

Radiomic analysis of pulmonary ground-glass opacity nodules for distinction of preinvasive lesions, invasive pulmonary adenocarcinoma and minimally invasive adenocarcinoma based on quantitative texture analysis of CT

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Abstract

Objective: To identify the differences among preinvasive lesions, minimally invasive adenocarcinomas (MIAs) and invasive pulmonary adenocarcinomas (IPAs) based on radiomic feature analysis with computed tomography (CT).

Methods: A total of 109 patients with ground-glass opacity lesions (GGOs) in the lungs determined by CT examinations were enrolled, all of whom had received a pathologic diagnosis. After the manual delineation and segmentation of the GGOs as regions of interest (ROIs), the patients were subdivided into three groups based on pathologic analyses: the preinvasive lesions (including atypical adenomatous hyperplasia and adenocarcinoma *in situ*) subgroup, the MIA subgroup and the IPA subgroup. Next, we obtained the texture features of the GGOs. The data analysis was aimed at finding both the differences between each pair of the groups and predictors to distinguish any two pathologic subtypes using logistic regression. Finally, a receiver operating characteristic (ROC) curve was applied to accurately evaluate the performances of the regression models.

Results: We found that the voxel count feature ($P < 0.001$) could be used as a predictor for distinguishing IPAs from preinvasive lesions. However, the surface area feature ($P = 0.040$) and the extruded surface area feature ($P = 0.013$) could be predictors of IPAs compared with MIAs. In addition, the correlation feature ($P = 0.046$) could distinguish preinvasive lesions from MIAs better.

Conclusions: Preinvasive lesions, MIAs and IPAs can be discriminated based on texture features within CT images, although the three diseases could all appear as GGOs on CT images. The diagnoses of these three diseases are very important for clinical surgery.

Keywords: CT; GGO; IPA; MIA; preinvasive lesions; radiomic analysis

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Introduction

Lung cancer is the leading cause of cancer-related deaths in the world (1), and adenocarcinomas as the most common

histological subtype were divided into preinvasive lesions, minimally invasive adenocarcinomas (MIAs) and invasive pulmonary adenocarcinomas (IPAs) collectively by the joint of International Association for the Study of Lung Cancer,

the American Thoracic Society and the European Respiratory Society (2) in 2011.

Preinvasive lesions include atypical adenomatous hyperplasia (AAH) and adenocarcinoma *in situ* (AIS). AAH characteristically appears as a very small and pure ground-glass opacity lesion (GGO) (3-6). AIS usually presents as a pure GGO but sometimes is partly solid or occasionally is a solid nodule (4,7-9). The imaging presentations of MIAs and IPAs can be pure GGOs or partly-solid nodules or even a solid nodule (9-14).

It can be seen that all of them could present as GGOs during CT examinations. Previous studies have shown that preinvasive lesions can be followed up for a long time and MIAs have either 100% or near 100% 5-year disease-free survival (DFS) after complete resection (2,11). In contrast, the 5-year DFS of an IPA of the pathological stage IA is 74.6% (15). Therefore, early diagnosis and exact discrimination of these diseases that presents as GGOs on CT are of great importance.

Previous studies regarding the identification of benign/malignant nodules among GGOs usually focused on lesion size (3-14), the appearance of the solid component (16), the detection of growth or the density change of GGOs (17). Many studies have suggested that the GGOs lesion sizes were significantly different among AIS, MIAs and IPAs (18-20), and the solid component size within GGO lesions could also discriminate IPAs from both MIAs and AIS (21). Changes in the size and density of GGOs were found in follow-up studies; however, they did not significantly help in differentiating preinvasive lesions and MIAs from IPAs (21-23).

However, now, except for these morphologic features, there are still no generally accepted quantitative guidelines that have been issued for GGO diagnosis. Radiomic analysis, as a more systematic approach, may provide more information regarding the discrimination of preinvasive lesions from IPAs or MIAs, as it is able to identify voxel-level changes within GGOs. Thus, our study presents the hypothesis that there may be some radiomic features that can be used to discriminate GGOs. The present study aims to find the differences among GGOs in CT images with pathological diagnoses in the IPAs, MIAs, and preinvasive lesion categories using radiomic analysis.

Materials and methods

The study was approved by the Medical Research Ethics

Committee and the Institutional Review Board of Tianjin Medical University Cancer Institute & Hospital.

Patient population

The inclusion criteria for the lung cancer patients in our study were as follows: 1) patients with a pathological diagnosis of cancer; 2) diagnostic CT scans before surgery; 3) CT images with a slice thickness of 2.5 mm or less; and 4) a single GGO. Patients were excluded if the lesions were either very small (<5 mm) or very large (>5 cm). Ultimately, there were 109 patients (mean age, 57.1 years; range, 38–77 years) qualified for our study.

