Development and validation of a nomogram based on CT texture analysis for discriminating minimally invasive adenocarcinoma from glandular precursor lesions in sub-centimeter pulmonary ground glass nodules

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Abstract. In a recent reclassification, adenocarcinoma in situ has been redefined as a glandular precursor lesion (GPL), alongside adenomatous hyperplasia. This updated classification necessitates corresponding adaptations in clinical diagnostic and therapeutic protocols. Consequently, the present study aimed to construct and validate a nomogram utilizing computed tomography (CT) texture features to effectively discriminate between minimally invasive adenocarcinoma (MIA) and GPL within sub-centimeter pulmonary ground glass nodules (GGNs). To achieve this objective, the present study employed rigorous statistical methodologies, including the Mann-Whitney U test and binary logistic regression analysis, to identify distinguishing features and establish predictive models. Subsequently, the diagnostic performance of these models underwent evaluation through receiver operating characteristic (ROC) curves. The area under the curve (AUC) in ROC curves was compared using DeLong's test. Additionally, the nomogram was constructed using R software and its diagnostic performance was validated through calibration curves. Within both the training and validation datasets, the AUCs were observed to be 0.992 [95% confidence interval (CI): 0.980-1.000] and 0.975 (95% CI: 0.935-1.000), respectively. DeLong's test revealed significant disparities in the

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AUCs between the nomogram and single-parameter models (P<0.001). Furthermore, calibration curves demonstrated concordance between the training and validation datasets. In conclusion, the application of a CT texture-based nomogram model has demonstrated aptitude in differentiating between MIA and GPL within sub-centimeter GGNs. This model streamlines the identification of optimal surgical interventions and enhances the sphere of clinical decision-making and management.

Introduction

The incidence and mortality rates of lung cancer surpass those of all other malignant tumors, accounting for ~20% of cancer-related deaths worldwide (1). Early diagnosis and treatment are pivotal in improving the prognosis and survival rates of lung cancer patients. In the 2021 WHO classification (2), adenocarcinoma in situ (AIS) has been reclassified as a glandular precursor lesion (GPL) along with adenomatous hyperplasia (AAH), while the subcategories of lung cancer now include minimally invasive adenocarcinoma (MIA) and invasive adenocarcinoma (IAC). This classification update necessitates corresponding adjustments to clinical diagnosis and treatment protocols. Currently, management follow-up strategies predominantly apply to AAH and AIS, whereas MIA warrants prompt surgical intervention (3,4). Furthermore, although some studies have indicated comparable long-term efficacy and 5-year survival rates between AAH, AIS and MIA surgeries, others have found that MIA exhibits higher Ki67 levels and EGFR mutation rates compared with AIS (5). Therefore, achieving precise discrimination of MIA from GPL during management follow-up and pre-surgical stages provides important insights for determining optimal clinical intervention timing, implementing surgical protocols and assessing prognosis.

Previous studies have demonstrated that high-resolution low-dose computed tomography (CT) is currently the most effective screening tool, reducing lung cancer mortality

Key words: computed tomography texture analysis, adenomatous hyperplasia, adenocarcinoma *in situ*, minimally invasive adenocarcinoma, nomogram

