

Texture analysis of early cerebral tissue damage in magnetic resonance imaging of patients with lung cancer

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Abstract. Primary tumors can secrete many cytokines, inducing tissue damage or microstructural changes in distant organs. The purpose of this study was to investigate changes in texture features in the cerebral tissue of patients with lung cancer without brain metastasis. In this study, 50 patients with lung cancers underwent 3.0-T magnetic resonance imaging (MRI) within 2 weeks of being diagnosed with lung cancer. Texture analysis (TA) was carried out in 8 gray matter areas, including bilateral frontal cortices, parietal cortices, occipital cortices and temporal cortices, as well as 2 areas of bilateral frontoparietal white matter. The same procedure was performed for 57 healthy controls. A total of 32 texture parameters were separately compared between the patients and controls in the different cerebral tissue sites. Texture features among patients based on histological type and clinical stage were also compared. Of the 32 texture parameters, 27 showed significant differences between patients with lung cancer and healthy controls. There were significant differences in cerebral tissue, both gray matter and white matter between patients and controls, especially in several wavelet-based parameters. However, there were no significant differences between tissue at homologous sites in bilateral hemispheres, either in patients or controls. TA detected overt changes in the texture features of cerebral tissue in patients with lung cancer without brain

metastasis compared with those of healthy controls. TA may be considered as a novel and adjunctive approach to conventional brain MRI to reveal cerebral tissue changes invisible on MRI alone in patients with lung cancer.

Introduction

According to the 2018 Global Cancer Statistics, lung cancer is the leading cause of cancer-associated mortality worldwide (1) and was so in China in 2014 (2). However, the cure for lung cancer remains elusive due to the aggressive and disseminative nature of tumors which restricts the efficacy of cancer treatments (3). The brain is a preferential site of metastasis in lung cancer (4). Among patients with lung cancer, ~30-50% develop brain metastasis (5-7), which substantially affects their quality of life. However, in the present study, even in the absence of brain metastasis, some individuals continued to exhibit paraneoplastic neurological syndromes, such as behavioral, memory and movement disorders as well as hypomnesia, dizziness and headaches (8). A number of previous studies have hypothesized that the pathophysiology underlying paraneoplastic syndromes is associated with the secretion of cytokines by the primary tumor (9-12). This may induce injury to distant organs and cause microstructural changes like those in the brain, which cause paraneoplastic syndromes or alter the homing and proliferation of metastatic cancer cells (9-13). However, the underlying mechanisms remain unclear.

Due to the potential ability to assess the characteristics of brain tissue noninvasively, magnetic resonance imaging (MRI) is currently the main method used for diagnosis, treatment guidance and monitoring of brain metastasis (14,15). However, identifying early changes in the cerebral tissue of patients with lung cancer without brain metastasis is difficult (16). Fortunately, MRI images of cerebral tissues contain a wealth of information invisible to the human eye that can be accessed by texture analysis (TA) techniques (17). Texture is a visual stimulus generated by repetitive image patterns, which can be depicted as regular or irregular, smooth or rough and fine or coarse. Although they can exhibit very complex patterns, some textures can be extracted relatively easily even by visual assessment due to the regular visual appearance of the patterns. However, most textures display random patterns because textural primitives are random, making them difficult

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to recognize and interpret. In medical images, these types of random textures are more common than regular patterns. As texture is an image feature defined by both pixel locations and brightness value, TA enables mathematical calculation of the patterns. Therefore, texture features provide the potential to distinguish and characterize properties of cerebral tissue (17).

The aim of the present study was to use texture analysis (TA) methods to describe early cerebral tissue damage caused by lung cancer without brain metastasis.

Materials and methods

Subjects. The present study was approved by the Ethics Committee of Shandong Provincial Hospital Affiliated to Shandong University (Jinan, China). Written informed consent was obtained from all patients prior to the study start. Patients with lung cancer were recruited from the Shandong Provincial Hospital between December 2017 and October 2018. The TA study included 50 consecutive patients (36 males, 14 females; mean age, 60 years; age range 47-75 years) who were all diagnosed by pathology and met the diagnostic criteria for primary lung cancer according to the World Health Organization (18). In addition to a detailed review of pathological specimens by pathologists upon admission to the hospital, all patients with lung cancer underwent a careful physical examination performed by experienced oncologists and neurologists from Shandong Provincial Hospital Affiliated to Shandong University, in order to determine the clinical stage, Karnofsky Performance Scale (KPS) score (19) and numeric rating scale score (NRS) (20). The demographic information and clinical characteristics of the patients are presented in Table I. The inclusion criteria were as follows: i) Age, 18-75 years; ii) histological type of small cell carcinoma, squamous cell carcinoma, adenocarcinoma and others; iii) KPS score >60 and iv) no neurological abnormalities. The exclusion criteria were as follows: i) Age, <18 years or >75 years; ii) systemic illness such as systemic lupus erythematosus, serious vascular disease, head trauma, epilepsy (healthy controls) or claustrophobia; iii) long-term use of psychoactive medication and iv) refusal to undergo MRI. The control group (CG) comprised of 57 age- and sex-matched healthy volunteers. The individuals in this group reported no types of malignant tumor or any systemic, neurological or psychiatric illnesses known to affect cerebral structure or function such as a long history of psychoactive medication use, head trauma, serious vascular disease, current depression, epilepsy or alcoholism.

