**Name of Journal: *World Journal of Radiology***

**ESPS Manuscript NO: 21175**

**Manuscript Type: ORIGINAL ARTICLE**

***Retrospective Study***

**Texture analysis on parametric maps derived from dynamic contrast-enhanced magnetic resonance imaging in head and neck cancer**

Jansen JFA *et al.*Texture analysis of DCE-MRI in HNSCC

**Jacobus FA Jansen, Yonggang Lu, Gaorav Gupta, Nancy Y Lee, Hilda E Stambuk, Yousef Mazaheri, Joseph O Deasy, Amita Shukla-Dave**

**Jacobus FA Jansen,** Department of Radiology, Maastricht University Medical Center, 6211 LK Maastricht, the Netherlands

**Yonggang Lu, Yousef Mazaheri, Joseph O Deasy, Amita Shukla-Dave,** Departments of Medical Physics, Memorial Sloan-Kettering Cancer Center, New York, NY 10065, United States

**Gaorav Gupta, Nancy Y Lee,** Departments of Radiation Oncology,Memorial Sloan-Kettering Cancer Center, New York, NY 10065, United States

**Hilda E Stambuk, Yousef Mazaheri, Amita Shukla-Dave,** Departments of Radiology, Memorial Sloan-Kettering Cancer Center, New York, NY 10065, United States

**Author contributions:** Jansen JFA, Lu Y and Shukla-Dave A designed the study and performed research and analyzed the data; Gupta G and Lee NY included patients; Stambuk HE performed the radiologic assessment; Mazaheri Y and Deasy JO provided support; Jansen JFA, Lu Y and Shukla-Dave A wrote the paper.

# Supported by The National Cancer Institute/National Institutes of Health, No. 1 R01 CA115895.

**Institutional review board statement:** Nineteen patients were enrolled in the institutional review board (IRB) of the Memorial Sloan-Kettering Cancer Center, New York, NY protocol titled “Dynamic Contrast Enhanced MRI and Magnetic Resonance Spectroscopy of Head and Neck Tumors” (IRB No. 06-007).

**Informed consent statement:** All patientsgave informed consent for their participation in the institutionalreview board–approved study. This study was also compliant with the Health Insurance Portability and Accountability Act.

**Conflict-of-interest statement:** All authors have no conflicts of interest with regard to this manuscript.

**Data sharing statement:** Upon formal request and with proper motivation, all original data in anonymized format is available from the corresponding author for local inspection, but cannot leave Memorial Sloan-Kettering Cancer Center.

**Open-Access:** This article is an open-access article which was selected by an in-house editor and fully peer-reviewed by external reviewers. It is distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited and the use is non-commercial. See: <http://creativecommons.org/licenses/by-nc/4.0/>

# Correspondence to: Amita Shukla-Dave, PhD, Director Quantitative Imaging, Associate Attending Physicist, Department of Medical Physics, Memorial Sloan-Kettering Cancer Center, 1275 York Avenue, New York, NY 10065, United States. [davea@mskcc.org](mailto:davea@mskcc.org)

**Telephone:** +1-212-6393184

**Fax:** +1-212-7173010

**Received:** June 29, 2015

**Peer-review started:** July 4, 2015

**First decision:** September 22, 2015

**Revised:** September 24, 2015

**Accepted:** November 23, 2015

**Article in press:**

**Published online:**

**Abstract**

**AIM:** To investigate the merits of texture analysis on parametric maps derived from pharmacokinetic modeling with dynamic contrast-enhanced magnetic resonance imaging (DCE-MRI) as imaging biomarkers for the prediction of treatment response in patients with head and neck squamous cell carcinoma (HNSCC).

**METHODS:** In this retrospective study, 19 HNSCC patients underwent pre- and intra-treatment DCE-MRI scans at a 1.5T MRI scanner. All patients had chemo-radiation treatment. Pharmacokinetic modeling was performed on the acquired DCE-MRI images, generating maps of volume transfer rate (Ktrans) and volume fraction of the extravascular extracellular space (ve). Image texture analysis was then employed on maps of Ktrans and ve, generating two texture measures: Energy (E) and homogeneity (H).

**RESULTS:** No significant changes were found for the mean and standard deviation for Ktrans and ve between pre- and intra-treatment (*P* > 0.09). Texture analysis revealed that the imaging biomarker energy (E) of ve was significantly higher in intra-treatment scans, relative to pretreatment scans (*P* < 0.04).

**CONCLUSION:** Chemo-radiation treatment in HNSCC significantly reduces the heterogeneity of tumors.

# Key words: Dynamic contrast-enhanced magnetic resonance imaging; Tumor heterogeneity; Image texture analysis; Head and neck squamous cell carcinomas

**© The Author(s) 2015.** Published by Baishideng Publishing Group Inc. All rights reserved.

**Core tip**: Head and neck squamous cell carcinoma (HNSCC) is a major form of cancer that still kills many cancer patients, and patients would certainly benefit with improved imaging methodology. The merits of texture analysis were investigated on parametric maps derived from pharmacokinetic modeling with dynamic contrast-enhanced magnetic resonance imaging as imaging biomarkers for the prediction of treatment response in patients with HNSCC, undergoing chemo-radiation treatment. Texture analysis revealed that the imaging biomarker energy (E) of parameter ve was significantly higher in intra-treatment scans, relative to pretreatment scans. This indicates that chemo-radiation treatment in HNSCC significantly reduces the heterogeneity of tumors.

