

Deep learning-assisted development and validation of an algorithm for predicting the growth of persistent pure ground-glass nodules

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Background: The prediction of the persistent pure ground-glass nodule (pGGN) growth is challenging and limited by subjective assessment and variation across radiologists. A chest computed tomography (CT) image-based deep learning classification model (DLCM) may provide a more accurate growth prediction.

Methods: This retrospective study enrolled consecutive patients with pGGNs from January 2010 to December 2020 from two independent medical institutions. Four DLCM algorithms were built to predict the growth of pGGNs, which were extracted from the nodule areas of chest CT images annotated by two radiologists. All nodules were assigned to either the study, the inner validation, or the external validation cohort. Accuracy, sensitivity, specificity, receiver operating characteristic (ROC) curves, and areas under the ROC curve (AUROCs) were analyzed to evaluate our models.

Results: A total of 286 patients were included, with 419 pGGN. In total, 197 (68.9%) of the patients were female and the average age was 59.5±12.0 years. The number of pGGN assigned to the study, the inner validation, and the external validation cohort were 193, 130, and 96, respectively. The follow-up time of stable pGGNs for the primary and external validation cohorts were 3.66 (range, 2.01–10.08) and 4.63 (range, 2.00–9.91) years, respectively. Growth of the pGGN occurred in 166 nodules [83 (43%), 39 (30%), and 44 (45%) in the study, inner and external validation cohorts respectively]. The best-performing DLCM algorithm was DenseNet_DR, which achieved AUROCs of 0.79 [95% confidence interval (CI): 0.70, 0.86] in predicting pGGN growth in the inner validation cohort and 0.70 (95% CI: 0.60, 0.79) in the external validation cohort.

Conclusions: DLCM algorithms that use chest CT images can help predict the growth of pGGNs.

Keywords: Pure ground-glass nodules (pGGNs); deep learning classification model (DLCM); predicting model; follow-up; growth

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Introduction

Lung cancer is the leading cause of cancer mortality worldwide (1). Extensive use of high-resolution computed tomography (CT) has resulted in frequent identification of ground-glass nodules (GGNs) (2,3). Most persistent pure GGNs (pGGNs) represent early-stage lung cancers; in particular, persistent lung GGNs may denote atypical adenomatous hyperplasia, adenocarcinoma in situ (AIS), minimally invasive adenocarcinoma (MIA), and invasive adenocarcinoma (IAC) (4,5). However, pGGNs include some rare cases of metastasis of gastric, colorectal, pancreatic, breast, and cholangiocarcinoma (6-10). Therefore, the careful follow-up and the early detection of the growth of tumor is important.

Although pGGNs have similar characteristics on CT imaging, their biological characteristics vary (11,12). Most pGGNs remain stable for long periods, whereas others are more invasive and prone to rapid growth (4,5,13). Therefore, most clinicians adopt a wait-and-see strategy incorporating regular follow-up. Follow-up has a crucial role in clinical decision-making and determining whether surgery is indicated and is therefore increasingly recommended by thoracic and pulmonary specialists. As with most guidelines, the Fleischner Society Guidelines for the management of pGGNs recommend follow-up intervals ranging from 3 to 6 months initially depending on the nodule characteristics, then every 2 years for a total of 5 years, if stable, and surgery if the pGGN or the solid

Highlight box

Key findings

• Deep learning (DL)-assisted algorithm could predict the growth of persistent pure ground-glass nodules (pGGNs).

What is known and what is new?

- DL algorithm has been used to diagnose pulmonary nodules.
- We developed and validated a new algorithm to predict growth of pGGNs.

What is the implication, and what should change now?

• By our algorithm, we could classify high and low risk pGGNs, which may allow clinicians to develop earlier treatment strategies for patients.

component grows (14). The increased number of detected pGGNs requiring the long-term follow-up results several problems, such as the economic burden, overuse of medical resources, and overexposure to radiation. Therefore, it is necessary and reasonable to extend the follow-up intervals once stable pGGNs have been identified.

Most previous studies have used quantitative CT imaging to determine the size, density, and volume of GGNs and have estimated their invasiveness based on these characteristics (4,5). However, both the assessment of these characteristics and forecasting of pGGN growth have been inaccurate (3,13).

Artificial intelligence in the form of automatic quantitative chest CT imaging has recently been shown to reliably diagnose pulmonary nodules and accurately predict lymph node metastasis (15-18). Artificial intelligence has proven so versatile and efficient in developing deep learning (DL) algorithms from otherwise uninterpretable neural network systems and medical imaging findings that it has been dubbed "Black Box" medicine. The present study aimed to construct and validate a DL classification model (DLCM) based on chest CT imaging findings for predicting the growth of pGGNs. We present this article in accordance with the TRIPOD reporting checklist (available at https://tlcr.amegroups.com/article/view/10.21037/tlcr-23-666/rc).

