Interval changes of histogram-derived diffusion indices predict treatment response to induction chemotherapy in head and neck cancer: a feasibility study

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**Background:** This retrospective study investigated whether the interval change of apparent diffusion coefficient (ΔADC) [baseline and after the first cycle of induction chemotherapy (ICT)] can be used as a valid predictive imaging biomarker of the treatment response to ICT in head and neck cancer (HNC).

**Methods:** A total of 19 consecutive patients with HNC who underwent diffusion-weighted magnetic resonance imaging (DWI) at baseline and after the first cycle of ICT were included. Whole-tumor ADC histogram parameters (mean, median, kurtosis, skewness, entropy, minimal, maximum, 25th percentile, and 75th percentile) were obtained. The correlations of ΔADC histogram parameters, volume, T-stage, N-stage, and age with the treatment response were examined using the Mann–Whitney U test. The predictive value of histogram parameters was examined using receiver operating characteristic (ROC) curves.

**Results:** Responders showed significantly higher values of ΔADC<sub>25</sub> (0.19±0.23) and ΔADC<sub>min</sub> (1.78±2.98) than non-responders (−0.09±0.15 and −0.73±0.36; P=0.035 and 0.009, respectively). When ΔADC<sub>25</sub> and ΔADC<sub>min</sub> were used for predicting the treatment response, the area under the ROC curve was 0.850/0.933 with a sensitivity of 73.3%/80.0% and specificity of 100%/100% (P=0.036 and 0.009, respectively).

**Conclusions:** ΔADC<sub>25</sub> and ΔADC<sub>min</sub> derived from whole-tumor histogram analysis are valuable imaging biomarkers for the early prediction of the ICT response in HNC.

**Keywords:** Head and neck cancer (HNC); magnetic resonance imaging (MRI); diffusion; histogram

Submitted Mar 20, 2022. Accepted for publication Aug 09, 2022.
doi: 10.21037/qims-22-263

View this article at: https://dx.doi.org/10.21037/qims-22-263

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Introduction

The Global Burden of Disease study estimated 8,900,000 new head and neck cancer (HNC) cases worldwide in 2017, representing 5.3% of all cancer cases (1). Although radiation therapy or concurrent chemoradiation therapy (CCRT) remains the mainstay of treatment for HNC, induction chemotherapy (ICT) can reduce loco-regional relapse and distant metastasis. Numerous studies have investigated the role of ICT in HNC, and available evidence of whether ICT is superior to standard care (concurrent chemoradiotherapy) is inconclusive (2). In addition to the controversial role of treatment efficacy, the adverse events of ICT, such as hematological and renal toxicity, are the main concerns. In some studies, ICT treatment-related mortality has been reported to be as high as 6–7% (3,4). Therefore, early reliable predictive biomarkers that can reveal whether HNC patients would benefit from ICT are urgently required.

The apparent diffusion coefficient (ADC) map derived from diffusion-weighted magnetic resonance imaging (DWI) provides physiological information on tumor cellularity (5). Water diffusivity within the tumor can reflect changes in tumor cellularity, which occur after ICT, radiotherapy, or concurrent chemoradiotherapy. Pathophysiological changes revealed through DWI are usually evident before morphologic changes (6), with the potential use of these changes in early response assessments. A systematic review demonstrated that high pretreatment ADC and a low increase in early intratreatment or posttreatment ADC were potential indicators of loco-regional failure in patients with HNC receiving chemoradiation therapy (7). Several studies (8-10) have investigated treatment responses to concurrent chemoradiation therapy or radiotherapy at different time points by using the mean or median value of ADC. However, HNC is typically heterogeneous; thus, ADC assessments are insufficient to reveal the diverse conditions of HNC. Recently, histogram analysis (11), which reflects the distribution and variation of all voxels and considers tumor heterogeneity, has been used in ADC studies, showing promising results for the prediction of the treatment response (12-14). Studies evaluating the predictive value of DWI for treatment response early after ICT are limited, and the optimal parameters of ADC analysis for predicting the treatment response have not been well established. Thus, reliable surrogate biomarkers should be identified that can be used for the individualization of treatment strategies.