Jansen JFA, Lu Y, Gupta G, Lee NY, Stambuk HE, Mazaheri Y, Deasy JO, Shukla-Dave A. Texture analysis on parametric maps derived from dynamic contrast-enhanced magnetic resonance imaging in head and neck cancer. *World J Radiol* 2015; In press

# INTRODUCTION

Dynamic contrast-enhanced magnetic resonance imaging (DCE-MRI) has become an important tool for the non-invasive quantification of tumor-associated vasculature[1]. By applying pharmacokinetic modeling on sequential T1-weighted images acquired after administration of a contrast agent, one can yield relevant quantitative tumor biomarkers, such as volume transfer rate (Ktrans) and volume fraction of the extravascular extracellular space (ve)[2,3]. For patients with advanced head and neck squamous cell carcinoma (HNSCC), DCE-MRI has shown potential for assessment of treatment response and outcome[4-6]. More specifically, measures derived from DCE-MRI, such as Ktrans and ve, were demonstrated to provide helpful additional information regarding characterization and prognosis of HNSCC with neck nodal metastases[7,8]. HNSCC has been shown to be heterogeneous, due to a chaotic and poorly organized tumor vasculature. Furthermore, tumor heterogeneity is linked with tumor malignancy[9,10]. Tumor heterogeneity of the blood supply may underlie treatment resistance as it prevents therapeutic efficacy. Thus, tumor heterogeneity is highly relevant for the assessment of tumor malignancy and the prediction of treatment outcome. Most studies often report summarizing measures, including mean, median, or standard deviation based on a selection of pixel-by-pixel values, to characterize the whole tumor. However, these often utilized parameters do not always completely characterize the morphologic nature (*i.e.*, heterogeneity) of tumors. It is clinically highly relevant to discover or improve on imaging biomarkers that properly reflect tumor heterogeneity that may assist HNSCC treatment.

A suitable method to assess tumor heterogeneity is image texture analysis[11-15]. During texture analysis, an algorithm is utilized that quantifies spatial intensity coherence of an image, which yields a number of textural features (related to heterogeneity), that are independent of the above mentioned summarizing measures such as mean and standard deviation. An example is the gray-level co-occurrence matrix (GLCM), which is a popular algorithms for texture analysis[11]. Image texture analysis has shown potential in the application of tumor differentiation and treatment response prediction[16-18]. For example, Karahaliou *et al*[17] investigated the feasibility of using texture analysis to quantify the heterogeneity of lesion kinetics and differentiate malignant and benign breast lesions. El Naqa *et al*[12] employed the texture features from PET images to predict treatment response in cervix and head and neck cancers. Alic *et al*[11] found that the tumor response group in limb sarcoma had a high feature of image coherence in pretreatment DCE-MRI Ktrans maps. However, the texture analysis of DCE-MRI pharmacokinetic maps has not been investigated in head and neck cancers for predicting treatment response yet. The purpose of the current study is to assess the merits of GLCM-based texture analysis of parametric maps derived from DCE-MRI pharmacokinetic modeling for the prediction of treatment response in patients with HNSCC.

**MATERIALS AND METHODS**

***Patients***

Our institutional review board approved and issued a waiver of informed consent for this retrospective study, which was compliant with the Health Insurance Portability and Accountability Act. Nineteen head and neck cancer patients with nodal metastases (M/F: 16/3; age: 41-74 years old; primary tumor site: oropharynx) with histopathologically confirmed HNSCC were selected for this study. The characteristics of the patients are listed in Table 1. All patients were treated with intensity-modulated radiation therapy with dose prescriptions of 70 Gy for gross disease, 59.4 Gy for high-risk regions, and 50 to 54 Gy for low-risk regions[19,20]. Patients received cetuximab (400 mg/m2 loading dose, followed by 250 mg/m2 weekly), bevacizumab (15 mg/kg, days 1 and 22), and cisplatin (50 mg/m2, days 1, 2, 22, and 23). Locoregional control (LC) was determined by clinical and radiographic examination (MRI and/or positron emission/computed tomography [PET/CT]) using established criteria[21,22] with a median followup of 32 mo (range: 14.7-76.3). Patients with residual morphologic abnormalities on followup imaging were not classified as a locoregional failure (LF) unless recurrence was proven by biopsy or the abnormality progressed in size on serial imaging.

## DCE-MRI

Patients underwent examination with MRI before (pretreatment) and 10 to 14 d (intra-treatment) after the commencement of chemo-radiation treatment. MRI was performed on a 1.5-Tesla scanner (General Electric, Milwaukee, WI, United States) using an 8-channel neurovascular phased-array coil. The MRI protocol consisted of the standard anatomic MRI scans (T1/T2 weighted images) and DCE-MRI scans.

A two-dimensional spoiled gradient echo (2D-SPGR) pulse sequence was used for DCE-MRI image acquisition. The data acquisition parameters for the 2D-SPGR pulse sequence were: repetition time (TR) = 7.8 ms, echo time (TE) = 1.9 ms, temporal resolution = 6 s, phases = 50-60, number of excitation (NEX) = 1, flip angle (α) = 30°, receive bandwidth = 15.63-kHz, field of view (FOV) = 18-20 cm, slice thickness = 5-6 mm, yielding 3-8 slices, zero gap and a 256 × 128 matrix (zero-filled to 256 × 256 during image reconstruction).

The contrast of Gd-DTPA (Gadopentetic Diethylene Triamine Penta Acetic acid) (Magnevist; Berlex Laboratories, Wayne, NJ, United States) was delivered by antecubital vein catheters at a bolus of 0.1 mmol/kg and 2 cc/s, followed by saline flush using a MR-compatible, programmable power injector (Spectris; Medrad, Indianola, PA, United States). The total acquisition time for obtaining the DCE-MRI data was approximately 5 min.

***Pharmacokinetic modeling***

Pharmacokinetic modeling was performed using the Tofts model[23]:



[1]

where Ct (mM) is the concentration of contrast agent in the tumor tissue; Cp (mM) is the concentration of contrast agent in the blood plasma [known as the arterial input function (AIF)]; Ktrans (min-1) is the volume transfer rate between the blood plasma and extravascular extracellular space (EES); ve is the volume fraction of EES; T is the acquisition time course (min); and t is the integration variable.