MRI examinations. Data were acquired within 2 weeks of diagnosis of lung cancer primarily due to some patients being diagnosed in other hospitals and then transferred to Shandong Provincial Hospital for further antitumor treatment. Following diagnosis, all patients underwent conventional MRI using a 3.0-T scanner (Philips Medical Systems B.V.) equipped with an 8-channel head and neck coil. The MRI machine was included in a detailed quality control plan requiring daily, monthly and quarterly inspections. The radio frequency amplifier properties and main magnetic field homogeneity of the MRI machine were measured and controlled quarterly. The following conventional MRI sequences were employed: Sagittal, coronal and axial T2-weighted imaging (T2WI), T1-weighted imaging

Table I. Clinical characteristics of the studied groups.

Characteristics	Patient, n=50	Control, n=57
Age in years, mean \pm standard deviation	60.0 \pm 7.2	59.1 \pm 6.5
Age range, years	47-75	48-75
Gender, Female vs. Male	36/14	37/20
Histological type, n		
Adenocarcinoma	29	-
Squamous	17	-
Small cell	4	-
Clinical stage		
Early	18	-
Advanced	32	-
Karnofsky performance scale		
≥ 70	49	-
<70	1	-
Numeric rating scales		
≤ 3	37	-
>3	13	-

(T1WI), fluid-attenuated inversion recovery imaging, diffusion-weighted imaging, apparent diffusion coefficient and enhanced T1WI. The sequences employed in the MRI protocol are presented in Table II.

MR image segmentation. Image selection was carried out using a DICOM viewer (version 3.0; Philips Medical Systems B.V.). For each MR image, regions of interest (ROIs) were drawn symmetrically by hand on bilateral sides of the hemispheres at each level of interest (21,22). In the present study, 16x16-pixel ROIs were manually drawn on the frontoparietal white matter in the centrum semiovale (Fig. 1A), frontal cortices and parietal cortices at the level of the body of the lateral ventricles (Fig. 1B), occipital cortices at the level of the basal ganglia (Fig. 1C) and temporal cortices at the level of the midbrain (Fig. 1D). The ROIs were carefully placed by two experienced neuroradiologists from Shandong Provincial Hospital Affiliated to Shandong University, to avoid overlap with any areas of microhemorrhage, hyperintensity or macroscopic hemosiderin depositions which may occur in some patients. Fig. 2 presents an array of texture blocks extracted from the cerebral tissues of a patient with lung cancer, without metastasis and a healthy volunteer.

TA analysis technique. The axial T2WI sequence was selected from the whole MRI study for TA as T2-weighted MR imaging has been demonstrated to be sensitive to tissue abnormalities in the human brain (23). Only single slices were evaluated to obtain a maximum in-plane resolution by avoiding the lower resolution or interpolation, typically caused by 3-dimensional TA (24). Further analysis was performed using 4 MRI image slices selected from the T2WI sequence at 4 interest levels. The texture features in 10 ROIs were assessed based on the histogram, gray-level co-occurrence matrix (GLCM),

Table II. MRI sequences included in the magnetic resonance protocol of the present study.

Sequence	TR	TE	TI	Slice/gap	Matrix	FOV	Flip angle
Axial T2WI	4,000	100	0	6.0/1.0	384x384	230	90
Axial T1WI-IR	3,000	43	1,150	6.0/1.0	264x264	230	-
Axial FLAIR	9,000	148	2,500	6.0/1.0	284x284	230	-
DWI	2,257	71	0	6.0/1.0	152x152	230	90
Axial T1W-STIR	3,000	43	1,150	6.0/1.0	264x264	230	-

TR, repetition time; TE, echo time; TI, inversion time; FOV, field of view; T1WI, T1-weighted imaging; T2WI, T2-weighted imaging; DWI, diffusion-weighted imaging; FLAIR, fluid-attenuated inversion recovery; IR, inverse recovery; STIR, short time inversion recovery.

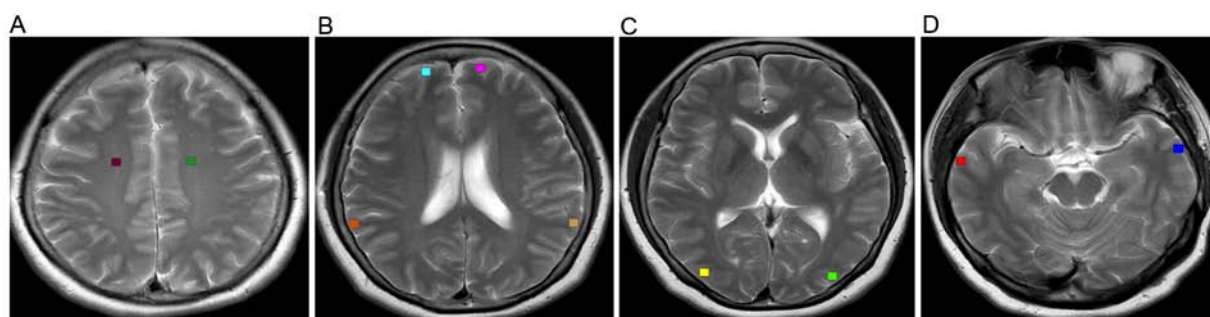


Figure 1. ROIs delineated at selected levels. (A) ROIs placed in frontoparietal white matter in the centrum semiovale. (B) ROIs placed in frontal and parietal cortices at the level of the body of the lateral ventricles. (C) ROIs placed in occipital cortices at the level of the basal ganglia. (D) ROIs placed in the temporal cortices at the level of midbrain. ROIs, regions of interest.

