Correlations of tumour permeability parameters with apparent diffusion coefficient in nasopharyngeal carcinoma

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ABSTRACT

Background and Purpose: Functional imaging has an established role in therapeutic monitoring of cancer treatments. This study evaluated the correlations of tumour permeability parameters derived from dynamic contrast-enhanced magnetic resonance imaging (DCE-MRI) and tumour cellularity derived from apparent diffusion coefficient (ADC) in nasopharyngeal carcinoma (NPC).

Material and Methods: Twenty NPC patients were examined with DCE-MRI and RESOLVE diffusion-weighted MRI (DW-MRI). Tumour permeability parameters were quantitatively measured with Tofts compartment model. Volume transfer constant (Ktrans), volume of extravascular extracellular space (EES) per unit volume of tissue (Vve), and the flux rate constant between EES and plasma (Kep) from DCE-MRI scan were measured. The time-intensity curve was plotted from the 60 dynamic phases of DCE-MRI. The initial area under the curve for the first 60 s of the contrast agent arrival (IAUC60) was also calculated. They were compared with the ADC value derived from DW-MRI with Pearson correlation analyses.

Results: Among the DCE-MRI permeability parameters, Kep had higher linearity in inverse correlation with ADC value (r = -0.60, p < 0.05), Ktrans (r = -0.60, p < 0.05) and IAUC60 (r = -0.64, p < 0.05) also had significant inverse correlations with ADC. Vve showed a significant positive correlation with ADC (r = 0.63, p < 0.05).

Conclusions: Nasopharyngeal tumour vascular permeability parameters derived from DCE-MRI scan were correlated linearly with tumour cellularity measured by free water diffusability with ADC. The clinical implementations of these linear correlations in the quantitative assessments of therapeutic response for NPC patients may be worth to further explore.

1. Introduction

Nasopharyngeal carcinoma (NPC) is a prevalent malignant disease with remarkably high incidence in southern China and Southeast Asia. Intensity-modulated radiotherapy is the standard treatment and has shown excellent tumour control and disease-free survival in the treatment of NPC [1–3]. Concurrent chemotherapy with or without an induction course is also required for locally advanced disease [4].

When a tumour grows larger than 2–3 cubic millimetres in size, oxygen and nutrient supplies by simple diffusion are no longer efficient enough to support the rapid cell proliferation [5]. The tumour oxygen tension triggers the hypoxia-inducible factor 1-alpha (HIF-1α) expression, which in turn induces the release of pro-angiogenic factors. A concentration gradient of pro-angiogenic factors is developed surrounding the hypoxic tumour. When they reach the nearby blood vessel, they trigger basement membrane degradations and start tip cell migrations. The tip cells migrate towards the hypoxic tumour and form a new blood vessel [6]. This newly formed blood vessel supports the hypoxic tumour with oxygen and nutrients to continue the tumour growth. Hence, aggressive tumours are often highly vascularised [7–9]. On the other hand, tumour plasma perfusion affects the delivery of chemotherapy and thus can affect the treatment outcome for head and neck tumours.
neck cancers. Both perfusion computed tomography (CT) and dynamic contrast-enhanced magnetic resonance imaging (DCE-MRI) are available diagnostic scans for assessing tumour plasma perfusion. Studies have shown that baseline tumour plasma perfusion is predictive in chemotherapy treatment response for patients with head and neck cancers [10,11]. Studies have indicated that tumour permeability is associated with the diffusivity and clinical stage of NPC [12,13]. Reduced diffusivity is associated with tumour cellulularity which can be used as an indicator for the evaluation of treatments [14]. Diffusion-weighted magnetic resonance imaging (DW-MRI) is one of the standard diagnostic examinations for detecting the diffusivity within a tumour. Apparent diffusion coefficient (ADC) measures the magnitude of diffusion of water molecules within tissue and is inversely proportional to the tumour cellularity [14]. Studies have shown the values of detecting tumour proliferation potential and predicting early treatment response by analysing the pre-treatment ADC in head and neck cancers [15,16]. Studies have indicated that pre-treatment ADC is a prognostic factor for local relapse-free survival and disease-free survival in NPC [17,18].