Pharmacokinetic measures (Ktrans, ve) were estimated by using a nonlinear least-square fitting method, of which the optimization procedure consisted of using a Matlab build-in subspace trust region algorithm[24]. Three-dimensional regions of interest (ROIs) were delineated for the metastatic node on all the 2D-SPGR slices containing the tumor by a neuroradiologist with more than 10 years of experience, as described previously[25]. The data was fitted on a voxel-by-voxel basis within the ROI, yielding values of average (mean) and standard deviation (std), as well as parametric maps of Ktrans and ve.

***Image texture analysis***

Image texture analysis was performed on parametric maps of Ktrans and ve at the tumor’s central slice. First, the noise on the maps was removed with a noise thresholding method which is based on the Ostu’s algorithm. Histogram equalization with 64 discrete levels was performed to enhance map contrast. GLCM was calculated with 16 gray levels by setting the distance of pixel of interest to the left of its neighbor to be one pixel[11]. Based on the GLCM obtained, two texture measures of parametric maps were computed:

[2]



Energy (E) =

Where *p*(i, j) is the [i,j]th entry in a gray-tone spatial dependence matrix (I = 1:N; j = 1:N), and N is the number of distinct gray levels in the quantized image. E returns the sum of squared elements in the GLCM. E ranges from 0 to 1. For a constant image, E equals 1.

[3]



Homogeneity (H) =

H returns a value that measures the closeness of the distribution of elements in the GLCM to the GLCM diagonal. H ranges from 0 to 1. For a diagonal GLCM, H equals 1.

***Statistical analysis***

The measures derived from DCE-MRI pharmacokinetic modeling and image texture analysis were used for statistical analysis. The Lilliefors test was used to test the normality of all DCE-MRI-derived measures from the study patients. The differences in these measures between pretreatment and intra-treatment DCE-MRI scans were tested using a paired student’s *t-*test. The non-parametric Mann-Whitney U test was performed to assess the metric differences between LC and LF groups. The measures from pretreatment and intra-treatment scans, and the metric difference between pretreatment and intra-treatment scans, were used for the Mann-Whitney *U* test. To determine the measures that provide the best prediction of outcome, a forward sequential feature selection algorithm was used, followed by logistic regression analysis, which determined the probability of prediction. Statistical analysis was performed using Matlab R2008a on a Microsoft Windows system.

**RESULTS**

The clinical outcome for 19 patients with cancer of the oropharynx was assessed: 17 patients had local control of the disease (LC), and 2 patients had local failure (LF).

Figure 1 displays DCE-MRI images (pretreatment and intratreatment scans) and derived parametric maps (Ktrans and ve) for a representative patients with locoregional failure (male, 50 years). For comparing the difference of tumor volumes between pretreatment and intratreatment studies in 19 patients, there was a statistical significance (9.02 ± 6.87 cm3 *vs* 7.23 ± 6.22 cm3; *P* < 0.02, Table 2), showing tumor volumes were smaller after the commencement of treatment. No significant changes were found for the common summarizing measures (mean and standard deviation) for Ktrans and ve (*P* > 0.09). However, texture analysis revealed that the imaging biomarker energy (E) of ve was significantly higher in the intra-treatment scans than in the pretreatment scans [0.41 ± 0.22 *vs* 0.30 ± 0.11 (no unit); *P* < 0.04], as shown in Table 2 and Figure 2.

No significant difference was found when comparing the measures between the LC and LF groups, either in the pretreatment or intra-treatment scans, as seen in Table 3. However, there was a trend towards greater elevation in the energy (E) of ve in the LF patients after treatment, relative to the LC patients [0.37 ± 0.23 *vs* 0.07 ± 0.16 (no unit); *P* = 0.07].

DISCUSSION

In this study, we assessed the benefit of GLCM texture analysis on pharmacokinetic DCE-MRI maps for the prediction of treatment response in patients with HNSCC with neck nodal metastasis. An early and reliable predictor of treatment outcome could be especially valuable in the management of advanced HNSCC. Whereas common summarizing measures (mean and standard deviation) for Ktrans and ve did not yield significant differences, a biomarker derived from texture analysis, energy (E) of ve, did yield significant results. Specifically, we observed a significantly higher intra-treatment energy E of ve,compared to pretreatment scans.

The GLCM-derived biomarker, E, can be interpreted as an inverse heterogeneity measure, in which lower values are indicative of greater heterogeneity[11]. Hence, as the energy E of ve was higher during treatment, compared with before treatment, it seems that treatment reduces the heterogeneity of the tumor. However, the treatment-induced reduction in heterogeneity seems more pronounced for patients with local failure, rather than patients with local control, which might indicate that ineffective treatment counter-intuitively yields more homogeneous, rather than heterogeneous, tumor characteristics. It is important to note that the low number of patients with LF observed in our cohort limits the reliability of this analysis. The low number of LF patients in this study is because at our Center, patients with advanced locoregional cancer of the oropharynx who undergo chemo-radiation treatment have an approximately 90% local control rate. Future studies would require larger patient populations. If treatment-induced reduction of tumor heterogeneity in patients with local failure is validated in subsequent studies, this finding may have implications for the future design of adaptive chemo-radiation therapy trials in advanced head and neck cancers.

DCE-MRI provides a non-invasive method to probe the properties of tumor vasculature, such as perfusion, permeability of blood vessels, and volume of extracellular space. The ability of DCE-MRI to predict a patient’s response to chemo-radiation treatment has been previously investigated in head and neck cancers[5,7,8]. In a study by Cao *et al*[7] that followed head and neck cancer patients for a median of 10 months (range, 5-27 mo), the blood volume in the primary gross tumor volume was increased significantly in local control patients after 2 wk of chemo-radiation, compared with local failure patients (*P* < 0.03)[7]. In a study of 33 patients with head and neck cancer treated with neoadjuvant chemo-radiation therapy, Kim *et al*[8] found that the average pretreatment Ktrans value of a group with complete response to treatment was significantly higher than that of a group with partial response (*P* = 0.001) at a 6 mo follow-up. Our findings add to these studies that in addition to standard descriptive measures of pharmacokinetic maps, texture analysis may provide potential biomarkers related to local disease control, and thus may help guide clinical management.