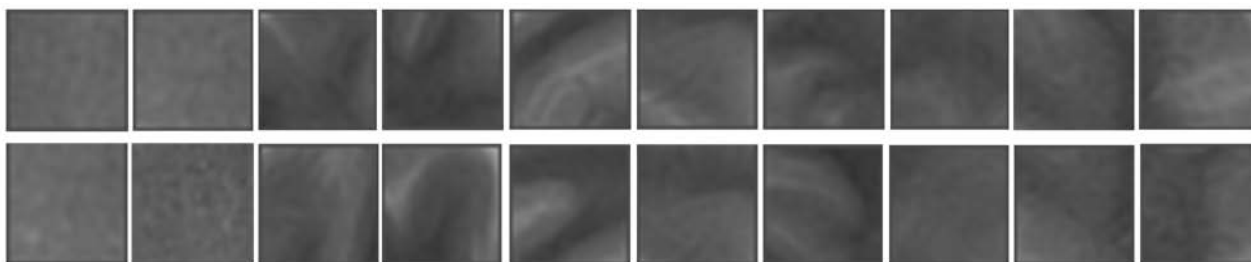


Figure 2. Examples of textures. Top row, the brain of a patient with lung cancer, without metastasis; bottom row, a healthy brain. Each block contains 16x16 pixels.

gray-level run length matrix (GLRLM) and wavelet transform, as summarized in Table III (17,25,26). The wavelet transform of the three dimensional image was the decomposition of the image along the x, y and z directions using low-pass filter (L) and high-pass filter (H), and each layer of wavelet decomposition decomposed the image data into eight different frequency bands (LLH, HLL, HLH, HHL, LHL, LHH, HHH and LLL), which included seven high frequency bands (LLH-HHH) and one low frequency band (LLL).

The first order statistics image properties derived from the gray-scale frequency histogram depend solely on individual pixel values. Second order statistics are properties of pixel pairs and describe the relationship between pixels and their gray levels. However, the first and second order statistics of images in texture analysis only contained the amplitude information of the image, ignoring the phase information, thus the wavelet transform of the image was also analyzed. TA was performed using the Python language (version 3.6)

on the JetBrains PyCharm platform (version 2018.1.3). Feature calculations were performed within the Pyradiomics package (version 2.2.0), which is an open-source Python package designed for the extraction of radiomic features from medical images (27). As the Fisher criterion can produce an array of features with highly discriminative power, a feature selection method was applied using the Fisher coefficient provided by the Python package, in order to identify the six texture features with the highest discriminatory potential for classification (17).

Statistical analysis. In total, 32 texture parameters were tested to determine which and how many features in each ROI demonstrated significant differences between patients with lung cancer, without brain metastasis and healthy controls. Statistical analyses were performed for each texture feature. Differences in texture features between the two groups were analyzed via the Wilcoxon rank sum test or Mann-Whitney U test, using SPSS software (version 22; IBM Corp.). The Fisher

Table III. List of texture features.

Histogram	Gray-level co-occurrence matrix	Gray-level run length matrix	Wavelet
Mean	Joint energy	Long run emphasis	HHH_kurtosis
Skewness	Inverse difference	Run length non-uniformity	HHL_maximum
Deviation	Correlation	Low gray-level run emphasis	HLH_skewness
Variance	Difference average	Short run low gray-level emphasis	LLL_10 percentile
Kurtosis	Difference entropy	Long run low gray-level emphasis	LLL_median
	Inverse difference normalized	Short run high gray-level emphasis	LLL_minimum
	Joint entropy	Long run high gray-level emphasis	LLL_variance
	Sum average	Short run emphasis	
	Sum entropy	Run gray-level non-uniformity	
		Run percentage	
		High gray-level run emphasis	

LLL, low frequency-low frequency-low frequency output; HHH, high frequency-high frequency-high frequency output; HHL, high frequency-high frequency-low frequency output; HLH, high frequency-low frequency-high frequency output.

coefficient was used to select the six most discriminative parameters. $P < 0.05$ was considered to indicate a statistically significant difference.

Results

Analysis workflow. Given brain MRI data sets of patients with lung cancer without brain metastasis and healthy controls, four interest levels were selected, and then texture blocks of 16x16 pixels were manually placed in a symmetrical manner on bilateral sides of the hemispheres at each level of interest. Finally, texture features were calculated based on the image histogram, the co-occurrence matrix, the run-length matrix and wavelets. The system diagram presented in Fig. 3 represents the process followed for MRI analysis.

Brain MRI of patients with lung cancer and healthy controls. In this study, conventional MRI enabled detection of lesions related to brain metastasis or primary brain tumors only when gross structural abnormalities could be visually detected. Based on the conventional MR images, no significantly different signals were found in any sequence in patients with lung cancer without metastasis compared with healthy volunteers (Table II). Fig. 4 shows conventional MR images of a 45-year-old female patient who presented with lung cancer without brain metastasis (top row) and a 54-year-old male healthy control (bottom row).