by 20% (6). Over 90% of these cases represent early-stage lung cancer, with pulmonary ground glass nodules (GGNs) being the primary manifestation on CT scans. While pathological biopsy serves as the gold standard for diagnosing lung cancer types, frozen section analysis has emerged as a valuable method for rapid intraoperative assessment of nodules, distinguishing between benign and malignant lesions and determining histological subtypes. This information plays a crucial role in guiding surgical strategies for lung nodules (7,8). However, due to the small size and low density of lung nodules, sampling often yields suboptimal results, resulting in a concordance rate of only ~68% between intraoperative frozen section analysis and postoperative paraffin pathology (8). Some researchers have explored the use of traditional CT imaging features (such as nodule size, density and solid component proportion) to differentiate between MIA and GPL (9,10). However, these features exhibit significant overlap across different nodule subtypes, leading to low diagnostic efficacy (11). Moreover, the extraction of these features heavily relies on the subjective interpretation skills and clinical experience of the radiologist (8). In addition, traditional techniques often struggle to identify nodules with smaller volumes and mixed densities. The increasing prevalence of GGNs with diameters <10 mm, detected through low-dose CT screening for early lung cancer, presents new challenges in clinical diagnosis. Some studies have proposed a critical value of 10 mm diameter for distinguishing between glandular precursor lesions and invasive lesions (9,10,12). Nevertheless, clinical practice has revealed numerous GGNs ≤ 10 mm confirmed as MIA or IAC (10), leading to continuing debates about the management strategies for GGNs of this size (13). Consequently, there is an urgent need for innovative and accurate techniques to enable the precise preoperative diagnosis of MIA and GPL.

Radiomics is a powerful technique that enables the extraction and analysis of numerous radiomics features from medical imaging data with high efficiency. This approach holds significant promise in distinguishing between different pathological subcategories of lung nodules, assessing the extent of infiltration and evaluating prognostic outcomes (14). Texture analysis, on the other hand, involves the extraction and quantitative analysis of non-macroscopic and deep-level CT image features that reflect the tumor's heterogeneity to some extent. It has demonstrated utility in various aspects, including tumor differential diagnosis, prognostic evaluation, treatment response prediction and monitoring (15-17). CT texture analysis is widely utilized for identifying benign and malignant solitary lung nodules and evaluating invasiveness, exhibiting exceptional performance (18,19). To date, there have been no reports on the application of texture analysis for the identification of MIA and GPL in sub-centimeter GGNs. Therefore, the present study aimed to develop and validate a nomogram based on CT quantitative parameters and texture features for improving the ability to discriminate MIA from AAH/AIS, thereby providing important guidance for formulating clinical treatment plans and optimizing the timing of surgical interventions.

Materials and methods

Patient and nodule selection. The present retrospective study received approval from the ethics committee and written

informed consent was waived. Patients with sub-centimeter GGNs who underwent high-resolution CT (HRCT) were enrolled at the First People's Hospital of Foshan between January 2019 and February 2022. The flowchart illustrating patient inclusion and exclusion is shown in Fig. 1, which was in accordance with previous studies (17,19,20). The inclusion criteria were as follows: i) GGNs with a maximum diameter ≤10 mm, evaluated using lung window settings [level: -600 Hounsfield units (HU); width: 1,500 HU]; ii) confirmation of AAH, AIS or MIA through surgical excision and pathology examination referring to the 2021 WHO classification (2); iii) availability of chest HRCT examination conducted within one month prior to surgical treatment; iv) absence of prior history of puncture, chemotherapy, or radiotherapy; v) absence of lung cancer or other malignant tumors. The exclusion criteria consisted of: i) inability to accurately delineate the region of interest due to the presence of artifacts or other lesions; ii) inaccurate identification or segmentation of CT images, including incomplete delineation or failure to avoid blood vessels, bronchi, pleura and other structures. The final selected cases were divided into two groups: 49 cases with AAH/AIS and 62 cases with MIA. The recruited patients were randomly assigned to a training set (78 nodules) and a validation set (33 nodules) at a ratio of 7:3.

CT image acquisition. The CT examinations for all patients included in the study were conducted using Philips CT scanners: the Philips Ingenuity 64-slice CT scanner (Philips Medical Systems, Inc.) and the Philips Brilliance iCT 256-slice CT scanner (Philips Ultrasound, Inc.). The patients were positioned in the supine (lying face-up) posture and the scanning range extended from the apex to the base of the lungs. Scans were performed at the end of a deep inspiration, followed by breath-holding to ensure stability. A tube voltage of 120 kV was used and the tube current employed automatic milliamp-second technology. The pitch value was set to 1.0, the collimation was 0.625x1.25 mm and the field of view (FOV) was set to 350x350 mm, with a pixel size of 512x512. The acquired images were reconstructed using both the standard algorithm and the high-resolution algorithm. The reconstructed slices had a thickness of 1 mm and a spacing of 1 mm.