The results of our study are not necessarily consistent with other studies that applied image texture analysis on DCE-MRI-derived parametric maps[11,26]. For example, Alic *et al*[11] used the heterogeneity features (i.e. coherence and fractal dimension) from texture analysis on parametric maps derived from DCE-MRI scans in 18 patients with limb sarcomas and found that tumors that responded to treatment had a high coherence on the pretreatment scans of those patients[11]. Similarly, Connor *et al*[26] found that microvascular uniformity (assessed with the fractal measure box dimension) in pretreatment scans correlated with treatment response in 10 patients with colorectal cancer with liver metastases. However, different tumor types and treatment regimens may impact the results of DCE-MRI-derived parametric maps in different ways. Validation studies with histopathology[27] or other measures of tumor micro-environment[10,25,28,29] open up scope for future studies in this area.

The limitations of this study are that: (1) the study had a small cohort of patients with only 2 patients with loco-regional failures, therefore an investigation with a large population is needed; (2) in this study, we made an assumption of an accurate T1 and AIF measurement and ignorable patient movements, which may affect the values of pharmacokinetic metrics and generated texture features; and (3) We did not compare the sensitivity and specificity of our method with the results from other investigators. All these issues will be addressed in future studies.

Image texture analysis can provide biomarker’s of heterogeneity for the analysis of DCE-MRI parametric maps of head and neck cancers. Specifically, we observed that chemo-radiation treatment significantly reduces the heterogeneity of tumors. The findings from this study may have potential value in stratifying patients and designing individualized treatment plans during the early management of HNSCC patients.

ACKNOWLEDGEMENTS

The authors would like to thank the MRI technologists for their great efforts to help perform all MRI examinations and Ms. Dara Srisaranard for her kind contribution to patient enrollment and data management. We thank Ms. Sandhya George for editing the manuscript.

**COMMENTS**

***Background***

Head and neck squamous cell carcinoma (HNSCC) is a major form of cancer that still kills vast numbers of cancer patients, and patients would certainly benefit with improved imaging methodology for a better staging/diagnosis, treatment evaluation, and treatment response prediction.

***Research frontiers***

Although magnetic resonance imaging (MRI) is often standardly applied in the clinical workup of patients newly diagnosed with HNSCC, advanced MRI methods [such as dynamic contrast-enhanced (DCE)-MRI] and advanced data analysis methods are currently being developed to optimize the added value of imaging.

***Innovations and breakthroughs***

In this study, novel texture analysis was applied to parametric maps derived from DCE-MRI. The results are very promising.

***Applications***

In principle, the applied image analysis can be readily applied on any parametric map, hence they can be incorporated in many clinical cancer trials that incorporate MRI imaging.

***Terminology***

AIF: Arterial input function; DCE: Dynamic contrast-enhanced; EES: Extravascular extracellular space; Gd-DTPA: Gadopentetic diethylene triamine penta acetic acid; GLCM: Gray-level co-occurrence matrix; HNSCC: Head and neck squamous cell carcinoma; LC: Locoregional control; LF: Locoregional failure; MRI: Magnetic resonance imaging; NEX: Number of excitation; PET: Positron emission tomography; ROI: Region of interest; SPGR: Spoiled gradient echo;TR: Repetition time; TE: Echo time.

***Peer-review***

This paper describes a study to assess the treatment response in patients with HNSCC using quantitative texture metrics of energy (E) and homogeneity (H). The authors observed that the heterogeneity of the tumors has been reduced after the treatment and the texture biomarkers can be used to evaluate the treatment response. Overall, the paper is well written and relative easy to follow.

# REFERENCES

1 **Hylton N**. Dynamic contrast-enhanced magnetic resonance imaging as an imaging biomarker. *J Clin Oncol* 2006; **24**: 3293-3298 [PMID: 16829653 DOI: 10.1200/JCO.2006.06.8080]

2 **Padhani AR**, Husband JE. Dynamic contrast-enhanced MRI studies in oncology with an emphasis on quantification, validation and human studies. *Clin Radiol* 2001; **56**: 607-620 [PMID: 11467863 DOI: 10.1053/crad.2001.0762]

3 **Padhani AR**, Khan AA. Diffusion-weighted (DW) and dynamic contrast-enhanced (DCE) magnetic resonance imaging (MRI) for monitoring anticancer therapy. *Target Oncol* 2010; **5**: 39-52 [PMID: 20383784 DOI: 10.1007/s11523-010-0135-8]

4 **Asaumi J**, Yanagi Y, Konouchi H, Hisatomi M, Matsuzaki H, Kishi K. Application of dynamic contrast-enhanced MRI to differentiate malignant lymphoma from squamous cell carcinoma in the head and neck. *Oral Oncol* 2004; **40**: 579-584 [PMID: 15063385 DOI: 10.1016/j.oraloncology.2003.12.002]

5 **Shukla-Dave A**, Lee NY, Jansen JF, Thaler HT, Stambuk HE, Fury MG, Patel SG, Moreira AL, Sherman E, Karimi S, Wang Y, Kraus D, Shah JP, Pfister DG, Koutcher JA. Dynamic contrast-enhanced magnetic resonance imaging as a predictor of outcome in head-and-neck squamous cell carcinoma patients with nodal metastases. *Int J Radiat Oncol Biol Phys* 2012; **82**: 1837-1844 [PMID: 21601373 DOI: 10.1016/j.ijrobp.2011.03.006]

6 **Chawla S**, Kim S, Dougherty L, Wang S, Loevner LA, Quon H, Poptani H. Pretreatment diffusion-weighted and dynamic contrast-enhanced MRI for prediction of local treatment response in squamous cell carcinomas of the head and neck. *AJR Am J Roentgenol* 2013; **200**: 35-43 [PMID: 23255739 DOI: 10.2214/AJR.12.9432]