Methods

Study design and patients

The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013). This twoinstitution (Institution 1: Beijing Chaoyang Hospital; Institution 2: People's Liberation Army General Hospital), retrospective cohort study was approved by the institutional review board of Beijing Chaoyang Hospital (IRB No. 2021ke-135), and written informed consent for this retrospective analysis was waived. The People's Liberation Army General Hospital was informed and agreed with this study. None of the patients who participated in this study were previously reported. We initially collected consecutive patients with at least two chest-CT scans for pGGNs performed at Institution 1 or Institution 2 from January 2010 to



Figure 1 Cohort selection flowchart. Flow diagram shows the method for persistent pGGN patient inclusion and exclusion. CT, computed tomography; pGGN, pure ground-glass nodule.

December 2020.

The inclusion criteria were as follows: (I) patients with persistent stable pGGNs and a follow-up of ≥ 2 years and (II) growing GGNs, no limitation of the follow-up duration. The exclusion criteria were as follows: (I) a confirmed solid nodule on initial chest CT image; (II) transient pGGNs; (III) disappeared pGGNs and suspicion of interstitial lung disease, fibrotic changes, or bronchiolitis; and (IV) cancer history. A flowchart of participant enrollment is shown in *Figure 1*.

pGGN identification, growth analysis, follow-up time calculation, and region of interest (ROI) placement

pGGNs were identified by one reviewer at two institutions (X.T. and Y.T., 4/12 years of experience of chest images)

who was not involved in image interpretation. Consecutive institutional retrospective searches of imaging databases were performed on or before May 31, 2021.

To assess intraobserver reproducibility, two radiologists reviewed all patients' CT images (initial and last) twice in 1 week [Reader 1 (Tang Y, 12 years of experience of chest images) and Reader 2 (Jin X, 12 years of experience chest images)] and determined whether each pGGN had grown or was stable. Interobserver (Reader 1 vs. Reader 2) reproducibility concerning the status of all pGGNs was also evaluated. The nodules were defined as having grown only when both readers had reached this conclusion twice. Interand intraclass correlation coefficients were used to evaluate the agreement between judging that growth had occurred by Reader 1 (twice) and by Reader 2 (twice).



Figure 2 Overview of DLCM predicting pulmonary nodules. CT, computed tomography; DLCM, deep learning classification model.

The follow-up time for stable pGGNs was calculated as the interval between the initial and last CT scan. For pGGNs that had grown, the follow-up time was defined as the interval between the initial CT scan and the CT scan in which growth was identified. ROIs the whole pGGNs of all layers, from pGGNs were first segmented by a radiologist (X.T.) by ITK-SNAP software (version 3.6; Penn Image Computing and Science Laboratory at the University of Pennsylvania, Philadelphia, PA, USA; http://www.itksnap. org/pmwiki/pmwiki.php) and later modified by another radiologist (X.J.).

Dataset establishment

All patients in Institution 1 were randomly allocated to the study cohort or inner validation cohorts at a ratio of 4:1, keeping the ratio of stable and growth pGGN data in these cohorts at 1:1. Patients in Institution 2 were all assigned to an independent external validation cohort. In patients who had multiple pGGNs, 40×40 pixel areas of each nodule on each lung CT were extracted separately. Subgroups of patients with pGGNs that had grown in the inner validation cohort were formed based on the duration of follow-up [0–1 year follow-up (D0–1), 1–2 years follow-up (D1–2), and more than 2 years follow-up (D2+)].

Model development, evaluation, and interpretability

We implemented four DLCMs, including Inceptionv3 (19), ResNet18 (20), DenseNet121 (21), and DenseNet_DR. All DLCMs were trained to predict whether pGGNs would grow (22).

We quantified all the DLCM predictive performances

by using the receiver operating characteristic (ROC) curve and the area under the ROC curve (AUROC) in the inner validation and external validation cohorts. Moreover, we calculated each model's AUROC for the following subgroups in the inner validation cohort: D0–1, D1–2, and D2+.

In the testing phase, the DLCM predicted all images of each pGGN and calculated the average predicted probability to obtain the prediction of the growth state of nodules, as shown in *Figure 2*. By comparing the predictive performance of all models, the model with the best predictive performance was established as the final model in this study. In-depth information about the selected model development is provided in the supplementary material.