Thirty-two (29.4%) of the patients had preinvasive lesions (including 22 patients who were diagnosed with AAH and 10 patients who were diagnosed with AIS), 37 (33.9%) had MIAs, and 40 (36.7%) had IPAs.

CT examinations

In our study, all the CT examinations were performed without contrast enhancement using a high-definition CT system (Discovery CT750HD, GE Healthcare, Milwaukee, Wisconsin, USA), and the scanning parameters were as follows: DFOV = 40 cm; tube voltage = 120 kVp; slice thickness = 1.25 mm; reconstruction interval = 1.25 mm; gantry rotation speed = 0.8 s; and a scanning range from the apex of the lung to the base to lung.

Delineation of ROI

The ROI in the present study was manually delineated and segmented slice-by-slice on the CT images. The ROI covered the whole lesion, and the large vessels and arteries in the ROI were excluded. Finally, the seed ROI was checked slice-by-slice in every subject by another radiologist to ensure that the ROI in each subject satisfied the lesion boundary definition.

Computerized texture analysis based on ROI

Radiomic analysis was performed using the 3D slicer software (Version 4.6.2; Surgical Planning Laboratory, Brigham and Women's Hospital, MA, USA; <http://www.slicer.org>) (24). Then, the texture features were calculated and extracted automatically using the module called "Heterogeneity CAD". The radiomic features (a total of 62 features; *Supplementary Table S1*) were divided into six categories, including: 1) first-order and distribution

statistics; 2) shape and morphology metrics; 3) Renyi dimensions; 4) geometrical measures; 5) the gray-level co-occurrence matrix (GLCM); and 6) the gray-level run length matrix (GLRL). The details of the significantly different features that were found to distinguish any two pathologic subtypes in the present study are shown in *Table 1* and all other feature instructions can be found in the 3D slicer software (<https://www.slicer.org/wiki/Documentation/Nightly/Modules/HeterogeneityCAD>).

The overall procedure of this analytical scheme was performed by two radiologists with more than 3 years of experience in chest CT imaging. Finally, we computed the means of each of the CT texture feature values measured by the two independent observers. The interobserver agreement regarding the texture features of the GGO ROIs was calculated using the interclass correlation coefficient analysis (ICC) with SPSS software.

Statistical analysis

Features of the study population were reported as means followed by standard deviations or medians followed by ranges in light of their normality for the continuous variables, and as frequencies for the categorical variables. Initially, Shapiro-Wilk tests and Bartlett’s tests were performed to confirm the normality and variance homogeneity of the data regarding the features. Subsequently, a Kruskal-Wallis rank sum test followed by Dunn-Bonferroni *post hoc* test or an analysis of variance (ANOVA) followed by Tukey’s HSD *post hoc* test where appropriate, was used to indicate the significant differences among the three groups. The data were corrected by Bonferroni’s approach (P<0.05) with two-sided to control for the type-1 errors.

Logistic regression analysis was conducted to find better predictors in distinguishing any two of the pathologic

subtypes. Features with P<0.05 in univariable model were entered into the multiple logistic regression analysis. The stepwise model selection that mixed both forward and backward methods of variable selection using Akaike’s information criterion (AIC) was used to select the final predictive model, and feature with P>0.05 was eliminated. Receiver operating characteristic (ROC) curves for each model were constructed. AUC and its 95% confidence interval (95% CI) estimated using DeLong’s method were calculated to evaluate the performances of the regressive models. P<0.05 was considered statistically significant.

All statistical analyses were conducted with Microsoft R Open (version 3.2.2; <https://mran.microsoft.com/>) in Microsoft Visual Studio Professional 2015 (version 14.0.25431.01 Update 3), and with IBM SPSS Statistics (Version 21.0; IBM Corp., New York, USA).

Results

Basic characteristics of patients and texture features of GGOs

Patients’ basic characteristics and lesion information are shown in *Table 2* and *Supplementary Figure S1*. The interobserver agreement regarding the texture features of the GGO ROI was generally acceptable (the value ranged from 0.417 to 0.999) (*Supplementary Table S2*).

Differences between preinvasive lesions and IPAs

Significant differences between the preinvasive lesions (including AAH and AIS) and the IPAs appeared in most of the texture features of the first category (first-order and distribution statistics) (*Table 3*), the second category (shape and morphology metrics) (*Table 4*) and the texture matrix categories, such as the GLCM (*Table 5*), the GLRL (*Table 6*), and other categories (*Table 7*).