CT image segmentation and feature extraction. The CT images in DICOM format were imported into the uAI-ChestCare software (version 0130; Shanghai United Imaging Healthcare Co., Ltd.), which was used for image segmentation and extraction of texture features (20,21). This software facilitated the automatic delineation of the complete 3D region of interest (ROI) for the identified lesions by outlining the tumor boundary on consecutive axial lung window images (with a window width of 1,500 HU and a window level of -600 HU). Subsequently, various quantitative and texture features were computed, encompassing maximum diameter (MD), solid volume (SV), solid volume rate (SVR), solid quality (SQ), solid quality rate (SQR), maximum computed tomography attenuation (CT_{max}), minimum computed tomography attenuation (CT_{min}), mean computed tomography attenuation (CT_{mean}), median computed tomography attenuation (CT_{median}), variance, kurtosis, skewness and entropy. To evaluate the accuracy of



Figure 1. Flowchart of patient selection. AAH, atypical adenomatous hyperplasia; AIS, adenocarcinoma *in situ*; MIA, minimally invasive adenocarcinoma; GPL, glandular precursor lesion; HRCT, high-resolution computed tomography.

the software's automated nodule delineation, two radiologists with over 10 years of experience in thoracic imaging diagnosis independently assessed the results. They excluded cases where the nodule contour was incomplete or where blood vessels, bronchi, pleura and similar structures were not properly avoided. In instances where disagreements arose regarding the exclusion of certain cases, they negotiated to precisely delineate nodule boundaries while avoiding the inclusion of bronchi, large vessels, vacuole and normal tissue beyond the pleura in accordance with previous studies (22,23). A senior physician with 15 years of experience in thoracic imaging diagnosis verified the preceding two radiologists' segmentation results and provided the final confirmation of the results (19). The CT quantitative and texture features extraction of AIS (Fig. 2A) and MIA (Fig. 2B) were taken as examples.

Nomogram model building. The clinical, CT quantitative and texture features of both the training and validation groups were subjected to analysis using the Mann-Whitney U test to identify the effective distinguishing features. Subsequently, binary logistic regression analysis was employed to develop both a single-parameter model and a combined multi-parameter nomogram model. The diagnostic performance of these prediction models was assessed through the construction of receiver operating characteristic (ROC) curves (24) and the comparison of area under the curve (AUC) of ROC curves was performed using DeLong's test (25). Furthermore, the construction of the nomogram was accomplished using the

rms package in the R software (version 4.0, R Foundation; http://www.Rproject.org). To evaluate the diagnostic performance of the nomogram prediction model, a calibration curve was employed.

Statistical analysis. The statistical analysis was conducted using SPSS version 20 software (IBM Corp.). Continuous variables were summarized as mean \pm standard deviation (SD) or median with the full range. Categorical variables were presented as frequencies and percentages. Differences in age and sex between the two groups were assessed using the independent sample t-test and the χ^2 test, respectively. To identify significant variables as predictive indicators of MIA the Mann-Whitney U test was applied based on the CT quantitative and texture features. Variables with a significance level of P<0.05 were selected as significant predictors for constructing both single-parameter and combined multi-parameter prediction models through binary logistic regression analysis. The diagnostic performance of each model was then evaluated by comparing the AUC, specificity and sensitivity of the ROC curve. AUC values greater than 0.5 were considered predictive. The significance of differences between ROC curves was determined using DeLong's test. Furthermore, the nomogram was constructed using the rms package in the R software. To verify the diagnostic performance of the nomogram prediction model, a calibration curve was employed. P<0.05 was considered to indicate a statistically significant difference.