Score-CAM (23) was used to interpret the DLCM. Score-CAM incorporated an increase in confidence in the design of weight for each activation to eliminate the dependence on gradients. The specific process is described in *Figure 3*.

Statistical analysis

Statistical analyses were conducted with R Studio (version 1.3.959; R Project for Statistical Computing, Vienna, Austria; https://www.r-project.org) and Python (version 3.6; Bioinformatics Lab at University of Ljubljana, Ljubljana, Slovenia; https://www.python.org/downloads/ release/python-360/), with P<0.05 considered statistically significant. The data are expressed as the means \pm standard deviations (SDs) for continuous variables if normally distributed and percentages for categorical variables. Clinical data were analyzed using the Pearson χ^2 test for categorical data and the Wilcoxon rank sum test for continuous data. Kappa consistency analysis was used for intra- and intergroup consistency analysis.



Figure 3 Overview of the interpretable analysis of Score-CAM algorithm. Feature maps were first extracted by DenseNet_DR and worked as masks on the original CT image. The result could be generated by a linear combination of feature weights and feature maps. CT, computed tomography.

Results

Clinical characteristics

We enrolled 286 eligible patients, who had 419 pGGNs, including 253 persistent stable pGGNs and 166 with pGGNs that had grown (Institution 1, primary cohort, 206 patients, 323 pGGNs; Institution 2, external validation cohort, 80 patients, 96 pGGNs). The primary cohort comprised 58 males and 148 females with a mean age of 59.53 (range, 18–93) years, and the external validation cohort comprised 31 males and 49 females with a mean age of 59.26 (range, 33–83) years.

pGGN growth analysis, follow-up time calculation, and ROI placement

Dataset establishment

Among all 419 pGGNs, 168 were classified as growth pGGNs by Reader 1 and 166 by Reader 2; 251 were classified as stable pGGNs by Reader 1 and 253 by Reader 2. In total, two cases with different growth statuses were finally identified as stable. Satisfactory interobserver and intraobserver reproducibility regarding whether growth had occurred was achieved, with the intraobserver and interobserver agreement of 0.96, 0.97, and 0.99, respectively. The follow-up times of stable pGGNs for the primary and external validation cohorts were 3.66 (range, 2.01–10.08) and 4.63 (range, 2.00–9.91) years, respectively.

The maximum diameter, volume, and density of all

pGGNs were calculated by the mask placement of the ROI on ITK-SNAP software. The growth pGGN with bigger diameter (8.23 vs. 13.06 mm, P<0.001), larger size (208.52 vs. 830.55 mm³, P<0.001) had a higher CT value [-672.59 vs. -615.89 Hounsfield unit (HU), P<0.001] than the stable pGGN group, similar as previous study (24). Patient characteristics in the two institutions and pGGNs in the study cohort, inner validation cohort, and external validation cohorts are listed respectively in *Tables 1,2*.

Model establishment and prediction in the internal validation cohort

Four models were used to predict the growth of pulmonary nodules: Inceptionv3 (19), ResNet18 (20), DenseNet121 (21), and DenseNet_DR. The performance of the four feature extractors was as follows: the AUROC was 0.54 for Inceptionv3, 0.63 for ResNet18, 0.76 for DenseNet121, and 0.79 for DenseNet_DR (*Figure 4A*). The performance of DenseNet_DR was significantly better than that of the other model and established as the predicting model in our study. The detailed number of nodules images used in the training and validation set is provided in the supplementary material.

Prediction in subgroups, in external validation cobort, and interpretability of model prediction decision

The use of DenseNet_DR for predicting the growth of pulmonary nodules was more accurate in the D0-1

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Table 1 Th	e patients-based	clinicopathologic	characteristics
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Characteristics	Institution 1 (n=206)	Institution 2 (n=80)	Total (n=286)	P value
Age (years)	59.5±12.7	59.3±10.0	59.5±12.0	0.77
Gender				0.08
Male	58 (28.2)	31 (38.8)	89 (31.1)	
Female	148 (71.8)	49 (61.3)	197 (68.9)	
Surgery				<0.01
No surgery	180 (87.4)	37 (46.3)	217 (75.9)	
Surgery	26 (12.6)	43 (53.8)	69 (24.1)	
Multiple/solitary nodules				0.01
Multiple nodule	48 (23.3)	8 (10.0)	56 (19.6)	
Solitary nodule	158 (76.7)	72 (90.0)	230 (80.4)	
Nodule number	1.6±1.6	1.2±0.7	1.5±1.4	0.01
CT follow-up times, year	4.2±2.4	3.9±1.8	4.1±2.3	0.53

Values are expressed as n (%) or mean ± SD. Institution 1: Beijing Chaoyang Hospital; Institution 2: People's Liberation Army General Hospital. CT, computed tomography; SD, standard deviation.