Table 1 Feature instructions with significant differences that served to distinguish each pair of groups of preinvasive lesions, MIAs and IPAs

Feature category	Feature name	Feature instructions
First-order and distribution statistics	Voxel count	Total number of voxels within the ROI of the grayscale image or parameter map
Shape and morphology metrics	Surface area (mm ²)	Surface area of the specified ROI of the image in square millimeters
GLCM	Correlation	A value between 0 (uncorrelated) and 1 (perfectly correlated) showing the linear dependency of gray level values in the GLCM
Geometrical measures	Extruded surface area	Surface area of the binary object when the image ROI is “extruded” into 4D, where the parameter or intensity value defines the shape of the Fourth dimension

MIAs, minimally invasive adenocarcinomas; IPAs, invasive pulmonary adenocarcinomas; GLCM, gray-level co-occurrence matrix; ROI, region of interest.

Table 2 Basic characteristics of patients (N=109)

Characteristics	n (%)		
	Preinvasive lesions (N=32)	MIAs (N=37)	IPAs (N=40)
Female	19 (25.7)	27 (36.5)	28 (37.8)
Male	13 (37.1)	10 (28.6)	12 (34.3)
Age (year)			
$\bar{x}\pm s$	53.4±11.6	57.5±9.2	59.8±7.9
Range	38–76	38–77	42–76
Lesion location			
Upper lobe (right/left)	20 (29.4)	24 (35.3)	24 (35.3)
Middle lobe (right)	2 (20.0)	5 (50.0)	3 (30.0)
Lower lobe (right/left)	10 (32.3)	8 (25.8)	13 (41.9)

MIAs, minimally invasive adenocarcinomas; IPAs, invasive pulmonary adenocarcinomas.

Table 3 Differences in texture features in first-order and distribution statistics category between each pair of groups of preinvasive lesions, MIAs and IPAs

Texture features	P value		
	Preinvasive lesions vs. MIAs	Preinvasive lesions vs. IPAs	MIAs vs. IPAs
Gray levels	>0.05	0.001	<0.001
Voxel count	>0.05	<0.001	<0.001
Energy	>0.05	0.004	<0.001
Entropy	>0.05	0.003	<0.001
Minimum intensity	>0.05	>0.05	>0.05
Maximum intensity	>0.05	0.028	>0.05
Mean intensity	>0.05	0.034	>0.05
Median intensity	>0.05	>0.05	>0.05
Range	>0.05	>0.05	>0.05
Mean deviation	>0.05	0.008	>0.05
Root mean square	>0.05	>0.05	>0.05
Standard deviation	>0.05	0.007	>0.05
Skewness	>0.05	>0.05	>0.05
Kurtosis	>0.05	>0.05	>0.05
Variance	>0.05	0.007	>0.05
Uniformity	>0.05	0.002	<0.001

MIAs, minimally invasive adenocarcinomas; IPAs, invasive pulmonary adenocarcinomas. P value was calculated with Kruskal-Wallis rank sum test followed by Dunn-Bonferroni *post hoc* test.

Table 4 Differences in texture features in shape and morphology metrics category between each pair of groups of preinvasive lesions, MIAs and IPAs

Texture features	P value		
	Preinvasive lesions vs. MIAs	Preinvasive lesions vs. IPAs	MIAs vs. IPAs
Volume (mm ³)	>0.05	0.001	<0.001
Surface area (mm ²)	>0.05	0.002	<0.001
Surface: volume ratio	>0.05	0.008	<0.001
Compactness 1	>0.05	0.002	<0.001
Compactness 2	>0.05	0.012	0.045
Spherical disproportion	>0.05	0.001	<0.001
Sphericity	>0.05*	0.007*	0.038*
Maximum 3D diameter	>0.05	0.001	<0.001
Volume cc	>0.05	0.001	<0.001

MIAs, minimally invasive adenocarcinomas; IPAs, invasive pulmonary adenocarcinomas; Volume cc, volume in cubic centimeters; *, statistical analysis using analysis of variance (ANOVA) followed by Tukey's HSD *post hoc* test, otherwise using a Kruskal-Wallis rank sum test followed by Dunn-Bonferroni *post hoc* test.

Differences between IPAs and MIAs

Similarly, there were also significant differences between the IPAs and MIAs in terms of many of the texture features of the GGOs, which were found in the first category (first-order and distribution statistics) (*Table 3*), the second category (shape and morphology metrics) (*Table 4*) and the texture matrix categories, such as the GLCM (*Table 5*), the GLRL (*Table 6*) and other categories (*Table 7*).

Differences between preinvasive lesions and MIAs

Among the GGOs, the differences between the preinvasive lesions and MIAs in terms of the texture features were only found in the correlation feature of the GLCM category (*Table 5*).