7 **Cao Y**, Popovtzer A, Li D, Chepeha DB, Moyer JS, Prince ME, Worden F, Teknos T, Bradford C, Mukherji SK, Eisbruch A. Early prediction of outcome in advanced head-and-neck cancer based on tumor blood volume alterations during therapy: a prospective study. *Int J Radiat Oncol Biol Phys* 2008; **72**: 1287-1290 [PMID: 19028268 DOI: 10.1016/j.ijrobp.2008.08.024]

8 **Kim S**, Loevner LA, Quon H, Kilger A, Sherman E, Weinstein G, Chalian A, Poptani H. Prediction of response to chemoradiation therapy in squamous cell carcinomas of the head and neck using dynamic contrast-enhanced MR imaging. *AJNR Am J Neuroradiol* 2010; **31**: 262-268 [PMID: 19797785 DOI: 10.3174/ajnr.A1817]

9 **Jackson A**, O'Connor JP, Parker GJ, Jayson GC. Imaging tumor vascular heterogeneity and angiogenesis using dynamic contrast-enhanced magnetic resonance imaging. *Clin Cancer Res* 2007; **13**: 3449-3459 [PMID: 17575207 DOI: 10.1158/1078-0432.CCR-07-0238]

10 **Jansen JF**, Koutcher JA, Shukla-Dave A. Non-invasive imaging of angiogenesis in head and neck squamous cell carcinoma. *Angiogenesis* 2010; **13**: 149-160 [PMID: 20383743 DOI: 10.1007/s10456-010-9167-z]

11 **Alic L**, van Vliet M, van Dijke CF, Eggermont AM, Veenland JF, Niessen WJ. Heterogeneity in DCE-MRI parametric maps: a biomarker for treatment response? *Phys Med Biol* 2011; **56**: 1601-1616 [PMID: 21335648 DOI: 10.1088/0031-9155/56/6/006]

12 **El Naqa I**, Grigsby P, Apte A, Kidd E, Donnelly E, Khullar D, Chaudhari S, Yang D, Schmitt M, Laforest R, Thorstad W, Deasy JO. Exploring feature-based approaches in PET images for predicting cancer treatment outcomes. *Pattern Recognit* 2009; **42**: 1162-1171 [PMID: 20161266 DOI: 10.1016/j.patcog.2008.08.011]

13 **Alic L**, van Vliet M, Wielopolski PA, ten Hagen TL, van Dijke CF, Niessen WJ, Veenland JF. Regional heterogeneity changes in DCE-MRI as response to isolated limb perfusion in experimental soft-tissue sarcomas. *Contrast Media Mol Imaging* 2012; **8**: 340-349 [PMID: 23613437 DOI: 10.1002/cmmi.1528]

14 **Davnall F**, Yip CS, Ljungqvist G, Selmi M, Ng F, Sanghera B, Ganeshan B, Miles KA, Cook GJ, Goh V. Assessment of tumor heterogeneity: an emerging imaging tool for clinical practice? *Insights Imaging* 2012; **3**: 573-589 [PMID: 23093486 DOI: 10.1007/s13244-012-0196-6]

15 **Dominietto M**, Lehmann S, Keist R, Rudin M. Pattern analysis accounts for heterogeneity observed in MRI studies of tumor angiogenesis. *Magn Reson Med* 2013; **70**: 1481-1490 [PMID: 23280475 DOI: 10.1002/mrm.24590]

16 **Just N**. Improving tumour heterogeneity MRI assessment with histograms. *Br J Cancer* 2014; **111**: 2205-2213 [PMID: 25268373 DOI: 10.1038/bjc.2014.512]

17 **Karahaliou A**, Vassiou K, Arikidis NS, Skiadopoulos S, Kanavou T, Costaridou L. Assessing heterogeneity of lesion enhancement kinetics in dynamic contrast-enhanced MRI for breast cancer diagnosis. *Br J Radiol* 2010; **83**: 296-309 [PMID: 20335440 DOI: 10.1259/bjr/50743919]

18 **Yang X**, Knopp MV. Quantifying tumor vascular heterogeneity with dynamic contrast-enhanced magnetic resonance imaging: a review. *J Biomed Biotechnol* 2011; **2011**: 732848 [PMID: 21541193 DOI: 10.1155/2011/732848]

19 **Setton J**, Caria N, Romanyshyn J, Koutcher L, Wolden SL, Zelefsky MJ, Rowan N, Sherman EJ, Fury MG, Pfister DG, Wong RJ, Shah JP, Kraus DH, Shi W, Zhang Z, Schupak KD, Gelblum DY, Rao SD, Lee NY. Intensity-modulated radiotherapy in the treatment of oropharyngeal cancer: an update of the Memorial Sloan-Kettering Cancer Center experience. *Int J Radiat Oncol Biol Phys* 2012; **82**: 291-298 [PMID: 21167652 DOI: 10.1016/j.ijrobp.2010.10.041]

20 **Lee N**, Harris J, Garden AS, Straube W, Glisson B, Xia P, Bosch W, Morrison WH, Quivey J, Thorstad W, Jones C, Ang KK. Intensity-modulated radiation therapy with or without chemotherapy for nasopharyngeal carcinoma: radiation therapy oncology group phase II trial 0225. *J Clin Oncol* 2009; **27**: 3684-3690 [PMID: 19564532]

21 **Goenka A**, Morris LG, Rao SS, Wolden SL, Wong RJ, Kraus DH, Ohri N, Setton J, Lok BH, Riaz N, Mychalczak BR, Schoder H, Ganly I, Shah JP, Pfister DG, Zelefsky MJ, Lee NY. Long-term regional control in the observed neck following definitive chemoradiation for node-positive oropharyngeal squamous cell cancer. *Int J Cancer* 2013; **133**: 1214-1221 [PMID: 23436584 DOI: 10.1002/ijc.28120]