Figure 2. CT images were used for accurate nodule delineation and quantitative texture feature extraction, complemented by corresponding hematoxylin-eosin stained pathological images at x40 magnification. (A) In the first case, a 69-year-old female had a 10x8 mm GGN in the upper lobe of the right lung with limited solid features. (B) In the second case, a 63-year-old female had a 9x7 mm GGN in the lower lobe of the right lung, mainly solid. CT, computed tomography; GGN, ground glass nodules; SV, solid volume; SVR, solid volume rate; SQ, solid quality; SQR, solid quality rate; CT_{max} , maximum computed tomography attenuation; HU, Hounsfield units.

Results

Patient characteristics. A total of 109 patients with 111 GGNs were ultimately included in the present study. The cohort consisted of 35 males and 74 females, with ages ranging 19-78 years. Among these patients, 49 GGNs in 47 patients were classified as GPL (AAH/AIS), with a mean age of (54.26 ± 12.24) years. Additionally, 62 GGNs in 62 patients were categorized as MIA, with a mean age of (52.84 ± 11.47) years. There were no significant differences observed in terms of sex (P=0.564) or age (P=0.522) between the two groups. Utilizing stratified random sampling, the training set consisted of 78 cases (34 AAH/AIS and 44 MIA), while the validation set included 33 cases (15 AAH/AIS and 18 MIA).

CT quantitative parameters and texture feature extraction. The results of patient characteristics, CT quantitative parameters and texture features were conducted using the Mann-Whitney U test and the results are presented in Table I. Notably, significant statistical differences (P<0.05) were observed between the two groups in seven parameters, including SV, SVR, SQ, SQR, CT_{max} , CT_{mean} and entropy.

Model construction and diagnostic validation based on CT quantitative parameters and texture features. The diagnostic models were constructed and the corresponding ROC curves are displayed in Fig. 3. Notably, the combined multi-parameter model exhibited superior predictive ability compared with each individual single-parameter model. The performance of the nomogram prediction model is illustrated in Table II for the training set and Table III for the validation set. In the training set, the nomogram achieved an AUC of 0.992 (95% CI: 0.980-1.000), a sensitivity of 0.907, a specificity of 1.000 and an accuracy of 0.948. For the validation set, the AUC was 0.975 (95% CI: 0.935-1.000), with a sensitivity of 0.842, a specificity of 0.941 and an accuracy of 0.912. The results of DeLong's test demonstrated significant statistical differences in AUCs between the nomogram and single-parameter models (P<0.001).

Nomogram and calibration curve. A nomogram (Fig. 4A) was constructed utilizing five parameters (SV, SVR, CT_{max} , CT_{mean} and entropy). The model formula was derived as follows: Total Points=0.576 x SV -1.131 x SVR - 0.162x CT_{max} - 0.136 x CT_{mean} + 21.533 x entropy. Each parameter in the nomogram corresponds to a specific score on the top 'Points' axis. The cumulative sum of these scores corresponds to the values displayed on the bottom 'Total Points' axis, representing the diagnostic probability of MIA: Two cases of AIS and MIA (Fig. 4B and C) provided illustrative examples from the collected database. The calibration curve (Fig. 5) demonstrates a favorable agreement between the predicted model and the observed data, as the scatter plot closely aligns with the ideal curve.

Discussion

The present study retrospectively collected surgically resected and pathologically confirmed cases of GPL and MIA that presented as sub-centimeter GGNs on CT images, then extracted and analyzed their CT quantitative and texture features. Subsequently, a nomogram model incorporating the five most informative identification indicators (SV, SVR,