Comparison of texture parameters in bilateral hemispheres between patients with lung cancer without brain metastasis and healthy controls. For the comparison of bilateral hemispheres of 10 ROIs, 32 features were tested for each texture block to determine which and how many of the parameters differed statistically between patients with primary lung cancer without brain metastasis and healthy controls. The number of texture parameters ($n=32$) which were significantly different were analyzed using the Wilcoxon test (Table IV). The feature vector included mean and variance features from histogram statistics, energy and correlation features from a co-occurrence matrix,

short-run and long-run emphasis features from a run-length matrix and phase information features from wavelet transform. The 32 texture parameters included; Five histogram-based parameters, nine GLCM-based parameters, 11 GLRLM-based parameters and seven wavelet-based parameters. Among the five histogram-based parameters, which depended solely on individual pixel gray-level values, the following four parameters; mean, skewness, deviation and variance were significantly different ($P < 0.05$), and only the kurtosis parameter demonstrated no significant difference ($P = 0.686$). Among the nine GLCM-based parameters, seven parameters were significantly different; however, the parameters for correlation ($P = 0.435$) and normalized inverse difference ($P = 0.156$) demonstrated no significant differences. All 11 GLRLM-based parameters were significantly different ($P < 0.05$). Among the seven wavelet-based parameters, five parameters were significantly different ($P < 0.05$), whereas wavelet-HHH_kurtosis and wavelet-HLH_skewness failed to demonstrate statistically significant differences ($P > 0.05$, respectively) (Table IV). There were 27 texture parameters with a statistically significant difference including 4 histogram-based parameters, 7 GLCM-based parameters, 11 GLRLM-based parameters and 5 wavelet-based parameters. The feature parameters with statistically significant differences were in first order features, second order features and wavelets. For instance, mean indicated the average density among the texture blocks; deviation calculated the statistical variability of density among the pixels in the blocks; sum average indicated the total number of correlated pixel pairs with the same sum in the block; gray-level non-uniformity referred to the orderliness or randomness of the pixel densities among the blocks which could denote the degree of structure, and gray-level run length emphasis measured consecutive pixels of the same density along particular orientations as another representation of structure.

Identification of the six most discriminative parameters between patients with lung cancer without brain metastasis and healthy controls. When the texture parameters of 10 ROIs were compared in bilateral hemispheres between the lung cancer group and the healthy control group, 27 of the

Table IV. Comparison of texture features between patients with lung cancer and healthy controls.

Feature	Patient, mean (SD)	Control, mean (SD)	P-value
Kurtosis	3.649 (1.88)	3.707 (2.57)	0.686
Mean	544.176 (167.18)	688.647 (209.26)	<0.001
Skewness	0.614 (0.61)	0.465 (0.67)	0.001
Deviation	64.877 (42.55)	78.146 (47.07)	<0.001
Variance	6,016.221 (460.30)	8,319.324 (434.89)	<0.001
Correlation	0.806 (0.14)	0.800 (0.15)	0.435
Difference average	0.873 (0.43)	1.063 (0.51)	<0.001
Difference entropy	1.584 (0.41)	1.749 (0.45)	<0.001
Inverse difference	0.674 (0.09)	0.631 (0.09)	<0.001
Inverse difference normalized	0.941 (0.01)	0.939 (0.01)	0.156
Joint energy	0.073 (0.06)	0.057 (0.06)	<0.001
Joint entropy	4.772 (1.16)	5.193 (1.22)	<0.001
Sum average	11.151 (5.34)	14.097 (8.38)	<0.001
Sum entropy	3.791 (0.85)	4.044 (0.90)	<0.001
Gray-level non-uniformity	22.665 (9.16)	21.697 (10.55)	<0.001
High gray-level run emphasis	52.799 (65.99)	86.091 (127.81)	<0.001
Long run emphasis	4.592 (2.71)	3.605 (1.98)	<0.001
Long run high gray-level emphasis	135.634 (99.71)	195.944 (295.67)	<0.001
Long run low gray-level emphasis	0.640 (0.94)	0.403 (0.59)	<0.001
Low gray-level run emphasis	0.108 (0.08)	0.086 (0.07)	<0.001
Run length non-uniformity	80.196 (33.18)	96.709 (34.77)	<0.001
Run percentage	0.618 (0.11)	0.673 (0.10)	<0.001
Short run emphasis	0.694 (0.10)	0.741 (0.09)	<0.001
Short run high gray-level emphasis	43.297 (60.87)	71.963 (107.37)	<0.001
Short run low gray-level emphasis	0.067 (0.04)	0.058 (0.04)	<0.001
wavelet-HHH_kurtosis	8.636 (7.28)	8.702 (7.19)	0.932
wavelet-HHL_maximum	35.940 (30.20)	45.996 (40.29)	<0.001
wavelet-HLH_skewness	-0.224 (1.21)	-0.263 (1.34)	0.930
wavelet-LLL_10 percentile	1,358.451 (402.93)	1,689.975 (524.57)	<0.001
wavelet-LLL_median	1,544.083 (19.83)	1,927.296 (585.72)	<0.001
wavelet-LLL_minimum	1,257.445 (375.73)	1,538.135 (509.09)	<0.001
wavelet-LLL_variance	46,637.723 (84568.84)	64,229.389 (80765.98)	<0.001

A total of 32 texture parameters were evaluated. The parameters that demonstrated statistically significant differences between patients with lung cancer and health controls are presented in bold. $P < 0.05$ was considered to indicate a statistically significant difference. LLL, low frequency-low frequency-low frequency output; HHH, high frequency-high frequency-high frequency output; HHL, high frequency-high frequency-low frequency output; HLH, high frequency-low frequency-high frequency output.

32 were significantly different. Among the 27 significantly different parameters, six texture features were identified using the Fisher coefficient provided by the Python package. The Fisher criterion could produce a set of features with a high discriminatory potential for separation and classification which were also highly associated with each other. Based on the Fisher coefficient and the values obtained by the Wilcoxon test, the six most discriminative parameters for differences between patients and healthy controls were selected. These six texture parameters were the wavelet-LLL_median, mean, wavelet-LLL_10 percentile, wavelet-LLL_minimum, run percentage, and run length non-uniformity. The six most discriminative texture features were extracted from the first-order statistics, second-order statistics and wavelets.