22 **Mutter RW**, Lok BH, Dutta PR, Riaz N, Setton J, Berry SL, Goenka A, Zhang Z, Rao SS, Wolden SL, Lee NY. Constraining the brachial plexus does not compromise regional control in oropharyngeal carcinoma. *Radiat Oncol* 2013; **8**: 173 [PMID: 23835205 DOI: 10.1186/1748-717X-8-173]

23 **Tofts PS**, Brix G, Buckley DL, Evelhoch JL, Henderson E, Knopp MV, Larsson HB, Lee TY, Mayr NA, Parker GJ, Port RE, Taylor J, Weisskoff RM. Estimating kinetic parameters from dynamic contrast-enhanced T(1)-weighted MRI of a diffusable tracer: standardized quantities and symbols. *J Magn Reson Imaging* 1999; **10**: 223-232 [PMID: 10508281 DOI: 10.1002/(SICI)1522-2586(199909)10: 3<223: : AID-JMRI2>3.0.CO; 2-S]

24 **More JJ**, Sorensen DC. Computing a Trust Region Step. *SIAM J Sci Statist Comput* 1983; **4**: 553-572

25 **Jansen JF**, Schöder H, Lee NY, Stambuk HE, Wang Y, Fury MG, Patel SG, Pfister DG, Shah JP, Koutcher JA, Shukla-Dave A. Tumor metabolism and perfusion in head and neck squamous cell carcinoma: pretreatment multimodality imaging with 1H magnetic resonance spectroscopy, dynamic contrast-enhanced MRI, and [18F]FDG-PET. *Int J Radiat Oncol Biol Phys* 2012; **82**: 299-307 [PMID: 21236594 DOI: 10.1016/j.ijrobp.2010.11.022]

26 **O'Connor JP**, Rose CJ, Jackson A, Watson Y, Cheung S, Maders F, Whitcher BJ, Roberts C, Buonaccorsi GA, Thompson G, Clamp AR, Jayson GC, Parker GJ. DCE-MRI biomarkers of tumour heterogeneity predict CRC liver metastasis shrinkage following bevacizumab and FOLFOX-6. *Br J Cancer* 2011; **105**: 139-145 [PMID: 21673686 DOI: 10.1038/bjc.2011.191]

27 **Jansen JF**, Carlson DL, Lu Y, Stambuk HE, Moreira AL, Singh B, Patel SG, Kraus DH, Wong RJ, Shaha AR, Shah JP, Shukla-Dave A. Correlation of a priori DCE-MRI and (1)H-MRS data with molecular markers in neck nodal metastases: Initial analysis. *Oral Oncol* 2012; **48**: 717-722 [PMID: 22366441 DOI: 10.1016/j.oraloncology.2012.02.001]

28 **Jansen JF**, Schöder H, Lee NY, Wang Y, Pfister DG, Fury MG, Stambuk HE, Humm JL, Koutcher JA, Shukla-Dave A. Noninvasive assessment of tumor microenvironment using dynamic contrast-enhanced magnetic resonance imaging and 18F-fluoromisonidazole positron emission tomography imaging in neck nodal metastases. *Int J Radiat Oncol Biol Phys* 2010; **77**: 1403-1410 [PMID: 19906496 DOI: 10.1016/j.ijrobp.2009.07.009]

29 **Jansen JF**, Parra C, Lu Y, Shukla-Dave A. Evaluation of Head and Neck Tumors with Functional MR Imaging. *Magn Reson Imaging Clin N Am* 2015 [DOI: 10.1016/j.mric.2015.08.011]

**P-Reviewer:** Xian J, Zhang T **S-Editor:** Ji FF **L-Editor: E-Editor:**

**Table 1 Patient characteristics**

|  |  |
| --- | --- |
| Characteristics | Value |
| Total patients | 19 |
| **Demographics** |  |
| Mean age (yr) | 57 |
| Age range (yr) | 41-74 |
| Male/female | 16/3 |
| **Location of primary tumor** |  |
| Oropharynx | 19 |
| **Stages** |  |
| Stage III | 15 |
| Stage IV | 4 |
| **Tumor size (cm3)** | 0.615-14.879 |
| **Therapy type**  *Outcome*  Locoregional control  Locoregional failure | Chemo-radiation  17  2 |

**Table 2 Paired student’s *t* test on tumor volume, Ktrans and ve between pretreatment and intra-treatment scans (*n* = 19)**