Early changes in tumour permeability kinetics have clinical values in monitoring chemotherapy response for NPC [19]. The pre-treatment ADC value was also found to be a clinical indicator of chemotherapy response in patients with NPC [20]. Both tumour permeability kinetics and cellularity can be crucial factors affecting chemotherapy efficacy. To our understanding, the correlations between the tumour permeability kinetics and ADC specific to NPC have not been established. The purpose of this study was to analyse the association between tumour permeability parameters and tumour diffusivity in NPC patients, so as to establish the quantitative relation between angiogenesis and cellularity for the potential therapeutic response monitoring.

2. Materials and Methods

This prospective study was approved by the hospital research ethics committee (REC-2017-06) in compliance with the ethical standards for human research in the Declaration of Helsinki and the Good Clinical Practice (GCP). Informed consent was obtained from the patients.

2.1. Patients

Twenty patients with histopathologically confirmed NPC (Table 1) who underwent helical tomotherapy were examined with DCE-MRI and DW-MRI before starting treatments. There were 16 males (80 %) and 4 females (20 %) with a median age of 47 ± 11.6 years (Range 26–66 years). All of them were classified based on the 2017 American Joint Committee of Cancer (AJCC) TNM classification of malignant tumours (8th Edition) [21] as stage II or above who were eligible for chemoradiation treatments.

2.2. MR imaging

The MR scans were originally used for radiation therapy planning. For that reason, all patients were immobilised in thermoplastic casts and scanned with a 1.5 T MAGNETOM Aera MR Simulator (Siemens Healthcare, Erlangen, Germany) dedicated to radiotherapy planning with gradient performance at maximum amplitude 45mT/m and maximum slew rate of 200mT/m/ms. It is equipped with a 70 cm wide bore and an external laser system for precise patient positioning. The scans were done on a flat couch embedded with a posterior 32-channel spine coil array and a pair of 4-channel large flex receiver coils with interfaces which were wrapped around the head region with a pair of bilateral coil holders (Orfit Industry, Wijnegem, Belgium). In order to avoid any retained contrast agent from DCE-MRI which might affect the accuracy of DW-MRI measurement, the DW-MRI was scanned before the DCE-MRI.

2.2.1. DW-MRI

The primary nasopharyngeal lesion of the patient was localised based on the non-contrast morphological images and previous diagnostic radiological reports. The axial DW-MRI images were acquired in an interleaved fashion using a 2D diffusion-weighted multi-shot readout-segment echo-planar imaging (RS-EPI) RESOLVE sequence with GRAPPA acceleration technique. The imaging parameters of DW-MRI were set as FOV 220 × 220 mm; matrix size of 112 × 112; slice thickness of 3.5 mm with 10 % gap; voxel size of 2 × 2 × 3.5 mm; slice number of 20; phase encoding in anteroposterior direction; TR/TE1/TE2 of 3890/51/78 ms (TE1 as the segment echo and TE2 as the navigator echo for phase correction between shots). Diffusion sensitivities (b-values) at high (800 sec/mm²) with average factor of 4 and low (50 sec/mm²) with average factor of 2 were selected. The DW-MRI scans were acquired in 6 min 27 s. The scan volume was set as contiguous axial sections through the primary nasopharyngeal lesion. The rate of diffusion is evaluated by these images with different b-values and quantitatively represented by ADC map.

2.2.2. DCE-MRI

In order to measure the contrast enhancement pattern in the tumour, two axial pre-contrast 3D T1-weighted spoiled gradient-echo (GRE) VIBE sequences using flip angles of 2 and 15 degrees were performed respectively for baseline mapping. These flip angles were chosen as this combination achieved the smallest overestimation for Ktrans in primary tumours of head and neck cancers [22]. The T1 mapping was scanned with FOV of 260 × 260 mm; matrix size of 192 × 154; slice thickness of 3.5 mm; voxel size of 1.4 × 1.4 × 3.5 mm; slice number of 30; TR/TE of 3.8/1.34 ms. Once T1 mapping was performed, the acquisition of DCE-MRI images with the same coverage was started for 3 dynamic phases before the administration of gadobutrol contrast agent. A dose of 0.1 mmol/kg of gadobutrol (Gadovist, Bayer Vital GmbH, Leverkusen, Germany) contrast agent was injected using a power injector at a constant flow rate of 2.5 mL/sec followed by a 20 mL normal saline flushed at the same flow rate. The DCE-MRI was performed using an axial 3D T1-weighted TWIST VIBE sequence with the following parameters: FOV of 260 × 260 mm; matrix size of 192 × 154; slice thickness of 3.5 mm; voxel size of 1.4 × 1.4 × 3.5 mm; slice number of 20; TR/TE of 6.28/2.38 ms; and flip angle of 9 degrees. A total of 60 dynamic phases with temporal resolution of 4.7 s were acquired. The total acquisition time of approximately 4 min 42 s.