This study evaluated the usefulness of interval change of ADC (ΔADC; derived from histogram analysis) between pretreatment and posttreatment (i.e., after one cycle of ICT) for the early prediction of treatment response to ICT in patients with HNC and the ability of derived indices to provide prognostic information for the patients. We present the following article in accordance with the STARD reporting checklist (available at https://qims.amegroups.com/article/view/10.21037/qims-22-263/rc).

Methods

Patients

The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013). In our institution, pre- and post-ICT MRI scans were standard of care protocol in our clinical practice for HNC patients. And informed consent was also signed by the patients before the image studies. In this study, we retrospectively analyzed these MRI images to identify the radiomic features and the study was also approved by the local institutional review board of Chung Shan Medical University Hospital (No. CS1-21105).

Twenty-six consecutive patients with histologically confirmed primary HNC undergoing two MR examinations before and 1 week after the first cycle of ICT were enrolled between June 2018 and January 2019. Seven patients who received other treatments before ICT, who had poor-quality MR images due to a low signal-to-noise ratio or artifacts, or who could not complete ICT because of side effects were excluded. Eventually, 19 patients were eligible for inclusion in the present analysis (Figure 1). Patient characteristics are displayed in Table 1. Staging was performed for each patient according to the 8th edition of the American Joint Committee on Cancer (AJCC) Cancer Staging Manual, Head and Neck Section. All participants were followed up for at least 2 years.

ICT and treatment response evaluation

The ICT regimen, which typically consists of docetaxel plus cisplatin and 5-fluorouracil (TPF), was modified according to the EORTC 24971/TAX 323 study (15). The modified TPF regimen consisted of 20 mg/m² docetaxel administered as a 1-h infusion on days 1 and 8, 50 mg/m² cisplatin infusion administered 2 h on day 1, 200 mg/m² leucovorin infusion administered on day 1, and intravenous
bolus 5-fluorouracil at 400 mg/m$^2$ administered for 30 min on day 1, followed by intravenous continuous infusion of 1,200 mg/m$^2$ 5-fluorouracil for 48 h on day 1. The regimen was repeated every 2 weeks, and a total of three cycles of modified TPF was administered to patients. The Response Evaluation Criteria in Solid Tumors 1.1 (RECIST 1.1) criteria were used to evaluate the response to the modified TPF regimen (16).

**MR examination**

All data were acquired on a 3.0T clinical MR scanner (MAGNETOM® Skyra; Siemens Healthcare) with a 20-channel head and neck coil, covering the range from the level of the skull base to the thoracic inlet. Images were obtained with patients in the supine position. After three-plane tripilot imaging, turbo spin echo-based fat-saturated T2-weighted axial images (TR/TE =4,000/86 ms; matrix size =320×320; field of view =220×220 mm$^2$; slice thickness =5 mm; bandwidth =400 Hz/Px; acquisition time =3 min and 6 s) and T1-weighted axial images (TR/TE =602/13 ms; matrix size =320×320; field of view =230×230 mm$^2$; slice thickness =5 mm; bandwidth =240 Hz/Px; and acquisition time =1 min 52 s) were acquired. In addition, readout-segmented echo-planar DWI with two-dimensional navigator-based reacquisition (TR/TE =5,800/63 ms; matrix size =160×160 (zero-filled to 320×320); slice thickness =5 mm; no intersection gap; slice thickness =5 mm; number of slices =32; iPAT =2; bandwidth...
=919 Hz/Px; readout segments =5; echo spacing =0.34 ms; b value =0 and 800 s/mm²; and acquisition time =2 min 15 s] was performed. The motion-probing gradients were placed along three orthogonal axes with the same strength.