Logistic regression analysis

Univariate logistic regression analysis revealed that the voxel count feature (P<0.001) of the first category (first-order and distribution statistics) was a predictor of preinvasive lesions compared with IPAs (*Table 8*). For the preinvasive lesions and the MIAs, the correlation feature (P=0.046) of the GLCM category was the only independent

Table 5 Differences in texture features in GLCM category between each pair of groups of preinvasive lesions, MIAs and IPAs

Texture features	P value		
	Preinvasive lesions vs. MIAs	Preinvasive lesions vs. IPAs	MIAs vs. IPAs
Autocorrelation	>0.05	<0.001	<0.001
Cluster prominence	>0.05	<0.001	<0.001
Cluster shade	>0.05	<0.001	<0.001
Cluster tendency	>0.05	<0.001	<0.001
Contrast	>0.05	<0.001	<0.001
Correlation	0.019	>0.05	<0.05
Difference entropy	>0.05	0.003	<0.001
Dissimilarity	>0.05	<0.001	<0.001
Homogeneity 1	>0.05	0.006	<0.001
Homogeneity 2	>0.05	0.005	<0.001
IMC1	>0.05	0.017	0.001
IDMN	>0.05	0.001	<0.001
IDN	>0.05	0.001	<0.001
Inverse variance	>0.05	0.020	<0.001
Maximum probability	>0.05	<0.001	<0.001
Sum average	>0.05	<0.001	<0.001
Sum entropy	>0.05	0.005	<0.001
Sum variance	>0.05	0.001	<0.001
Energy (GLCM)	>0.05	0.001	<0.001
Entropy (GLCM)	>0.05	0.004	<0.001
Variance (GLCM)	>0.05	<0.001	<0.001

GLCM, gray-level co-occurrence matrix; MIAs, minimally invasive adenocarcinomas; IPAs, invasive pulmonary adenocarcinomas; IMC1, informational measure of correlation 1; IDMN, inverse difference moment normalized; IDN, inverse difference normalized. P value was calculated with Kruskal-Wallis rank sum test followed by Dunn-Bonferroni *post hoc* test.

predictors (Table 8). Between the MIAs and IPAs, the multiple logistic regression analysis showed that the surface area feature (P=0.040) of the secondary category (shape and morphology metrics) and the extruded surface area feature (P=0.013) of the fourth category (geometrical measures) were independent predictors. (Table 8)

ROC analysis

Based on the univariate and multiple logistic regression analyses, we investigated whether the features could accurately distinguish IPAs from MIAs or preinvasive

Table 6 Differences in texture features in GLRL category between each pair of groups of preinvasive lesions, MIAs and IPAs

Texture features	P value		
	Preinvasive lesions vs. MIAs	Preinvasive lesions vs. IPAs	MIAs vs. IPAs
RLN	>0.05	0.002	<0.001
RP	>0.05	0.005	0.002
LGLRE	>0.05	0.022	>0.05
HGLRE	>0.05	0.001	0.035
SRLGLE	>0.05	0.031	>0.05
SRHGLE	>0.05	0.001	0.042
LRLGLE	>0.05	0.002	0.013
LRHGLE	>0.05	<0.001	0.007
LRE	>0.05	0.003	0.011
GLN	>0.05	0.010	<0.001
SRE	>0.05	0.011	>0.05

GLRL, gray-level run length matrix; MIAs, minimally invasive adenocarcinomas; IPAs, invasive pulmonary adenocarcinomas; RLN, run length nonuniformity; RP, run percentage; LGLRE, low gray level run emphasis; HGLRE, high gray level run emphasis; SRLGLE, short run low gray level emphasis; SRHGLE, short run high gray level emphasis; LRLGLE, long run low gray level emphasis; LRHGLE, long run high gray level emphasis; LRE, long run emphasis; GLN, gray level nonuniformity; SRE, short run emphasis. P value was calculated with Kruskal-Wallis rank sum test followed by Dunn-Bonferroni *post hoc* test.

lesions. The results showed that the voxel count feature was significantly different between the IPAs and preinvasive lesions, exhibiting 82.5% sensitivity and 62.5% specificity (95% CI: 0.650, 0.871; Figure 1A). The correlation feature could predict preinvasive lesions and MIAs better, with a sensitivity and specificity of 81.1% and 53.1%, respectively (95% CI: 0.533, 0.795; Figure 1B). The surface area feature (95% CI: 0.671, 0.878) and the extruded surface area feature (95% CI: 0.648, 0.863) both achieved a sensitivity and specificity of more than 60% and 80%, respectively, in distinguishing IPAs from MIAs (Figure 1C, D). However, when incorporating the two features as regression variables in predicting the pathological subgroup jointly, the corresponding ROC analysis showed higher sensitivity and specificity of 67.5% and 86.5%, respectively.