The imaging protocols of the DW-MRI and DCE-MRI scans are summarized in Table 2.

### Table 1

Patient and tumour characteristics.

| Age | Range 26–66 | Median 47 |
|---|---|---|
| Sex | Male | 16 | 80 |
|   | Female | 4 | 20 |

| Pathologic T classification | T1 | 7 | 35 |
|---|---|---|---|
|   | T2 | 3 | 15 |
|   | T3 | 6 | 30 |
|   | T4 | 4 | 20 |

| Stage | II | 4 | 20 |
|---|---|---|---|
|   | III | 10 | 50 |
|   | IV | 6 | 30 |
2.3. Post-processing

All the DCE-MRI and DW-MRI image series were post-processed by the image reading application syngo.via (Siemens Healthcare, Erlangen, Germany) developed by MR Simulator vendor. Tumour permeability parameters were quantitatively measured with Tofts compartment model [23,24]. In the Tofts model, each voxel of tissue is differentiated into tissue parenchymal cells, blood vessels and tissue extracellular extravascular space (EES). The influx rate from blood vessels into EES can be measured as volume transfer constant ($K_{\text{trans}}$) and the reflux rate from EES back into plasma can be measured as flux rate constant ($K_{\text{ep}}$) with gadolinium contrast administration [24]. The volume of EES per unit volume of tissue ($V_e$) can also be depicted with the perfusion model. The arterial input function (AIF) was modelled to the gadobutrol contrast agent according to the Tofts model by population-based technique [25] with the AIF curve best fitted to either fast [26], intermediate [27] or slow [28] population-averaged AIF. The $K_{\text{trans}}$, $V_e$ and $K_{\text{ep}}$ from the DCE-MRI scan were measured by positioning a volume of interest (VOI) at the primary nasopharyngeal tumour localised by reviewing patient’s recent diagnostic MR and/or positron emission tomography (PET) scans at where distinct contrast enhancement and/or elevated standardised uptake value (SUV) were observed. The MR signal intensity time course in DCE-MRI was represented by the time-intensity curve (TIC). It was plotted from the 60 dynamic phases of DCE-MRI. The initial area under curve for the first 60 s of the contrast agent arrival (iAUC60) was also calculated. All $K_{\text{trans}}$, $V_e$, $K_{\text{ep}}$ and iAUC60 were compared with the ADC value from DW-MRI with Pearson correlation analyses. The $K_{\text{trans}}$ was compared with disease stages by Kruskal-Wallis test.

3. Results

Examples of the DCE-MRI and DW-MRI images are shown in Fig. 1. Patients with higher TNM-staged diseases tended to have slightly higher median $K_{\text{trans}}$ values. The median $K_{\text{trans}}$ for stage II, III and IV diseases were 0.10 min$^{-1}$ (range 0.06–0.23 min$^{-1}$), 0.13 min$^{-1}$ (range 2.3. Post-processing

Table 2

| Parameter          | DW-MRI     | DCE-MRI    |
|--------------------|------------|------------|
| Slice orientation  | Transversal| Transversal|
| TR (ms)            | 3890       | 6.28       |
| TE (ms)            | 51.78      | 2.38       |
| b values (s/mm$^2$)| 50(2), 800(4)|           |
| Voxel size (mm$^3$)| 2 $\times$ 2 $\times$ 3.5 | 1.4 $\times$ 1.4 $\times$ 3.5 |
| Field of view (mm$^2$)| 220 $\times$ 220 | 260 $\times$ 260 |
| Matrix             | 112 $\times$ 112 | 192 $\times$ 154 |
| Slice thickness (mm)| 3.5       | 3.5        |
| Slice gap (mm)     | 0.35       |            |
| No. of slices      | 20         | 30         |
| Fat suppression    | SPAIR      | CHESS      |
| Readout segments   | 7 shots    |            |
| GRAPPA acceleration factor | 2      |            |
| Flip angle         | 180°       | 9°         |
| Acquisition time (min:s) | 6:27   | 4:42       |

Fig. 2. The Box-and-whisker plot of $K_{\text{trans}}$ values from DCE-MRI among patients with different disease stages. The medians and ranges are shown. The bottom and top edges of boxes represent the 1st and the 3rd quartiles of values respectively.
0.04–0.18 min⁻¹) and 0.14 min⁻¹ (range 0.05–0.17 min⁻¹) respectively (Fig. 2). A mild increase of \( K_{\text{trans}} \) was observed versus increase in disease stage although statistical significances could not be obtained (\( p = 0.871 \)).