**Data analysis**

All measurements were performed by a neuroradiologist (with more than 10 years of experience in head and neck CT imaging), who was blinded to clinical and survival data. All primary tumors were evaluated through DWI performed from 1 to 3 days before the start of treatment and performed after the first cycle of ICT. The obtained data were transferred to an offline PC and analyzed using MATLAB 2018b software (MathWorks, Natick, MA, USA). All images in our study were accurately registered first, and then, ADC maps were calculated using signal intensities of the corresponding DWI images. Polygonal regions of interest (ROIs) were manually drawn on the derived ADC maps along the contours of the primary tumor on each slice (whole-lesion measure); the segmentation was aided with side-by-side visualization of T2-weighted and DWI images. ROIs in the lesions were carefully drawn with the inclusion of the solid portions of the lesions and the exclusion of any obviously cystic or necrotic areas with reference to T2WI (6,17). Subsequently, the mean, median, 25th percentile (ADC_{25}), 75th percentile (ADC_{75}), minimum, maximum, skewness, kurtosis, and entropy within the whole volume were derived through histogram analysis. Kurtosis indicates the histogram peakedness (the lower the kurtosis, the more flat the histogram); skewness is related to histogram symmetry (positive skewness indicates a right-tailed histogram); and entropy is a metric positively associated with image heterogeneity. The nth percentile is the point at which n% of the voxel values that form the histogram are found to the left. The interval changes (Δ) of the aforementioned parameters were generated by calculating the difference in pretreatment and posttreatment (after the first cycle of ICT) ADC values divided by the pretreatment ADC value. For example, ∆ADC = (ADC_{1st} − ADC_{pre})/ ADC_{pre}, where ADC_{pre} represents pretreatment ADC value, and ADC_{1st} represents the ADC value after the first cycle of ICT. In addition, ROIs were used to measure the whole-tumor volume. In each primary tumor, the whole-tumor volume was calculated by multiplying each cross-sectional area by the section thickness.

**Statistical analysis**

The histogram indices of the interval change of ADC (including ∆ADC_{min}, ∆ADC_{max}, ∆ADC_{mean}, ∆ADC_{median}, ∆ADC_{kurtosis}, ∆ADC_{skewness}, ∆ADC_{entropy}, ∆ADC_{25} and ∆ADC_{75}) and the interval change of the primary tumor volume (∆TV) as well as other clinical variables such as age, T-stage (T1–2 versus T3–4) and N-stage (N0–1 versus N2–3) were compared between responders and non-responders by using a Mann–Whitney U test. Overall survival (OS) was defined as the time from the start of ICT to the occurrence of any cause of death; the data of patients who were alive at the end of follow-up were censored on that date. The survival curve for OS was generated using the Kaplan–Meier method, and differences in survival curves between responders and non-responders were tested for significance by using the log-rank test. A receiver operating characteristic (ROC) analysis with the area under the curve was used to investigate the discriminatory capability of the significant predictive values of the responders. To calculate the sensitivity, specificity, and accuracy of the significant predictive value of the responders, the optimal threshold was determined by giving equal weighting to sensitivity and specificity on the ROC curve. Statistical calculations were performed using statistical analysis software (Statistical Package for the Social Sciences, Version 18.0; IBM, Armonk, New York, USA), and a P value of <0.05 was considered statistically significant after correction for multiple comparisons.

**Results**

**Patient characteristics**

In this study, the data of 19 patients were included in analysis (17 men and 2 women; mean age, 56 years; range, 43–75 years), with the majority having oral cavity cancers (10/19). All patients had completed three cycles of ICT, with a median follow-up of 25 months (range, 4–31 months). According to the RECIST 1.1 criteria, HNC patients showing at least a partial response to ICT were identified as responders (n=15), and those showing stable or progressive disease were classified as non-responders (n=4). The detailed demographic characteristics are displayed in Table 1.

**Analysis of histogram indices**

Figures 2,3 show two representative cases of responders and
non-responders, including the T2-weighted image, DWI image, ADC map, and ADC histogram acquired before (Figure 2A-2D) and after one-cycle (Figure 2E-2H) of ICT, respectively. Compared with non-responders, a decreased tumor volume was obtained on the T2-weighted and DWI images of responders, as well as an elevation of tumor water diffusion after ICT. Comparisons of histogram indices between responders and non-responders are summarized in Table 2. Although no significant differences were found in $\Delta$ADC$_{\text{mean}}$, $\Delta$ADC$_{\text{median}}$, $\Delta$ADC$_{\text{75}}$, $\Delta$ADC$_{\text{max}}$, $\Delta$ADC$_{\text{skewness}}$, $\Delta$ADC$_{\text{kurtosis}}$, $\Delta$ADC$_{\text{entrophy}}$, ATV, age, T-stage, and N-stage between responders and non-responders ($P>0.05$), significant differences were found in $\Delta$ADC$_{\text{min}}$ (1.78±2.98 vs. -0.73±0.36, $P=0.009$) and $\Delta$ADC$_{25}$ (0.19±0.23 vs. -0.09±0.15, $P=0.035$) between the responders and non-responders.