However, the six texture features were mainly based on the wavelets and GLRLM features (Fig. 5).

Comparison of the top six texture parameters between the left and right hemispheres in patients with lung cancer without brain metastasis and healthy controls. The comparison of bilateral hemispheres of patients with lung cancer and healthy controls, proved that TA had a high discriminatory potential, but its stability needed to be further confirmed. Firstly, the difference between bilateral hemispheres in the controls was compared using the six most discriminative parameters (Table V). For these six texture parameters, there were no significant differences between the bilateral hemispheres based on the Wilcoxon test. Subsequently, the six most

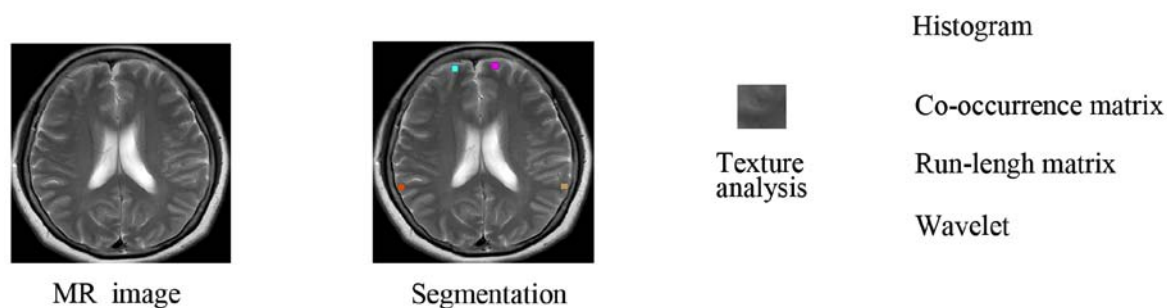


Figure 3. Workflow for brain MR image analysis. MR, magnetic resonance.

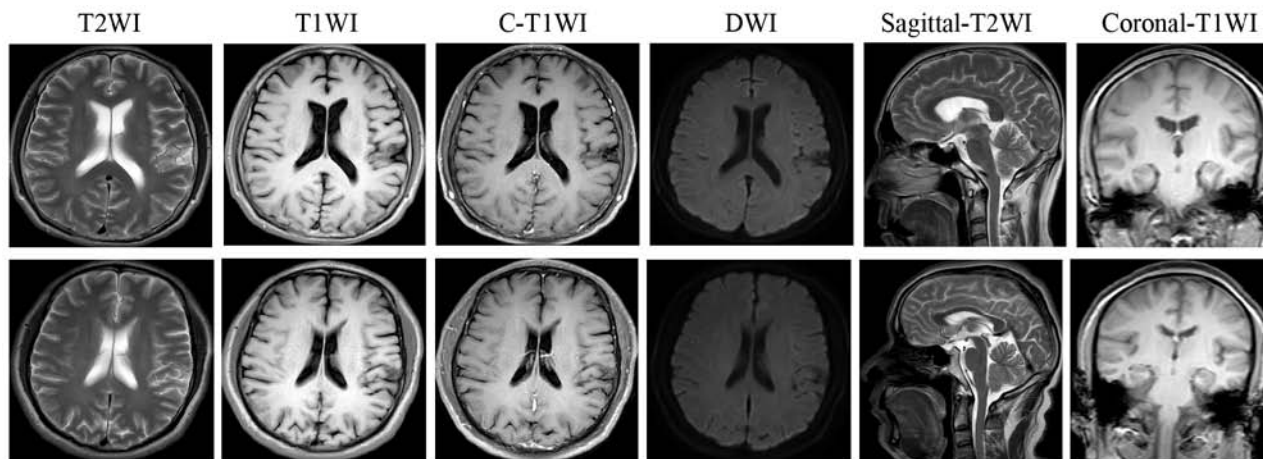


Figure 4. Different magnetic resonance sequences between patients and controls. Top row, the brain of a patient with lung cancer, without metastasis; bottom row, a healthy brain. T2WI, T2-weighted imaging; T1WI, T1-weighted imaging; C-T1WI, contrast-enhanced T1 weighted imaging; DWI, diffusion-weighted imaging.

