation between ADC values and tumor cellularity.\(^5\)

Beyond that, there are several studies, which emphasized the fact that ADC values can also predict several biological features of tumors, such as neoangiogenesis and proliferation potential.\(^4\)\(^,\)\(^6\)\(^-\)\(^8\)

Regarding proliferation potential, an overall mod-
erate inverse correlation was identified between ADC values and Ki 67 index pooled for various tumors indicating that ADC can predict the amount of proliferating cells in tissue.

Regarding HNSCC, the relationship between ADC values and Ki 67 index was also identified \( r = -0.61 \). In another study it was reported that ADC values were associated with CD3 positive stained cells \( r = -0.568, p = 0.009 \), which is a measurement of the influx of T-cells within the tumor. Therefore, it can be assumed that ADC values cannot only reflect the whole cell density within the tumor but are sensitive to smaller specific tumor cell fractions. Interestingly, skewness was only associated with the amount of positive PD-L1-stained immune cells. Taken together with the results that ADC values were associated with the amount of tumor invading T-cells within the tumor, there might be a possibility to reflect and predict tumor microenvironment in HNSCC.

The parameter skewness represents the asymmetry of the distribution within the histogram in regard of the ADC values. In short, tumors which lean to the left have consecutively lower ADC values and thus a higher malignancy, whereas tumors, which lean to the right have higher ADC values and thus lower malignancy. Regarding HNSCC, skewness was significantly higher in patients with HPV-mediated oropharyngeal cancer. Interestingly, there is an association between positive human papilloma patients and PD-L1 expression. This might explain why skewness derived from ADC values is related to both histopathological features of HNSCC.

In a recent excellent investigation, a direct correlation between skewness derived from apparent diffusion coefficient (ADC) and the immune cell score (ICS). The resulting Spearman’s correlation coefficient is
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r = -0.38, p = 0.04.
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FIGURE 3. Correlation analysis between skewness derived from apparent diffusion coefficient (ADC) and the immune cell score (ICS). The resulting Spearman’s correlation coefficient is \( r = -0.38, p = 0.04 \).

Recently, there is extensive interest about PD-1 and PD-L1-inhibitors as a form of immunotherapy in oncology. In melanoma patients, PD-1 and PD-L1 inhibitors showed the first very promising results. The effectiveness of this immunotherapy was shortly after also shown for non-small lung cancer and other tumor entities.

Regarding HNSCC, there were several trials establishing the effectiveness of this novel treatment in the recurrent setting. In these trials, PD-L1 status was defined as negative or positive with at least 1% of positive staining, well comparable with the approach in the present study.

In one study the probability of response was not significantly different between PD-L1 negative and positive patients, when only tumor cells were calculated but only when immune cells were also included into the analysis. Correspondingly to our results, skewness was only significantly associated with the scoring system regarding immune cells, reflecting these small, yet not negligible differences in PD-L1 staining scores. As shown in our results, there are significant differences between the scores, which needs to be considered. The immune cell score only measures the tumor associated PD-L1 cells and not the expression of the cancer cells themselves.

Moreover, it is well known that HNSCC is a highly heterogeneous tumor with different biological behavior comprising different localizations and HPV-mediated oropharyngeal cancer. Beyond that, even intratumoral heterogeneity was described as very important in HNSCC, which the amount of heterogeneity differing between primary localization. This might be important for biopsy sampling, as the tumor might significantly show different PD-L1 expression in several biopsy sites. This is, moreover, of special interest, as PD-L1 expression in head and neck cancer is localized at the tumor borders in HPV-positive cancers. Presumably, only imaging with a quantitative analysis approach can display this tumor.
heterogeneity completely during clinical routine. There might also be the opportunity for ADC histogram parameters to indicate treatment changes of the tumors during PD-L1 treatment.

To date, only one other study investigated possible associations between ADC values and PD-L1 expression in other tumors in oncology.\(^3\) In this study, there was a reported statistical difference between PD-L1 positive compared to negative tumors in regard of ADCmax in rectal cancers.\(^3\) Correspondingly, there was a statistical trend in the present study for ADCmax to correlate with the ICS, but it did not reach statistical significance, presumably due to small sample size. More studies are needed to investigate, whether ADCmax might be a promising biomarker to predict PD-L1 expression in oncologic imaging.

To overcome this limitation, further studies are needed with a multiparametric approach including MRI, CT and PET data guided with artificial intelligence. Presumably, non-invasive prediction of histopathology by imaging can be translated into clinical routine employing this approach.

There are several limitations of the present study to address. First, it is a retrospective analysis performed on a prospectively recruited patient sample. However, the imaging and pathology analyses were performed independently to each other to reduce possible bias. Second, the imaging analysis was performed as a whole tumor measurement, whereas the PD-L1 analysis was performed on biopptic specimens with possible spatial incongruencies. Yet, this approach represents daily clinical practice for which no whole lesion pathology-radiology correlation is possible. Third, the ADC measurement was performed by one reader with possible accompanied reader-bias. However, it is reported that ADC values have a good to excellent interreader variability in HNSCC, which might reduce this limitation. Fourth, because of the exploratory study design we did not correct for multiple tests, which has an impact on the presented results. Fifth, the biopsy was taken from the primary tumor before the MRI. This might cause biopsy-related artifacts of the tumor, such as hematoma or local inflammation, which can have an impact on the ADC values.

In conclusion, here is a weak association between skewness derived from ADC values and PD-L1 expression in HNSCC, which might not be strong enough to predict PD-L1 expression in clinical routine. Presumably, ADC values are more influenced by complex histopathology compartments, comprising cellular and extracellular aspects of tumors than only of a single subset of tumor associated cells.

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