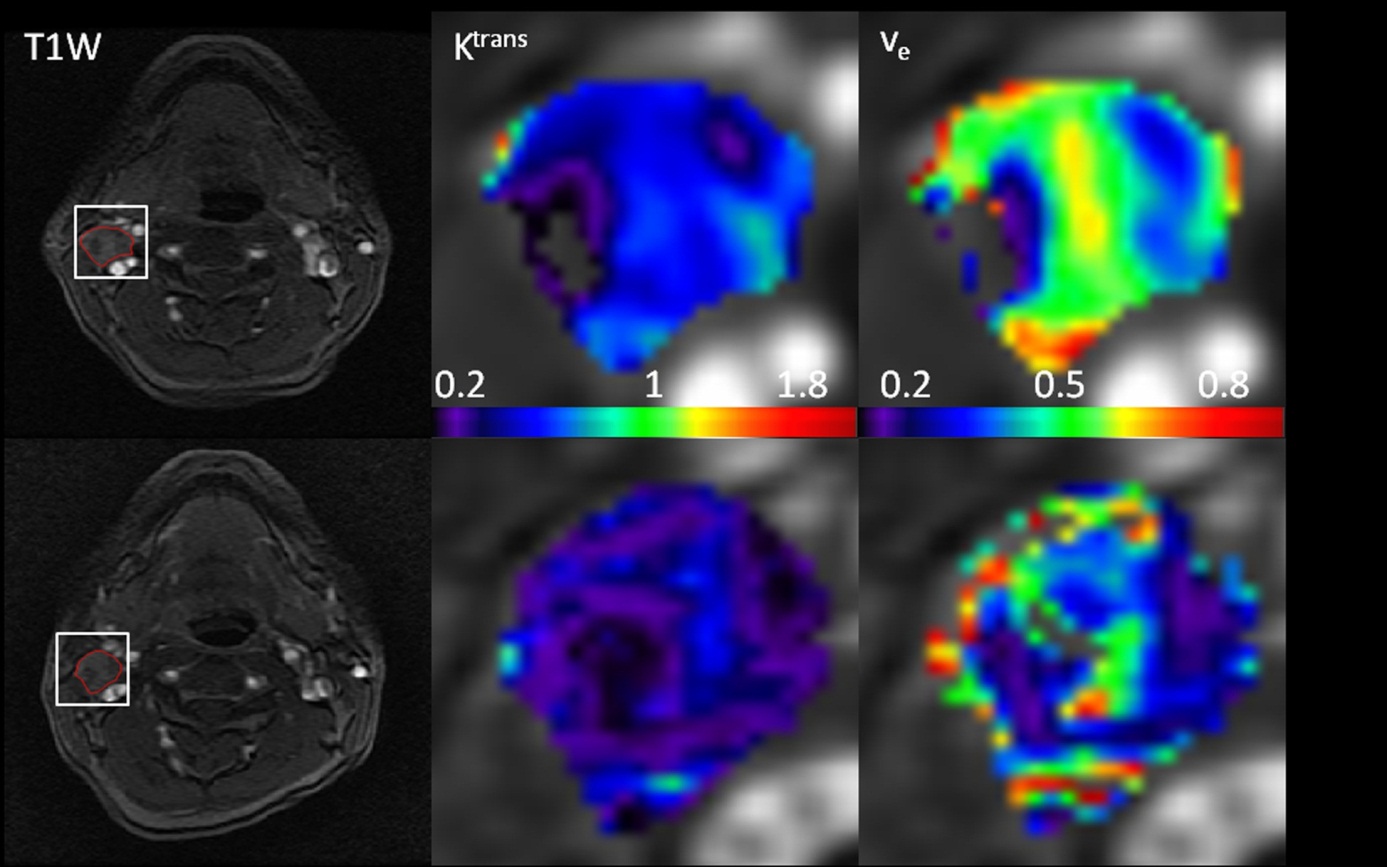
|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| Parameters | Measures | Pretreatment  (*n* = 19)  (mean ± SD) | Intra-treatment  (*n* = 19)  (mean ± SD) | *P* values |
| Tumor volume (cm3) |  | 9.02 ± 6.87 | 7.23 ± 6.22 | 0.02a |
| Ktrans  (volume transfer rate, min-1) | Mean | 0.34 ± 0.18 | 0.37 ± 0.22 | 0.36 |
| SD | 0.25 ± 0.11 | 0.25 ± 0.11 | 0.99 |
| E | 0.34 ± 0.14 | 0.38 ± 0.19 | 0.18 |
| H | 0.84 ± 0.04 | 0.83 ± 0.05 | 0.61 |
| ve  (volume fraction of the extravascular extracellular space) | mean | 0.29 ± 0.12 | 0.34 ± 0.19 | 0.09 |
| SD | 0.15 ± 0.04 | 0.17 ± 0.07 | 0.26 |
| E | 0.30 ± 0.11 | 0.41 ± 0.22 | 0.04a |
| H | 0.84 ± 0.04 | 0.83 ± 0.06 | 0.78 |

a*P* < 0.05 of comparison pre *vs* intra-treatment. SD: Standard deviation; E: Energy; H: Homogeneity.