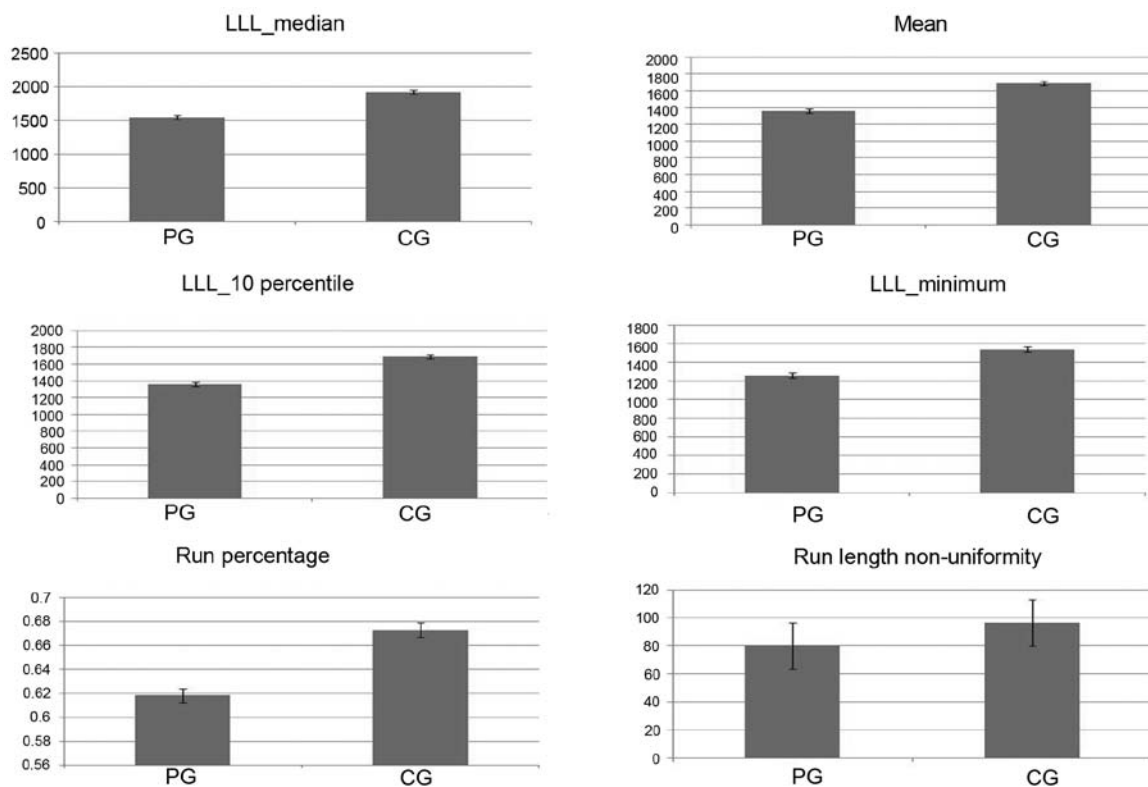


Figure 5. Comparison of different texture patterns between the PG and CG groups. The six features demonstrated significant differences between the two groups. PG, patient group; CG, control group; LLL, low frequency-low frequency-low frequency output.

discriminative parameters were used to find out if primary lung cancer had the same impact on the bilateral hemispheres. For these six texture parameters, there were no significant differences between the bilateral hemispheres in patients with lung cancer (Table V).

Comparison of the top six texture parameters between gray and white matter in patients with lung cancer without brain metastasis and healthy controls. The top six texture parameters were tested to find out how many and which of them differed statistically between gray and white matter. The texture features of gray and white matter in patients with lung cancer and healthy controls were also analyzed using the Wilcoxon test and based on the top six texture parameters. An identical trend between gray and white matter was observed in the patient and control groups. Among the six parameters, four parameters were significantly different between the patient and control groups. These four parameters were wavelet-LLL_median, wavelet-LLL_ 10 percentile, wavelet-LLL_ minimum, and run length non-uniformity. The parameters of mean and run percentage showed no significant difference (Table VI).

Comparison of texture parameters between patients with lung cancer without brain metastasis and healthy controls in cerebral gray and white matter. The aforementioned six most discriminative parameters were used to determine which differed statistically between patients with lung cancer without brain metastasis and healthy controls in different cerebral regions. In white matter regions, all six texture parameters analyzed with Wilcoxon test showed significant differences between patients and controls (Table VII). The six most discriminative parameters were subsequently tested to determine whether primary lung cancer has the same effect on the gray matter regions. For these six texture parameters, all showed significant differences between patients and controls (Table VII).

Comparison of the texture features by clinical stage in patients with lung cancer without brain metastasis. To investigate the relationship between the texture parameters of brain MR images and clinical lung cancer stages, which was a revised 2017 edition of the American Joint Committee on Cancer (AJCC) Cancer Staging Manual (28,29), patients with lung cancer, without brain metastasis were divided into two groups: Stage I+II and stage III+IV. The top six texture parameters were tested to find out how many and which of them differed statistically between the 2 patient groups. All of the six texture parameters were significantly different between the patient groups (Table VIII). Compared to the stage I+II group, all six parameters were significantly greater in the stage III+IV group (Table VIII).

Comparison of the top six texture features by histological type. The main histological types of lung cancer are squamous cell carcinoma, adenocarcinoma and small cell lung cancer (SCLC). The top six texture parameters were compared among these groups to establish associations between the brain MRI texture parameters and the histological types (Table IX). The top six texture parameters showed no significant differences between the squamous cell and adenocarcinoma groups but were all significantly different between the SCLC group

Table V. Comparison of the top six texture features between the left and right hemispheres in patients with lung cancer, without brain metastasis and healthy controls.

Feature	Patients		P-value	Controls		P-value
	Right, mean (SD)	Left, mean (SD)		Right, mean (SD)	Left, mean (SD)	
LLL_median	1,526.338 (442.99)	1,561.828 (478.29)	0.492	1,906.664 (580.25)	1,947.927 (591.44)	0.364
Mean	548.260 (160.31)	560.092 (173.88)	0.554	681.227 (206.92)	696.062 (211.67)	0.375
LLL_10 percentile	1,344.940 (387.64)	1,371.958 (417.95)	0.468	1,668.751 (520.21)	1,711.199 (528.96)	0.284
LLL_minimum	1,246.636 (365.12)	1,268.253 (386.43)	0.528	1,536.064 (488.56)	1,540.206 (529.67)	0.590
Run percentage	0.616 (0.11)	0.621 (0.11)	0.665	0.672 (0.10)	0.673 (0.10)	0.890
Run length non-uniformity	19.620 (32.86)	80.760 (33.54)	0.692	96.930 (35.30)	96.470 (34.29)	0.875

P<0.05 was considered to indicate a statistically significant difference. LLL, low frequency-low frequency output; HHH, high frequency-high frequency output; HHL, high frequency-high frequency-low frequency output; HLL, low frequency-low frequency-high frequency output; HHL, high frequency-high frequency-low frequency output.