The correlations of permeability parameters from DCE-MRI with ADC from DW-MRI are shown in regression plots in Fig. 3. All the permeability parameters acquired by DCE-MRI in NPC tumours showed significant linear correlations with ADC value. The \( K_{\text{trans}} \) (\( r = -0.60, p = <0.05 \)), \( K_{\text{ep}} \) (\( r = -0.69, p = <0.05 \)) and \( \text{iAUC60} \) (\( r = -0.64, p = <0.05 \)) showed significant inverse correlations with ADC value whereas the \( V_e \) showed a significant positive correlation with ADC (\( r = 0.63, p = <0.05 \)).

4. Discussions

This study has shown that tumour cellularity as reflected by ADC was linearly associated with the vascular permeability in nasopharyngeal tumours. In particular, we have identified the reflux rate which was the parameter with the strongest inverse linear correlation with the ADC. This linear correlation between tumour cellularity and vascular permeability in nasopharyngeal tumour might be worth to further investigate for its implementations in therapeutic response monitoring of NPC patients.

Our findings in tumour cellularity and permeability correlation were consistent with known nasopharyngeal tumour biology \([9,29,30]\). Advanced tumour proliferates vigorously, which needs steady supplies of oxygen and nutrients to support the active cell replications. This process is facilitated by tumour angiogenesis. The tumour grows by absorbing oxygen and nutrients for its rapid cell proliferation. In this study, we found that NPC patients with reduced ADC were associated with higher tumour permeability. The \( K_{\text{trans}} \) and \( K_{\text{ep}} \) were inversely correlated with the ADC value in nasopharyngeal tumours, which agreed with the phenomenon of active tumour angiogenesis for advanced tumours with high cellularity \([31–33]\). In particular, the \( K_{\text{ep}} \) exhibited stronger inverse correlation with ADC. This may be explained by the fact that increased vascular permeability promotes additional extravasation of plasma in combination with the effect that the increased tumour cellularity narrows the EES. Both factors contribute to an increase in EES pressure which potentiates a flux tension from EES to plasma. Therefore, the \( K_{\text{ep}} \), reflecting the flux rate constant between the EES and plasma, had higher linearity in inverse correlation.

Advanced tumours are usually densely packed with highly proliferative cells. The presence of numerous membranous and macromolecular structures impedes the free water diffusability within the tumour. The increased cellularity occupies the space within the tumour so that the volume of EES per unit volume of tissue decreases. The \( V_e \) in the Tofts model measures the EES per unit volume of tissue and is mathematically calculated by \( V_e = K_{\text{trans}} / K_{\text{ep}} \). The \( V_e \) represents the ratio of \( K_{\text{trans}} \) to \( K_{\text{ep}} \). We have found a significant positive correlation of the \( V_e \) with the ADC. This positive correlation between \( V_e \) and ADC in nasopharyngeal tumour agrees with permeability and diffusability study findings from other tumours including prostate and brain cancers \([34–36]\). However, it is not the same with some breast cancer studies in which negative correlations between \( V_e \) and ADC were reported \([37,38]\). One of their findings was that breast cancer tumours with high cellularity showed a negative correlation with ADC, whereas those with low cellularity showed a positive correlation.

Fig. 3. Regression plots of permeability parameters from DCE-MRI for \( K_{\text{trans}} \) (A), \( K_{\text{ep}} \) (B), \( V_e \) (C) and \( \text{iAUC60} \) (D) against ADC value from DW-MRI. The Box-and-whisker plots of the corresponding vascular permeability parameters and ADC are also shown to demonstrate their distributions. The bottom/left and top/right edges of boxes represent the 1st and the 3rd quartiles of values respectively.
permeability in nasopharyngeal tumours in our study cohort. Diffusion that advanced nasopharyngeal tumour with high cellularity tends to lower diffusability are leakier in vasculature than others [43]. It suggests restriction as measured by ADC has been used as an indicator of tumour characteristics of a tumour by summarising the contrast agent flush in and early postulations for this contrary correlation in breast cancer is that non-

vessels into EES and the reflux rate from EES back into plasma can be linear correlations in the quantitative assessments of therapeutic permeability and diffusability, the clinical implementations of these parameters derived from DCE-MRI scan were correlated linearly with cellular density measured by free water diffusibility with ADC. The clinical implementations of these linear correlations in the quantitative assessments of therapeutic response for NPC patients may be worth to further explore.