Discussion

IPAs, MIAs and preinvasive lesions (including AAH and AIS) can all appear as GGO lesions on CT scans. We used radiomic analysis to investigate the differences among these

Table 7 Differences in texture features in geometrical measures and Renyi dimensions categories between each pair of groups of preinvasive lesions, MIAs and IPAs

Category	Texture features	P value		
		Preinvasive lesions vs. MIAs	Preinvasive lesions vs. IPAs	MIAs vs. IPAs
Geometrical measures	Extruded surface area	>0.05	<0.001	<0.001
	Extruded volume	>0.05	0.002	<0.001
	Extruded surface: volume ratio	>0.05	>0.05	>0.05
Renyi dimensions	Box-counting dimension	>0.05*	>0.05*	>0.05*
	Correlation dimension	>0.05	0.001	<0.001

MIAs, minimally invasive adenocarcinomas; IPAs, invasive pulmonary adenocarcinomas; *, statistical analysis using analysis of variance (ANOVA) followed by Tukey's HSD *post hoc* test, otherwise using a Kruskal-Wallis rank sum test followed by Dunn-Bonferroni *post hoc* test.

Table 8 Logistic regression analyses of texture features between each pair of groups of preinvasive lesions, MIAs and IPAs

Features	Regression coefficients	P	OR	95% CI
Group of preinvasive lesions & Group of MIAs				
Correlation*	0.577	0.046	1.781	1.052–3.363
Group of preinvasive lesions & Group of IPAs				
Voxel count*	1.075	<0.001	2.930	1.704–5.451
Group of MIAs & Group of IPAs				
Surface area (mm ²)#	0.451	0.040	1.571	1.033–2.494
Extruded surface area#	-1.073	0.013	0.341	0.133–0.758

MIAs, minimally invasive adenocarcinomas; IPAs, invasive pulmonary adenocarcinomas; OR, odds ratio; 95% CI, 95% confidence interval; *, univariate logistic regression analysis; #, multiple logistic regression analysis entered with significant features (P<0.05) in univariable model. For univariate and multiple logistic regression analyses, the independent variable was each feature and the dependent variable was each group.

three diseases in the present study. We ultimately found that the extruded surface area feature and the surface area feature could be used to predict IPAs compared with MIAs. We also found that the correlation feature could distinguish the MIAs from the preinvasive lesions, and the voxel count feature could distinguish preinvasive lesions from IPAs better.

For a long time, the study of the assessment of GGOs using CT particularly used visual features such as lesion size and the presence of pleural retraction (25), the proportion of the solid component, the shape and size of the lesion, and the window width (26). However, it was not an easy or accurate method for identifying the different diseases. To improve the diagnoses of the three diseases and to address the challenge of differential diagnoses, many studies have adopted quantitative methodological analyses to evaluate GGOs. Texture features provide a more comprehensive assessment of GGOs. Some features have been found to be especially sensitive in the detection of tumor heterogeneity.

In recent years, several studies have applied texture analysis of GGOs. One study (27) found that higher kurtosis and smaller mass are significant differentiators of preinvasive lesions from IPAs; however, they only focused on partly-solid GGOs and few texture features, such as histogram features, volumetric features, and morphological features. Son *et al.* (28) studied nodule size, volume, density, mass, skewness/kurtosis, and the CT attenuation values of the histograms, in addition to the texture features (uniformity and entropy) of ground-glass opacity nodules with little or no solid component using CT. That study found that the CT attenuation values and entropy could be used as independent predictors of invasive adenocarcinomas. Quantitative analyses of preoperative CT imaging metrics can help to distinguish invasive adenocarcinomas from preinvasive lesions or MIAs.

Based on the preceding studies, the present studies focused on GGOs and obtained more than 50 texture features, including not only morphology but also matrix information regarding the GGOs. In our study, we focused