Table VI. Comparison of the top six texture features between gray matter and white matter in patients with lung cancer, without brain metastasis and healthy controls.

Feature	Patients			Controls		
	Gray matter, mean (SD)	White matter, mean (SD)	P-value	Gray matter, mean (SD)	White matter, mean (SD)	P-value
LLL_median	1,578.194 (470.68)	1,407.642 (392.67)	0.001	1,974.210 (599.28)	1,739.639 (486.80)	<0.001
Mean	567.995 (170.78)	498.900 (139.48)	<0.001	706.957 (213.88)	615.395 (171.85)	<0.001
LLL_10 percentile	1,362.905 (410.33)	1,340.638 (373.17)	0.608	1,694.792 (538.49)	1,670.707 (466.43)	0.661
LLL_minimum	1,250.471 (379.24)	1,285.339 (361.70)	0.389	1,521.724 (522.89)	1,603.779 (445.74)	0.124
Run percentage	0.647 (0.09)	0.504 (0.09)	<0.001	0.703 (0.08)	0.549 (0.09)	<0.001
Run length non-uniformity	88.046 (30.90)	48.790 (21.37)	<0.001	106.039 (30.97)	59.380 (21.65)	<0.001

P<0.05 was considered to indicate a statistically significant difference. LLL, low frequency-low frequency output; HHH, high frequency-high frequency output; HHL, high frequency-high frequency-low frequency output; HLH, high frequency-low frequency-high frequency output.

Table VII. Comparison of the top six texture features between cerebral gray and white matter in patients with lung cancer, without brain metastasis and healthy controls.

Feature	Gray matter			White matter		
	Patients, mean (SD)	Controls, mean (SD)	P-value	Patients, mean (SD)	Controls, mean (SD)	P-value
LLL_median	1,578.194 (470.68)	1,974.210 (599.28)	<0.001	1,407.642 (392.67)	1,739.639 (486.80)	<0.001
Mean	567.995 (170.78)	706.957 (213.88)	<0.001	498.900 (139.48)	615.395 (171.85)	<0.001
LLL_10 percentile	1,362.905 (410.33)	1,694.792 (538.49)	<0.001	1,340.638 (373.17)	1,670.707 (466.43)	<0.001
LLL_minimum	1,250.471 (379.24)	1,521.724 (522.89)	<0.001	1,285.339 (361.70)	1,603.779 (445.74)	<0.001
Run percentage	0.647 (0.09)	0.703 (0.08)	<0.001	0.504 (0.09)	0.549 (0.09)	<0.001
Run length non-uniformity	88.046 (30.90)	106.039 (30.97)	<0.001	48.790 (21.37)	59.380 (21.65)	<0.001

P<0.05 was considered to indicate a statistically significant difference. LLL, low frequency-low frequency output.

Table VIII. Comparison of the top six texture features between different clinical stages in patients with lung cancer, without brain metastasis.

Feature	Stage I+II, mean (SD)	Stage III+IV, mean (SD)	P-value
LLL_median	1,348.190 (196.51)	1,649.503 (525.46)	<0.001
Mean	482.068.995 (72.08)	592.987 (190.12)	<0.001
LLL_10 percentile	1,181.089 (183.65)	1,453.300 (455.26)	<0.001
LLL_minimum	1,093.485 (171.46)	1,344.760 (425.07)	<0.001
Run percentage	0.580 (0.10)	0.636 (0.11)	<0.001
Run length non-uniformity	70.024 (27.21)	85.705 (34.94)	<0.001

P<0.05 was considered to indicate a statistically significant difference. LLL, low frequency-low frequency-low frequency output.

compared with the other groups. Compared to the adenocarcinoma group, the SCLC group showed significantly increased texture parameters; similar results were observed between the squamous cell carcinoma and SCLC groups.

Discussion

In light of recent advances in the acquisition and analysis of MR images for the high-throughput extraction of imaging feature information, medical imaging can now be used to quantify the distinguishing features of tumor tissues (30-32). This approach is distinct from prior subjective or qualitative methods (14). Applying TA to medical imaging features has been a significant field of research and has generated an extensive body of literature (33-35). It has been demonstrated that MR images include tissue-specific texture features that can be extracted by mathematical methods (30). TA can be used to distinguish pathological from healthy human cerebral tissue and to classify different types of cerebral tissue (15,36,37). TA enables the discrimination of multiple sclerosis lesions from normal and normal-appearing white matter (38). However, previous studies have mainly focused on detecting small nodules, that is, the lesions themselves. This study is distinct in that it primarily characterizes early cerebral tissue damage caused by lung cancer before metastasis by applying TA methods. The TA of MRI features can also be used for classifying different cerebral tissues and structures in brain MR images (36,39).

MR images were selected in the current study to detect textural differences in cerebral tissue between patients with primary lung cancer and healthy controls. Of a total of 32 texture parameters tested, 27 parameters were significantly different between patients with lung cancer and healthy controls. The results of this study indicate that in patients with lung cancer, there are differences in the texture parameters of brain MR images of cerebral tissues. It can therefore be inferred that lung cancer can cause early brain tissue damage prior to metastasis.