### Analyses of ROC and OS

Furthermore, ROC analyses were used to investigate the feasibility of using interval changes of ADC to predict HNC patients’ responses to ICT. The area under the ROC curve of $\Delta$ADC$_{\text{min}}$ and $\Delta$ADC$_{25}$ was 0.933 [95% confidence interval (CI): 0.720–0.997] (Figure 4A) and 0.850 (95% CI: 0.614–0.970) (Figure 4B), respectively. The highest sensitivity (80.0%, 73.3%) and specificity (100%, 100%) was obtained using the cutoff probability based on the Youden index when $\Delta$ADC$_{\text{min}}$ and $\Delta$ADC$_{25}$ were selected. OS did not show significantly statistically differences between responders and non-responders (Figure 5).

### Discussion

The present study demonstrated the feasibility of interval change assessments in ADC through whole-tumor histogram analysis to predict the response to ICT in patients with HNC, contributing to the individualization of treatment strategies. Significantly higher $\Delta$ADC$_{\text{min}}$ and $\Delta$ADC$_{25}$ values were found in patients responding positively to ICT than...
in non-responders, probably because of elevated water diffusion resulting from low cellularity in the responsive tumor regions. Specifically, the results of ROC analysis revealed high AUC (0.933, 0.850) for $\Delta ADC_{\text{min}}$ and $\Delta ADC_{25}$, yielding high diagnostic sensitivity (80.0%, 73.3%) and specificity (100%, 100%), which suggests the superior discrimination ability of $\Delta ADC_{\text{min}}$ and $\Delta ADC_{25}$ in predicting therapeutic responses to ICT.

Several studies have indicated that the ADC change between pre- and intratreatment could be useful in predicting the treatment response to ICT, radiation therapy alone, or chemoradiotherapy in HNC. Better responses were revealed for tumors showing significant increases in ADC in the early phase of treatment compared with those with little or no ADC increase (6,7,18-21). Wong et al. (21) further demonstrated that a serial change in ADC was a stronger marker than single ADC measurement at pre- or intratreatment for the prediction of the treatment response in patients with locally advanced HNC after CRT. However, another report demonstrated no significant difference in ADC values between responders and non-responders after two cycles of ICT (19), implying the possible variation of mean ADC during the therapeutic period. Our finding is consistent with the findings of the study; no significant differences were obtained in the interval changes of the mean ADC and tumor volume between the two groups, supporting the possible limitation of conducting conventional comparisons of the mean ADC values.

Several investigators have used the mean (9,10) or median (8,22) ADC to predict the treatment response to ICT or chemoradiotherapy in HNC. However, HNC is typically heterogeneous, and this heterogeneity may be due

Figure 3 A non-responder of a 54-year-old man with histologically proven left maxillary cancer. (A-D) Before induction chemotherapy. (E-H) After one cycle of induction chemotherapy. (A) An axial T2-weighted image showing left maxillary cancer with intermediate signal intensity. (B) The corresponding DWI image with the identical lesion for the reconstruction of ADC measurements. (C) The corresponding ADC map with the identical lesion. (D) A whole-lesion histogram analysis based on the ADC map before induction chemotherapy. (E) An axial T2-weighted image showing stable size of left buccal cancer with intermediate signal intensity. (F) The corresponding DWI image with the identical lesion for the reconstruction of ADC measurements. (G) The corresponding ADC map with the identical lesion. (H) A whole-lesion histogram analysis based on the ADC map after one cycle of induction chemotherapy. DWI, diffusion-weighted magnetic resonance imaging; ADC, apparent diffusion coefficient.
Table 2 ΔADC parameters, ΔTV, T-stage, and N-stage of the primary tumor and patient age for prediction of treatment response