We have found that ADC is linearly associated with the vascular permeability in nasopharyngeal tumours in our study cohort. Diffusion restriction as measured by ADC has been used as an indicator of tumour cellularity and could explain the association with perfusion as demonstrated here, although we acknowledge that DW-MRI at low b-values is sensitive to perfusion effects, and that a low b-value of 50 s/mm² as chosen in our study, may not fully suppress the influence of perfusion on the measured ADC values. Despite this, our findings are in line with several prior studies supporting the hypothesis that proliferative tumours with higher cellularity are likely to have higher vascular permeability. Ni, Liu [13] reported in their study for NPC with DCE-MRI and DW-MRI scans using a 3T MR scanner, that a negative correlation was found with Ktrans and ADC in their cohort. Our results support their findings regarding Ktrans and ADC correlation. They also revealed that the clinical stage of NPC was positively correlated with Ktrans. We also observed a similar trend in our cohort though statistical significance was not achieved. Kooreman, et al. [44] reported an inverse correlation between diffusion coefficient (D) and perfusion fraction (f) in prostate cancer with intravoxel incoherent motion (IVIM) MRI using MR-Linac. Pre-treatment prostatic tumours were found to have lower D but higher f than non-cancerous prostates. Chu, et al. [36] also reported in their study for brain tumours with DCE-MRI and DW-MRI scans that a similar inverse correlation between tumour permeability and diffusability was found. They identified that brain tumours were characterised by lower ADC values but higher volume transfer constant Ktrans. In our cohort of nasopharyngeal tumours, albeit both Ktrans and Kep were found to have significant correlations with ADC, we have found that the flux rate constant Kep has a higher linearity in inverse correlation with ADC values than the Ktrans among the permeability parameters. Svolos, et al. [35] also investigated the permeability and diffusability of brain tumours based on multi-parametric MRI scans including DW-MRI, diffusion tensor imaging (DTI) and dynamic-susceptibility contrast imaging (DSC). They noticed that vascularity and cellularity were factors closely related in brain tumours but they were correlated in a non-linear manner. On the contrary, we observed that this vascularity and cellularity correlation, reflected by permeability and diffusability respectively, was relatively linear in our cohort of nasopharyngeal tumours.

The changes in these correlations between tumour permeability and diffusability after the commencement of chemotherapy treatments can reflect the status changes in angiogenesis and cellularity. Since we have found significant linear correlations between parameters of tumour permeability and diffusability, the clinical implementations of these linear correlations in the quantitative assessments of therapeutic response for NPC patients may be worthwhile to further explore. DCE-MRI is more informative than DW-MRI in which the influx rate from blood vessels into EES and the reflux rate from EES back into plasma can be differentiated for further studies in their individual value for therapeutic response monitoring of NPC. These permeability characteristics can be differentially visualised in parametric maps. As concurrent chemo-radiation is the standard treatment for locally advanced NPC patients, these parametric maps might be useful in monitoring angiogenic changes during chemo-radiation treatments [45]. Since we acquired the scans with an MR Simulator in which the patients were scanned in the same positions as their radiation treatment positions, the parametric maps had the same frame of reference as their radiation treatment plans. The parametric maps might be possible to use for radiation treatment customisations and adaptations with tumour biological changes [46].

Our study on the correlation of permeability parameters and ADC in NPC has limitations. The sample size is relatively small, and further recruitment of NPC patients in this study is needed to arrive at a more robust observation of permeability and diffusivity. Stratification of patients by different T-stages will also be possible with a larger cohort to see if there is any difference in correlations with different tumour sizes. In order to maintain a high temporal resolution for TIC, the DCE-MRI was scanned with limited anatomical coverage to the primary nasopharyngeal tumour only. Cervical lymphatics were not included in the DCE-MRI scans in our study. Further studies including the investigations in permeability and diffusion correlations at cervical lymph nodes are worth considering for a more comprehensive understanding of NPC.

In conclusion, nasopharyngeal tumour vascular permeability parameters derived from DCE-MRI scan were correlated linearly with cellularity measured by free water diffusibility with ADC. The clinical implementations of these linear correlations in the quantitative assessments of therapeutic response for NPC patients may be worth to further explore.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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