Among the 27 significantly different parameters, six texture features that best discriminated between patients and healthy controls were identified by calculating Fisher coefficients. These texture parameters were the LLL_median, mean, LLL_percentile, LLL_minimum, run percentage and run length non-uniformity. Using these six parameters, the two cerebral

hemispheres and gray and white matter were compared between and within the participant groups. First, the left and right hemispheres of cerebral tissue were compared in the controls, as well as the gray and white matter. It is well known that the human brain is symmetrical in structure and with respect to gray and white matter, there are obvious differences in both composition and function. Gray matter is mainly composed of the cell bodies of neurons, but white matter is principally surrounded by the myelin sheaths of neurons. In regard to functionality, the former contains nerve centers, which play a major role in generating neural activity; the latter, on the other hand, represents the main transmission pathways carrying the neural signals generated by the brain and the gray matter of the spinal cord (40,41). The results of the present study show that the texture parameters were not significantly different between the 2 hemispheres, but 4 of 6 texture parameters were significantly different between gray and white matter. It can therefore be concluded that TA applied to brain MR images is stable and reliable. This analysis was repeated in the patient group and an identical trend was observed, which may indicate that primary lung lesions can cause similar early damage in both hemispheres and that TA can be used to classify different structures. By comparison of different regions of gray and white matter between patients with lung cancer and healthy controls, all six texture parameters were significantly different in both gray matter and white matter. It can be assumed that the overt texture changes are caused by the lung cancer and this leads to similar impairments in both gray and white matter.

Clinical stage is used to describe the severity and range of involvement of malignant tumors according to the original tumor and degree of spread in an individual, which is the basis for understanding the extent of the disease. This information can help doctors develop appropriate treatment plans and understand the prognosis and outcome of the disease. The numerically higher the stage, the more advanced the tumor progression and the worse the prognosis (42). The present study analyzed associations between the clinical stages of lung cancer and texture parameters of MR images. The results revealed that, compared with the early-stage group (stage I+II), the advanced-stage group (stage III+IV) showed a more significant increase in all six parameters. This may indicate that increased parameters are related to more advanced clinical stages of lung cancer.

In addition to the clinical stage, the choice of tumor treatment is also related to the pathological type of the

Table IX. Comparison of the top six texture features between different histological types in patients with lung cancer, without brain metastasis.

Feature	Mean (SD)			P-value		
	Ad	Sq	Sm	Ad vs. Sq	Ad vs. Sm	Sq vs. Sm
LLL_median	1,450.823 (18.75)	1,544.848 (449.86)	2,565.255 (451.93)	0.334	<0.001	<0.001
Mean	519.909 (124.32)	555.842 (164.23)	921.111 (158.10)	0.234	<0.001	<0.001
LLL_10 percentile	1,274.496 (300.34)	1,362.517 (378.48)	2,268.574 (413.46)	0.141	<0.001	<0.001
LLL_minimum	1,176.569 (281.63)	1,267.834 (350.97)	2,084.727 (425.21)	0.051	<0.001	<0.001
Run percentage	0.603 (0.11)	0.623 (0.09)	0.759 (0.06)	0.145	<0.001	<0.001
Run length non-uniformity	75.889 (31.78)	80.426 (30.69)	126.191 (28.12)	0.158	<0.001	<0.001

Ad, adenocarcinoma; Sq, squamous cell; Sm, small cell; LLL, low frequency-low frequency output.

tumor (18,28,43-46). Associations with the histological type were also analyzed in this study. The results show that compared to adenocarcinoma group, the SCLC group showed significantly increased texture parameters; similar results were found between the squamous cell carcinoma and SCLC groups. However, the texture parameters assessed in the adenocarcinoma and squamous cell carcinoma groups were not significantly different. It is necessary to note that compared with adenocarcinoma and squamous cell carcinoma, SCLC has a higher degree of malignancy and a worse prognosis (47,48). The significant differences in texture parameters between SCLC and adenocarcinoma or squamous cell carcinoma are possibly attributable to these differences. It can be assumed that greater texture changes are caused by greater degrees of malignancy.

There are several limitations of this study. Firstly, as this is a preliminary experiment, the sample size was relatively small. Furthermore, a limited set of features, rather than a larger set of multiple hundreds or thousands (compound) of features was selected for analysis because the selected texture features have consistently been shown to be effective in the analysis of medical images (25,38,49-51) and robust against variations between scanners and protocol parameters (52) especially after normalization (53). Finally, in the absence of confirmation of the early damage to human brain tissue, further pathological investigation using an animal model is required.

In conclusion, this small pilot study indicates that TA with a variable set of texture features can serve as an adjuvant diagnostic tool to traditional MRI to detect early damage in patients with lung cancer without brain metastasis. TA may serve as a new method for the study of tumor metastasis and paraneoplastic syndromes.

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Availability of data and materials

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

Authors' contributions

JX acquired the data, performed the analyses and wrote the initial manuscript. XC and BW performed texture analysis. GW, MH, RL and YQ acquired the data. JX and QY selected ROIs of the MR images. ZL and MH made substantial contributions to the conception and design of the present study. All authors read and approved the final manuscript.

Ethical approval and consent to participate

The present study was approved by the Ethics Committee of Shandong Provincial Hospital Affiliated to Shandong

University (Jinan, China). Written informed consent was obtained from all patients prior to the study start.

Patient consent for publication

All patients signed consent for publication.

Competing interests

The authors declare that they have no competing interests.

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