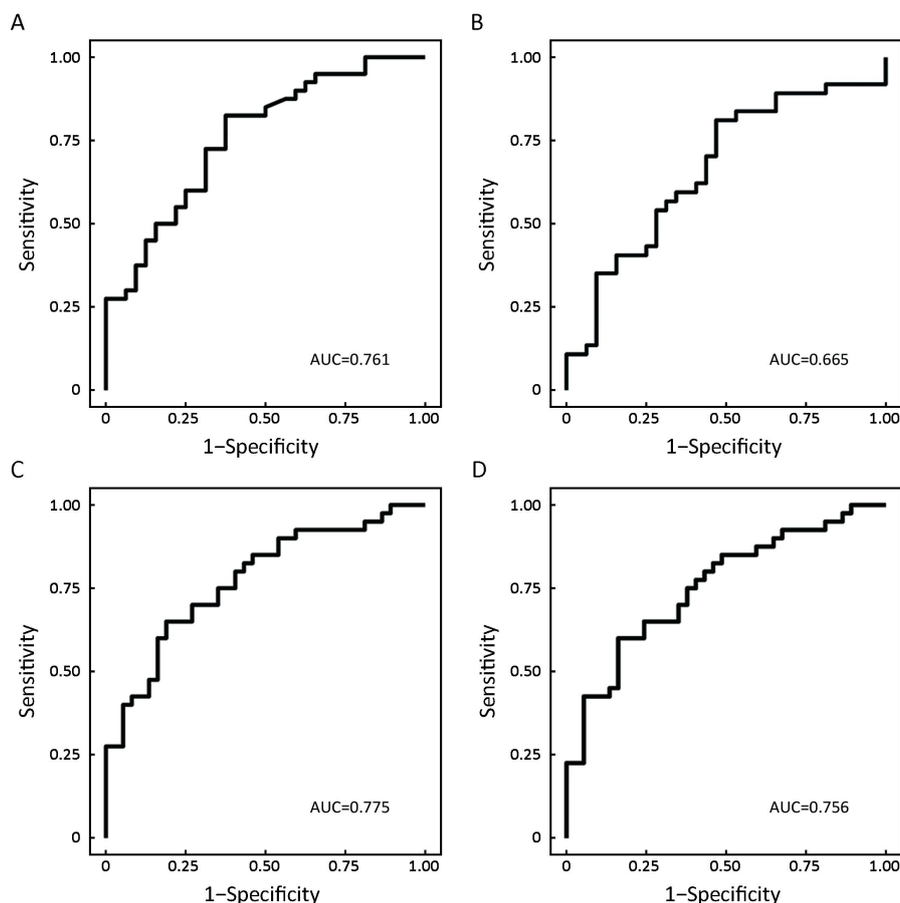


Figure 1 Receiver operating characteristic (ROC) analysis of significant differences between each pair of groups of preinvasive lesions, minimally invasive adenocarcinomas (MIAs) and invasive pulmonary adenocarcinomas (IPAs). (A) Voxel count [area under the curve (AUC)=0.761]; (B) Correlation (AUC=0.665); (C) Surface area (AUC=0.775); (D) Extruded surface area (AUC=0.756).

on not only partly-solid GGO lesions but also many of the texture features. We found that many texture features, including morphology and the matrix information, were significantly different between each pair of the groups of preinvasive lesions, MIAs and IPAs; however, we did not find that the kurtosis (27) or entropy (28) were significant predictors of preinvasive lesions, MIAs or IPAs. This may have been due to the different sample sizes and categories of lesions.

In our study, we found that the voxel count feature was a significant differentiator of preinvasive lesions from IPAs. The voxel count is the total number of voxels within an ROI, therefore, the voxel count feature reflects the size and mass of GGOs. We found that a larger voxel count increased the probability of IPAs. Therefore, it could be suggested that preinvasive lesions have smaller voxels; in others words, they have smaller sizes or masses. In fact,

many previous studies (20) involving the morphological assessment of GGOs revealed that preinvasive lesions could be distinguished from IPAs with smaller lesion sizes. In turn, the GGOs of larger sizes or with larger solid components had a higher likelihood of being invasive adenocarcinomas (MIAs and IPAs) (29,30). A study based on texture analysis of partly solid GGOs found that the preinvasive lesions had significantly smaller volumes and masses than IPAs (27). The findings of our study corresponded to the findings of previous studies well.

We also found that the correlation feature was a significant differentiator of preinvasive lesions from MIAs. Correlation is a value that shows the linear dependency of gray level values in the GLCM. This feature reflects the correlation of the gray level in part of the images; its value reflects the contrast of the gray levels in the GLCM. The greater the mean of the gray level value of the GLCM is,

the greater the value of the correlation is. In other words, when the gray level value of the GLCM is different, the value of the correlation is smaller. In our study, we found that a higher correlation was a significant differentiator of MIAs from preinvasive lesions. Several studies have suggested that increased heterogeneity is associated with malignant lung cancer (31) in GGOs and the solid portion within the lesion tends to be found in invasive lesions (32). Heterogeneous changes in the ROIs in our study may have been reflected by the gray level values and even by the correlation feature, which is the reflection of gray level values in the GLCM.

MIAs and IPAs are both invasive lesions, and in our study, we found that the surface area feature and the extruded surface area feature were both differentiators of IPAs from MIAs. The extruded surface area feature is the measurement of the surface area when the image ROI is “extruded” into 4D space, in which the feature or intensity value defines the shape of the fourth dimension. The surface area feature measures the area of ROI two-dimensionally. In other words, the surface area feature and the extruded surface area feature both reflect the macroscopic size and shape of the ROI mass. IPAs are more malignant than MIAs in terms of pathology and disease recovery. Many studies of GGO morphology have suggested that IPAs have greater diameters than MIAs [5 mm or less (10,11) in diameter (4,9,10,12-14)]. In our study, we focused on the texture features of GGOs. The surface area feature and the extruded surface area feature reflected more global information regarding the size and shape of the ROI; and thus we believe that the surface area feature and the extruded surface area feature are more suitable for the reevaluation of GGOs than the diameter of a nodule. More importantly, these features could reflect the homogeneity of ROIs using visualized data, and then be used to distinguish IPAs from MIAs.