Table I. Clinical characteristics, CT quantitative parameters and texture features between GPL and MIA groups.						
Clinical characteristic	AAH/AIS (n=49)	MIA (n=62)	P-value			
Age, years, mean (SD)	54.26 (12.24)	52.84 (11.47)	0.646ª			
Sex			0.521 ^b			
Female, n (%)	33 (70.2)	41 (66.1)				
Male, n (%)	14 (29.8)	21 (33.9)				
MD, mm, median (range) ^c	7.60 (5.02-10.00) ^c	8.20 (4.24-9.87)	0.080			
SV, mm ³ , median (range) ^c	59.00 (11.00-128.00)	87.00 (0.00-156.00)	<0.001 ^d			
SVR, %, median (range) ^c	38.00 (8.00-66.00)	45.00 (0.00-68.00)	0.021^{d}			
SQ, mg, median (range) ^c	49.00 (7.00-110.00)	72.00 (0.00-128.00)	0.001 ^d			
SQR, %, median (range) ^c	29.00 (6.00-66.00)	47.00 (0.00-66.00)	0.001^{d}			
CT _{max} , HU, median (range) ^c	-151.50 (-270.00-102.00)	-111.00 (-282.00-62.00)	0.034^{d}			
CT _{min} , HU, median (range) ^c	-784.50 (-915.00558.00)	-788.0 (-1007.00582.00)	0.935			
CT _{mean} , HU, median (range) ^c	-584.50 (-709.00348.00)	-530.00 (-737.00399.00)	0.002^{d}			
CT _{median} , HU, median (range) ^c	-588.50 (-710.00368.00)	-546.00 (-730.00419.00)	0.091			
Variance, median (range) ^c	137.00 (92.00-187.00)	139.00 (84.00-194.00)	0.429			
Kurtosis, median (range) ^c	3.50 (2.00-4.80)	3.30 (1.90-4.90)	0.797			
Skewness, median (range) ^c	0.39 (-0.19-0.86)	0.48 (-0.40-0.78)	0.327			

Tabl

Entropy, median (range)^c

^at-test; ^b χ^2 test; ^ccontinuous variables were represented as median (minimum, maximum); ^dP<0.05 ndicates a statistically significanct difference. HU, Hounsfifield unit; SD, stander deviation; MD, maximum diameter; SV, solid volume; SVR, solid volume rate; SQ, solid quality; SQR, solid quality rate; CT_{max}, maximum computed tomography attenuation; CT_{min}, minimum computed tomography attenuation; CT_{mean}, mean computed tomography attenuation; CT_{median}, median computed tomography attenuation.

5.50 (3.60-7.30)

Table II. The AUC values for each prediction model were calculated and evaluated on both the training and validation set.

Clinical characteristic	Training set		Validation set	
	AUC (95% CI)	P-value	AUC (95% CI)	P-value
SV, mm ³)	0.735 (0.622-0.848)	<0.001ª	0.547 (0.347-0.748)	0.640
SVR, %)	0.654 (0.529-0.779)	0.021ª	0.540 (0.331-0.750)	0.690
SQ, mg	0.732 (0.620-0.843)	0.001 ^a	0.509 (0.303-0.715)	0.931
SQR, %	0.729 (0.611-0.847)	0.001ª	0.523 (0.321-0.725)	0.822
CT _{max} , HU	0.641 (0.513-0.769)	0.034 ^a	0.575 (0.376-0.775)	0.456
CT _{mean} , HU	0.706 (0.577-0.834)	0.002 ^a	0.509 (0.305-0.721)	0.931
entropy	0.752 (0.640-0.866)	<0.001 ^a	0.521 (0.318-0.724)	0.835
nomogram	0.992 (0.980-1.000)	<0.001 ^a	0.975 (0.935-1.000)	<0.001 ^a

^aP<0.05 indicates a statistically significanct difference. AUC, area under the curve; CI, confidence interval; HU, Hounsfifield unit; SV, solid volume; SVR, solid volume rate; SQ, solid quality; SQR, solid quality rate; CT_{max}, maximum computed tomography attenuation; CT_{mean}, mean computed tomography attenuation.