| Variables               | All patients (n=19) | Responders (n=15) | Non-responders (n=4) | P value |
|-------------------------|---------------------|-------------------|----------------------|---------|
| ΔADC mean               | 0.09±0.20           | 0.12±0.21         | −0.04±0.09           | 0.271   |
| ΔADC median             | 0.11±0.19           | 0.15±0.20         | −0.04±0.08           | 0.089   |
| ΔADC 25                 | 0.13±0.24           | 0.19±0.23         | −0.09±0.15           | 0.035*  |
| ΔADC 75                 | 0.08±0.20           | 0.11±0.21         | 0.00±0.05            | 0.317   |
| ΔADC kurtosis           | 0.05±0.48           | 0.07±0.53         | 0.02±0.30            | 0.764   |
| ΔADC skewness           | −0.16±1.09          | −0.05±1.16        | −0.55±0.73           | 0.484   |
| ΔADC entropy            | 0.01±0.07           | 0.02±0.07         | −0.04±0.05           | 0.089   |
| ΔADC min                | 1.25±2.83           | 1.78±2.98         | −0.73±0.36           | 0.009*  |
| ΔADC max                | −0.03±0.24          | −0.08±0.19        | 0.17±0.31            | 0.110   |
| ΔTV                     | −0.28±0.31          | −0.34±0.23        | −0.04±0.48           | 0.271   |
| T stage (T1–2 vs. T3–4) | 7/12                | 6/9               | 1/3                  | 0.591   |
| N stage (N0–1 vs. N2–3) | 9/10                | 7/8               | 2/2                  | 0.908   |
| Age (years)             | 55.9±6.91           | 56.5±7.74         | 54.0±0.82            | 0.315   |

*, statistically significant after Mann–Whitney U test. ΔADC, interval changes of ADC values were generated by calculating the difference in pretreatment and posttreatment (after one cycle of induction chemotherapy) ADC values divided by the pretreatment ADC value; ΔTV, interval changes of primary tumor volume were generated by calculating the difference in pretreatment and posttreatment (after one cycle of induction chemotherapy) primary tumor volume divided by the pretreatment primary tumor volume; ADC, apparent diffusion coefficient; TV, primary tumor volume.

Figure 4 Results of ROC curves of ΔADC min and ΔADC 25. (A) The AUC for ΔADC min was 0.850 (P=0.036). (B) The AUC for ΔADC 25 was 0.933 (P=0.009). ROC curves show the good performance of ΔADC min and ΔADC 25. ROC, receiver operating characteristic; AUC, areas under the ROC curve; ADC, apparent diffusion coefficient.
to areas with different cellularity, necrosis, and stroma and areas with increased or decreased vascularity. The mean and median ADC may be suboptimal, and they are not always significantly sensitive to small changes or treatment effects, because when areas with different ADC values are included in ROI, the effects of heterogeneity are smoothed out. The histogram analysis of ADC values within the primary tumor has been frequently used to adaptively evaluate the heterogeneity of the primary tumor relative to the complex tumor microenvironment (11). Such analysis has proven helpful in predicting the histologic grade (23), HPV status (24), and treatment response (25,26) in patients with HNC. In the present study, although primary tumor $\Delta$ADC$_{\text{mean}}$ and $\Delta$ADC$_{\text{median}}$ values were higher in responders than in non-responders, which is compatible with the results of previous studies (18,20), this finding did not reach statistical significance. By contrast, $\Delta$ADC$_{\text{min}}$ and $\Delta$ADC$_{25}$ in whole-primary-tumor assessments showed significant differences between responders and non-responders, indicating the potential of using ADC histogram analysis to extract effective surrogate biomarkers for prognosis prediction.

$\Delta$ADC$_{\text{min}}$ or $\Delta$ADC$_{25}$ derived from histogram analysis has been applied for assessing chemotherapy responses in malignant bone tumors (27,28), pancreatic cancer (29), or gynecological cancer (30). Saleh et al. (27) and Oka et al. (28) have used $\Delta$ADC$_{\text{min}}$ to monitor responses to chemotherapy in osteosarcoma or Ewing’s sarcoma, demonstrating that $\Delta$ADC$_{\text{min}}$ values were significantly higher in patients with good responses than in those with poor responses. Kyriazi et al. (30) also found that changes in all ADC histogram parameters after the first and third cycles of chemotherapy were higher in responders than in non-responders, indicating the superior discrimination ability of $\Delta$ADC$_{25}$ in predicting responses to chemotherapy in patients with metastatic ovarian or primary peritoneal cancer. For responders, due to cell shrinkage and death after treatment, the increase in ADC would occur as a result of an increase in the fractional volume and diffusion of water molecules in the extracellular space. A strong negative correlation exists between ADC values and tumor cellularity (5), indicating that the tumor regions with the lowest or lower diffusivity (i.e., the highest or higher cellularity) may be more sensitive to chemotherapy or radiotherapy. This might explain why $\Delta$ADC$_{\text{min}}$ or $\Delta$ADC$_{25}$ is the most significant predictor of treatment response in cancer patients. The present study demonstrated the superior performance of $\Delta$ADC$_{\text{min}}$ and $\Delta$ADC$_{25}$ in predicting the response to ICT, which is in agreement with the previous report.