However, our study still had several limitations. First, the sample size of this study was not big enough. In future studies, we will enlarge the sample size and do some prospective research. Second, the manual delineation of the ROI may have been bound to have had some errors. In future studies, when the sample size is big enough, it will be necessary to consider automatic segmentation of the ROI and then manual correction slice-by-slice. Third, the texture features in this study did not cover a large scale of features within the GGOs. In future studies, more detailed information hidden in the lesion images needs to be obtained.

Conclusions

In our study based on texture features analyses of GGOs, the voxel count feature was a significant differentiator of preinvasive lesions from IPAs. The preinvasive lesions can be differentiated from MIAs using the correlation feature. The differences between MIAs and IPAs could be accurately reflected by the surface area feature and the extruded surface area feature. Further details and proper explanations of the texture features require more studies.

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Footnote

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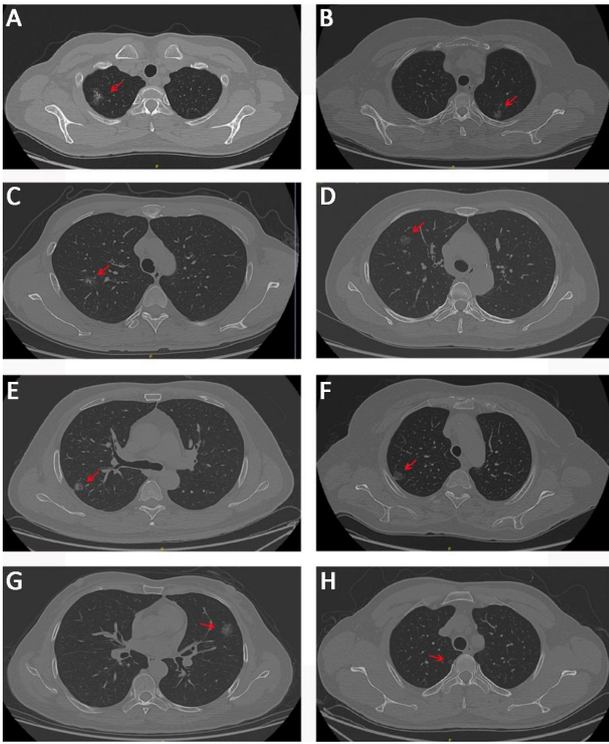


Figure S1 Computed tomography (CT) images of typical patients' lesions. A–H stand for eight different patients' ground-glass opacity (GGO) lesions on axial CT images.

Table S1 Quantitative feature categories and feature names

Feature category	Feature numbers and feature names		
First-order and distribution statistics	F1 Gray levels	F7 Mean intensity	F13 Skewness
	F2 Voxel count*	F8 Median intensity	F14 Kurtosis
	F3 Energy	F9 Range	F15 Variance
	F4 Entropy	F10 Mean deviation	F16 Uniformity
	F5 Minimum intensity	F11 Root mean square	
	F6 Maximum intensity	F12 Standard deviation	
Shape and morphology metrics	F17 Volume (mm ³)	F20 Compactness 1	F23 Sphericity
	F18 Surface area (mm ²)*	F21 Compactness 2	F24 Volume cc
	F19 Surface: volume ratio	F22 Spherical disproportion	F25 Maximum 3D diameter
GLCM	F26 Autocorrelation	F33 Homogeneity 1	F40 Sum entropy
	F27 Cluster prominence	F34 Homogeneity 2	F41 Sum variance
	F28 Cluster shade	F35 IMC1	F42 IDN
	F29 Cluster tendency	F36 IDMN	F43 Contrast
	F30 Variance (GLCM)	F37 Energy (GLCM)	F44 Dissimilarity
	F31 Correlation*	F38 Inverse variance	F45 Sum average
	F32 Difference entropy	F39 Maximum probability	F46 Entropy (GLCM)
GLRL	F47 RLN	F51 SRLGLE	F55 LRE
	F48 RP	F52 SRHGLE	F56 GLN
	F49 LGLRE	F53 LRLGLE	F57 SRE
	F50 HGLRE	F54 LRHGLE	
Geometrical measures	F58 Extruded surface area*	F59 Extruded surface: volume ratio	F60 Extruded volume
Renyi dimensions	F61 Box-counting dimension	F62 Correlation dimension	

GLCM, gray-level co-occurrence matrix; GLRL, gray-level run length matrix; RLN, run length nonuniformity; RP, run percentage; LGLRE, low gray level run emphasis; HGLRE, high gray level run emphasis; IMC1, informational measure of correlation 1; IDMN, inverse difference moment normalized; SRLGLE, short run low gray level emphasis; SRHGLE, short run high gray level emphasis; LRLGLE, long run low gray level emphasis; LRHGLE, long run high gray level emphasis; Volume cc, volume in cubic centimeters; IDN, inverse difference normalized; LRE, long run emphasis; GLN, gray level nonuniformity; SRE, short run emphasis; *, statistically significant to distinguish any two pathologic subtypes.