 $\text{CT}_{\text{max}},\,\text{CT}_{\text{mean}}$ and entropy) was developed to accurately differentiate MIA from GPL. The ROC curve revealed high predictive accuracy in distinguishing between MIA and GPL, with an AUC values of 0.992 and 0.975 for the training set and validation set, respectively. Early-stage lung cancers predominantly present as solitary GGNs, posing challenges in distinguishing benign nodules from malignant ones due to the diverse manifestations and overlapping features observed on CT images. In line with previous studies by Wu et al (9,10), who extensively analyzed the CT imaging features of sub-centimeter pure GGNs, the present study identified relevant morphological features such as lesion size, vessel changes and tumor-lung interface, which reflect the invasiveness of GGNs. Moreover, previous investigations (26-28) have established a close association between the size of the solid component and the average CT attenuation of GGNs with their invasiveness and pathology. The findings of the present study corroborated these conclusions, demonstrating significant differences in SV, SVR, SQ, SQR, CT_{max} and CT_{mean} between GPL and MIA. Notably, the values of these variables

6.30 (3.20, 8.00)

<0.001^d

Set	AUC	95% CI	Sensitivity	Specificity	Accuracy
Training	0.992	0.980-1.000	0.907	1.000	0.948
Validation	0.975	0.935-1.000	0.842	0.941	0.912

Table III. Performance of nomogram model on the training and validation set.

AUC, area under the curve; CI, confidence interval.



Figure 3. ROC curves evaluated model performance in both the training and validation sets. The nomogram outperformed all other models, with an AUC of 0.992 in the training set and 0.975 in the validation set. ROC, receiver operating characteristic; AUC, area under the curve; SV, solid volume; SVR, solid volume; SVR, solid volume; ate; SQ, solid quality; SQR, solid quality rate; CT_{max}, maximum computed tomography attenuation; CT_{mean}, mean computed tomography attenuation.

were consistently higher in the MIA group compared with the GPL group. These results confirm the importance of the solid component in predicting the behavior of GGNs. To summarize, the present study retrospectively analyzed surgically resected GGNs, extracting and analyzing their CT quantitative and texture features and achieved high predictive performance in discriminating between MIA and GPL through the development of a nomogram model incorporating five informative identification indicators. The challenges associated with differentiating benign and malignant nodules in early-stage lung cancers were underscored, emphasizing the significance of considering the presence and characteristics of the solid component in predicting invasiveness and pathological behavior of GGNs.

Texture analysis plays a crucial role in efficiently and accurately extracting biological information that reflects tumor heterogeneity, which may not be discernible by visual examination of images alone. It enables quantitative assessment of subtle changes in image pixel values and their arrangement. As a result, it holds great value in qualitative diagnosis, invasiveness assessment, prognostic prediction of tumors and informed clinical management. Qiu et al (17) demonstrated the independent prognostic significance of mean CT attenuation and entropy in evaluating the invasiveness of 428 cases of clinical stage IA lung adenocarcinoma. Zhu et al (19) employed the ANOVA test and the least absolute shrinkage and selection operator algorithm to identify 18 CT texture features, including entropy. They successfully developed a diagnostic model capable of distinguishing MIA from GPL presenting as pure GGNs, achieving high identification performance with an AUC of 0.884 in the training set and 0.872 in the validation set. In the present study, there



Figure 4. The nomogram helps differentiate MIA from AIS/AAH. (A) The 'Total Points' on the nomogram predicts the chance of MIA. (B) An AIS case had a 'Total Points' value of 193.588, meaning a MIA probability less than 0.1. (C) An MIA case has a 'Total Points' value of 202.369, corresponding to a MIA probability greater than 0.9. MIA, minimally invasive adenocarcinoma; AIS, adenocarcinoma *in situ*; AAH, adenomatous hyperplasia; SV, solid volume; SVR, solid volume rate; SQ, solid quality; SQR, solid quality rate; CT_{max} , maximum computed tomography attenuation; CT_{mean} , mean computed tomography attenuation.