Approximately 20–30% of patients do not respond to ICT (31-33), and our results showed similar results in that four of our enrolled patients (4/19, 21%) did not show responses to ICT. All of these four patients received concurrent chemoradiotherapy, but three died before the study endpoint due to disease progression, which implies that a poor response to ICT might be a poor prognostic factor of OS. Two (patient 1 and 2 in Table 3) patients received additional target therapy with cetuximab after ICT failure and showed longer survival than those without therapy. Several studies (34,35) have shown that cetuximab can improve survival outcomes in advanced HNC. Taken together, these findings indicate that the early identification of potential poor responders to ICT and an early shift in treatment regimens to target therapy or immune therapy may improve survival outcomes in HNC.

Our study has some limitations. First, this was a retrospective study with a small patient population with varying tumor locations. Although non-responders tended to have shorter survival times than responders, no significant difference was found, possibly resulting from the small and heterogeneous patient population in this study. The main purpose of this study was to identify early predictors of treatment response to avoid ineffective

Figure 5 Overall survival rates according to responsiveness to ICT in patients with HNC. HNC, head and neck cancer; ICT, induction chemotherapy.
ICT. A multicenter prospective study with a larger sample size should be conducted to validate the clinical effectiveness of our findings. Second, ROIs were manually drawn by one observer, which limits reproducibility. A comprehensive investigation of interobserver, intraobserver, or intersoftware variances is required. Third, the second DWI was performed after one cycle of ICT, which would not reveal the evolution of the ADC pattern during ICT. A longitudinal study with serial ADC measurements would be useful. Fourth, performing DW imaging in the regions of the head and neck is still challenging. Although the use of the segmented-readout EPI scheme could alleviate susceptibility artifacts, problems with patient movement from swallowing cannot be effectively resolved. Further reduction of the total acquisition time by using advanced imaging sequences would be helpful.

In conclusion, our findings demonstrated that ADC histogram analysis can be used to extract potential surrogate biomarkers for the early prediction of the treatment response to ICT. ∆ADCmin and ∆ADC25 before and after one cycle of ICT outperformed other histogram indices, showing promising diagnostic efficacy for predicting responsiveness to ICT. These early predictive biomarkers may help avoid ineffective treatments and unnecessary toxicity, enabling further individualization of treatment strategies.

Acknowledgments

This manuscript was edited by Wallace Academic Editing. Funding: This work was supported by a grant (#MOST 108-2221-E-040-007-MY3) at the Ministry of Science and Technology, Taiwan.

Footnote

Reporting Checklist: The authors have completed the STARD reporting checklist. Available at https://qims.amegroups.com/article/view/10.21037/qims-22-263/rc

Conflicts of Interest: All authors have completed the ICMJE uniform disclosure form (available at https://qims.amegroups.com/article/view/10.21037/qims-22-263/coif). PHT reports that this work was supported by a grant (#MOST 108-2221-E-040-007-MY3) at the Ministry of Science and Technology, Taiwan. The other authors have no conflicts of interest to declare.

Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013). The study was approved by the local institutional review board of Chung Shan Medical University Hospital (No. CS1-21105) and informed consent was signed by the patients before the image studies.

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Cite this article as: Cheng KL, Lu HJ, Lin X, Wang HY, Chou YH, Tyan YS, Tsai PH. Interval changes of histogram-derived diffusion indices predict treatment response to induction chemotherapy in head and neck cancer: a feasibility study. Quant Imaging Med Surg 2022;12(12):5383-5393. doi: 10.21037/qims-22-263