Table S2 Interobserver agreement regarding texture features of GGO ROIs using interclass ICC

Texture features	ICC (95% CI)	Texture features	ICC (95% CI)
Gray levels	0.908 (0.100–0.981)	Autocorrelation	0.997 (0.261–0.999)
Voxel count	0.999 (0.595–1.000)	Cluster prominence	0.908 (0.010–0.981)
Energy	0.714 (0.002–0.928)	Cluster shade	0.948 (0.018–0.989)
Entropy	0.838 (0.005–0.964)	Cluster tendency	0.512 (0.001–0.844)
Minimum intensity	0.417 (0.001–0.787)	Contrast	0.851 (0.006–0.967)
Maximum intensity	0.999 (0.650–1.000)	Correlation	0.829 (0.005–0.962)
Mean intensity	0.834 (0.005–0.963)	Difference entropy	0.999 (0.641–1.000)
Median intensity	0.984 (0.056–0.997)	Dissimilarity	0.998 (0.362–1.000)
Range	0.968 (0.029–0.994)	Homogeneity 1	0.992 (0.110–0.998)
Mean deviation	0.556 (0.001–0.866)	Homogeneity 2	0.998 (0.393–1.000)
Root mean square	0.721 (0.003–0.930)	IMC1	0.829 (0.005–0.962)
Standard deviation	0.707 (0.002–0.926)	IDMN	0.832 (0.005–0.962)
Skewness	0.952 (0.019–0.990)	IDN	0.997 (0.221–0.999)
Kurtosis	0.741 (0.003–0.937)	Inverse variance	0.996 (0.215–0.999)
Variance	0.998 (0.367–1.000)	Maximum probability	0.999 (0.425–1.000)
Uniformity	0.910 (0.010–0.981)	Sum average	0.999 (0.564–1.000)
Volume (mm ³)	0.999 (0.472–1.000)	Sum entropy	0.996 (0.198–0.999)
Surface area (mm ²)	0.994 (0.150–0.999)	Sum variance	0.886 (0.008–0.976)
Surface: volume ratio	0.999 (0.537–1.000)	Energy (GLCM)	0.999 (0.596–1.000)
Compactness 1	0.807 (0.004–0.956)	Entropy (GLCM)	0.998 (0.354–1.000)
Compactness 2	0.999 (0.602–1.000)	Variance (GLCM)	0.965 (0.027–0.993)
Spherical disproportion	0.832 (0.005–0.962)	RLN	0.999 (0.498–1.000)
Sphericity	0.893 (0.008–0.977)	RP	0.851 (0.006–0.967)
Maximum 3D diameter	0.528 (0.001–0.852)	LGLRE	0.991 (0.095–0.998)
Volume cc	0.900 (0.009–0.979)	HGLRE	0.948 (0.018–0.989)
Extruded surface area	0.984 (0.056–0.997)	SRLGLE	0.829 (0.005–0.962)
Extruded volume	0.968 (0.029–0.994)	SRHGLE	0.832 (0.005–0.962)
Extruded surface: volume ratio	0.720 (0.000–0.287)	Box-counting dimension	0.795 (0.004–0.952)
LRLGLE	0.982 (0.050–0.996)	LRHGLE	0.714 (0.002–0.928)
Correlation dimension	0.901 (0.009–0.979)	LRE	0.795 (0.004–0.952)
SRE	0.714 (0.002–0.928)	GLN	0.933 (0.014–0.986)

GGO, ground-glass opacity lesions; ROI, region of interest; ICC, correlation coefficient analysis; Volume cc, volume in cubic centimeters; LRLGLE, long run low gray level emphasis; SRE, short run emphasis; 95% CI, 95% confidence interval; IMC1, informational measure of correlation 1; IDMN, inverse difference moment normalized; IDN, inverse difference normalized; GLCM, gray-level co-occurrence matrix; RLN, run length nonuniformity; RP, run percentage; LGLRE, low gray level run emphasis; HGLRE, high gray level run emphasis; SRLGLE, short run low gray level emphasis; SRHGLE, short run high gray level emphasis; LRHGLE, long run high gray level emphasis; LRE, long run emphasis; GLN, gray level nonuniformity.