Figure 5. The calibration curve of the nomogram model was created and evaluated in both the training and validation sets.

was a statistically significant difference in entropy between GPL and MIA, which is consistent with previous findings. Moreover, the combined multi-parameter model exhibited superior predictive performance, with AUC values of 0.992 in the training set and 0.975 in the validation set, surpassing the research conducted by Zhu et al (19). This disparity may be attributed to differences in the composition of enrolled cases. The present study encompassed not only pure vitreous nodules but also mixed-density nodules. Previous studies (26,28) have indicated the diagnostic relevance of the solid component within GGNs. Another study (29) employed machine learning and deep learning methods to evaluate benign and malignant pulmonary nodules, revealing AUC values of 0.763 for the support vector machine (SVM) model and 0.723 for the convolutional neural networks (CNN) model in distinguishing GPL from MIA. Notably, the predictive performance of both SVM and CNN models was lower than that of the nomogram model established in the present study. These findings suggest that CT texture analysis may offer greater assistance and efficiency in achieving this objective. The patients in the present study were sourced from a single center, resulting in a relatively small sample size, but the parameters and quality of these CT images were highly standardized and homogenized. Conversely, in a previous multicenter study (29), although the sample size was larger, challenges related to CT image homogenization may have persisted. This discrepancy could explain the superior diagnostic effectiveness of the constructed model. Consequently, future studies should focus on expanding the sample size and analyzing data from different centers to obtain a more accurate and efficient predictive model.

Nomogram models are graphical representations employed to illustrate analytical outcomes derived from multifactorial logistic regression models or Cox proportional hazards models. These models use a set of parallel, non-intersecting lines within a coordinate plane to portray the quantitative analysis diagram, depicting the functional relationship between multiple variables. By employing intuitive symbols, nomograms facilitate the calculation of disease occurrence probability, recurrence risk and prognosis. Consequently, they find extensive application in clinical research pertaining to pulmonary GGNs (30). In the present study, a nomogram model for the identification of precursor lesions and MIA was established, based on five CT quantitative and texture features, namely SV, SVR, CT_{max} , CT_{mean} and entropy. This model serves as an effective and intuitive reference standard, enabling radiologists to achieve accurate and prompt diagnoses.

In conclusion, the use of CT quantitative and texture features offered significant utility in the differentiation of MIA from GPL. The nomogram model developed demonstrated superior discriminatory capabilities. This model's diagnostic efficacy, combined with its graphical representation, facilitates the precise classification of GGN types by radiologists, thereby aiding clinicians in making informed decisions regarding treatment and follow-up strategies for GGNs. Nonetheless, the present study has certain limitations that require acknowledgment. It was a retrospective, single-center study with a small sample size, which may introduce selection bias and potentially affect the precision and applicability of the model. In the future, efforts should be made to enhance the model's robustness and applicability by increasing the sample size, conducting multi-center studies and incorporating external test sets.

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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

CL and ZX participated in the design of the study. YJ and QD performed data analysis and prepared the figures. YY, RD and JZ participated in the analysis of the figures and data. CL and YJ prepared and revised the manuscript. AP, MG and ZX reviewed the results and revised the manuscript. CL, AP, MG and ZX confirm the authenticity of all the raw data. All authors have read and approved the final version of this manuscript.

Ethics approval and consent to participate

The present study was approved by the Institutional Research Ethics Committee of the First Peoples' Hospital of Foshan (approval no. 2021.02) and was conducted in accordance with the ethical principles of the Declaration of Helsinki and the ethical laws and regulations established in China. Written informed consent was waived.

Patient consent for publication

Not applicable.

Use of artificial intelligence tools

The uAI-ChestCare software is an artificial intelligence (AI) software based on deep learning method, specifically designed for applications related to chest or pulmonary care and have achieved good results in prior research endeavors. In the present study, CT lung window images, configured with a window width of 1,500 Hounsfield Units (HU) and a window level of -600 HU in DICOM format, were input into the uAI-ChestCare software. Subsequently, the software autonomously delineated the entire 3D ROI for identified lesions by outlining the tumor boundaries across consecutive axial slices. Following this, an array of quantitative and textural features were computed, encompassing parameters such as MD, SV, SVR, SQ, SQR, maximum computed tomography attenuation CT_{max} , CT_{min} , CT_{mean} , CT_{median}), variance, kurtosis, skewness and entropy.

Competing interests

The authors declare that they have no competing interests.

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