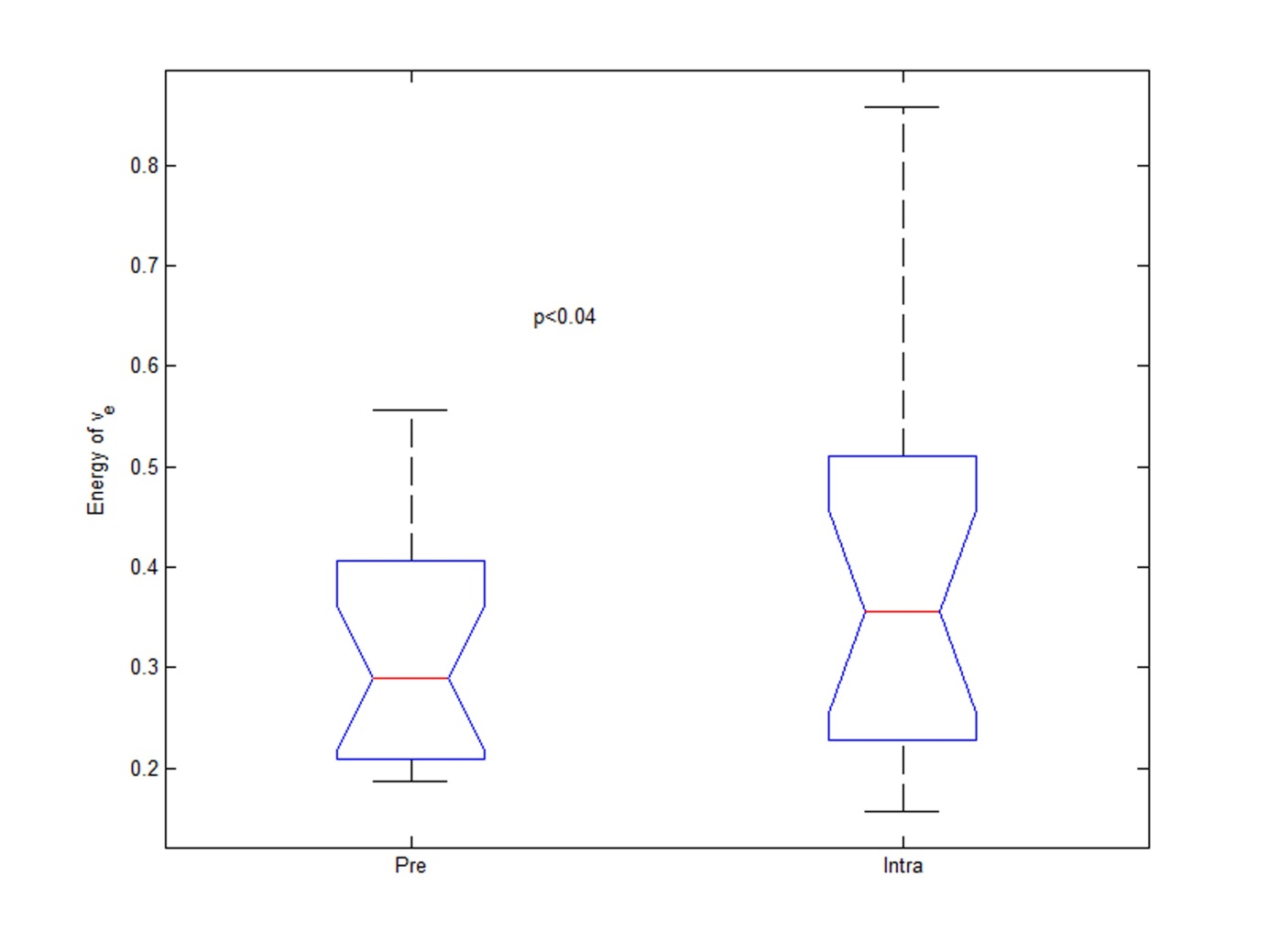
**Table 3 Mann-Whitney *U* test of tumor volume, Ktrans and ve between groups of patients with locoregional control (*n* = 17) and locoregional failure (*n* = 2)**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| Measures for patient groups (LC and LF) | | | Pretreatment | Intra-treatment | Difference between Intra-treatment and pretreatment |
| Tumor volume  (cm3) | | LC | 9.68 ± 7.03 | 7.70± 6.45 | -1.97 ± 3.13 |
| LF | 3.79 ± 0.73 | 3.42 ± 0.91 | -0.37 ± 0.17 |
| *P* value | 0.15 | 0.54 | 0.64 |
| Ktrans  (min-1) | Mean | LC | 0.34 ± 0.19 | 0.38 ± 0.23 | 0.04 ± 0.15 |
| LF | 0.34 ± 0.01 | 0.26 ± 0.13 | -0.08 ± 0.14 |
| *P* value | *P* = 0.84 | *P* = 0.42 | *P* = 0.42 |
| Std | LC | 0.26 ± 0.11 | 0.25 ± 0.11 | -0.007 ± 0.12 |
| LF | 0.20 ± 0.05 | 0.27 ± 0.16 | -0.08 ± 0.14 |
| *P* value | *P* = 0.49 | *P* = 0.74 | *P* = 0.49 |
| E | LC | 0.35 ± 0.15 | 0.38 ± 0.19 | 0.02 ± 0.11 |
| LF | 0.27 ± 0.08 | 0.41 ± 0.30 | 0.14 ± 0.22 |
| *P* value | *P* = 0.74 | *P* = 1.00 | *P* = 0.57 |
| H | LC | 0.84 ± 0.04 | 0.83 ± 0.06 | -0.01 ± 0.04 |
| LF | 0.82 ± 0.03 | 0.85 ± 0.04 | 0.03 ± 0.04 |
| *P* value | *P* = 0.35 | *P* = 0.57 | *P* = 0.35 |
| ve | Mean | LC | 0.29 ± 0.13 | 0.35 ± 0.20 | 0.05 ± 0.12 |
| LF | 0.28 ± 0.10 | 0.26 ± 0.01 | -0.01 ± 0.11 |
| *P* value | *P* = 0.94 | *P* = 0.42 | *P* = 0.35 |
| Std | LC | 0.16 ± 0.04 | 0.17 ± 0.07 | 0.01 ± 0.07 |
| LF | 0.14 ± 0.01 | 0.22 ± 0.01 | 0.08 ± 0.02 |
| *P* value | *P* = 0.23 | *P* = 0.14 | *P* = 0.10 |
| E | LC | 0.31 ± 0.12 | 0.38 ± 0.21 | 0.07 ± 0.16 |
| LF | 0.28 ± 0.02 | 0.65 ± 0.21 | 0.37 ± 0.23 |
| *P* value | *P* = 0.94 | *P* = 0.10 | *P* = 0.07 |
| H | LC | 0.84 ± 0.04 | 0.82 ± 0.06 | -0.01 ± 0.06 |
| LF | 0.84 ± 0.02 | 0.91 ± 0.06 | 0.06 ± 0.06 |
| *P* value | *P* = 0.74 | *P* = 0.14 | *P* = 0.18 |

LC: Locoregional control; LF: Locoregional failure.

****

**Figure 1 Pretreatment and intra-treatment dynamic contrast-enhanced magnetic resonance imaging images of patient with locoregional failure (male, 50 years).** The top row shows images of the pretreatment study, and the bottom row shows images from the intra-treatment study. From left to right, the columns show a T1-weighted image, Ktrans (min-1) map, and ve map. The white rectangles delineate the regions of interest (ROIs) at the metastatic nodes. Ktrans and ve maps are zoomed at the locations of ROIs. The corresponding texture measures for the Ktrans and ve maps for pre and intra-treatment are: Ktrans:Energy (pre: 0.25, intra: 0.38), Homogeneity (pre: 0.81, intra: 0.83), ve: Energy (pre: 0.27, intra: 0.70), Homogeneity (pre: 0.83, intra: 0.90).

****

**Figure 2 Box-and-whisker plot demonstrating the significant difference in the energy of ve between pretreatment and intra-treatmentscans (*P* < 0.04).** Pre: Pretreatment; Intra: Intra-treatment.