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Characteristics	Study cohort (n=193)	Inner validation cohort (n=130)	External validation cohort (n=96)	Total (n=419)	P value
Follow-up (years)					
Stable pGGNs	3.59±1.68, n=110	3.84±1.41, n=91	4.66±2.25, n=52	3.90±1.76, n=253	<0.01
Growth pGGNs	2.80±1.98, n=83	2.19±2.40, n=39	3.46±2.29, n=44	2.83±2.20, n=166	0.03
Diameter (mm)	9.4±4.1	10.7±15.0	10.8±3.9	10.1±9.0	<0.01
Volume (mm ³)	452.0±1,165.3	417.2±637.4	511.9±793.4	511.9±793.4	<0.01
CT value (HU)	-654.3±107.6	-654.4±109.7	-635.9±108.3	-635.9±108.3	0.14
Location					0.28
Left upper lobe	42 (21.8)	33 (25.4)	26 (27.1)	101 (24.1)	
Left lower lobe	19 (9.8)	23 (17.7)	7 (7.3)	49 (11.7)	
Right upper lobe	90 (46.6)	54 (41.5)	44 (45.8)	188 (44.9)	
Right middle lobe	14 (7.3)	5 (3.8)	7 (7.3)	26 (6.2)	
Right lower lobe	28 (14.5)	15 (11.5)	12 (12.5)	55 (13.1)	
Cell type					<0.01
AIS	4 (2.1)	4 (3.1)	4 (4.2)	12 (2.9)	
MIA	1 (0.5)	1 (0.8)	10 (10.4)	12 (2.9)	
IAC	9 (4.7)	7 (5.4)	31 (32.3)	47 (11.2)	
No surgery	179 (92.7)	118 (90.8)	51 (53.1)	51 (53.1)	

Values are expressed as n (%) or mean ± SD. pGGN, pure ground-glass nodule; CT, computed tomography; HU, Hounsfield unit; AIS, adenocarcinoma in situ; MIA, minimally invasive adenocarcinoma; IAC, invasive adenocarcinoma; SD, standard deviation.



Figure 4 The ROC curve of nodules in the inner validation cohort, D0–1, D1–2, and D2+ predicted by the DLCM. The ROC curve in the inner validation cohort of all nodules (A), D0–1 (B), D1–2 (C), and D2+ (D) as predicted by DLCM. AUROC, area under the ROC curve; ROC, receiver operating characteristic; D0–1, 0–1 year follow-up; D1–2, 1–2 years follow-up; D2+, more than 2 years follow-up; DLCM, deep learning classification model.

Table 3 The	performance	of DLCM or	1 internal	validation	cohort and	subgroups
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Subgroup	AUC	ACC	SENS	SPEC	PPV	NPV
Internal validation cohort	0.787	0.727	0.719	0.730	0.489	0.878
D0-1	0.855	0.743	0.813	0.731	0.351	0.956
D1-2	0.723	0.711	0.500	0.730	0.143	0.942
D2+	0.715	0.732	0.750	0.730	0.200	0.970

DLCM, deep learning classification model; AUC, area under the curve; ACC, accuracy; SENS, sensitivity; SPEC, specification; PPV, positive predictive value; NPV, negative predictive value; D0–1, 0–1 year follow-up; D1–2, 1–2 years follow-up; D2+, more than 2 years follow-up.

(AUROC =0.855, *Figure 4B*) than in the D1-2 (AUROC =0.723, *Figure 4C*), and D2+ (AUROC =0.715, *Figure 4D*) subgroups (*Table 3*). DenseNet_DR achieved an

AUROC of 0.70 [95% confidence interval (CI): 0.60, 0.79, *Figure 5*] in the external validation cohort. DenseNet_DR performed worse in the external validation cohort than in



Figure 5 The ROC curve of all nodules in the external validation cohort, with AUROC =0.704. AUROC, area under the ROC curve; ROC, receiver operating characteristic.

the internal validation cohort, but the AUROC was similar to that for the D2+ subgroup. Feature-weighted activation maps overlaid with CT images showed that DenseNet_ DR focused its attention on the pulmonary nodules and periphery of nodules (*Figure 6*).

Discussion

In our study, the DLCM accurately predicted the growth of pGGNs in the inner validation cohort (AUROC =0.79) and somewhat less accurately in the external validation cohort (AUROC =0.70). The performance of our proposed DLCM indicates that it could provide an alternative to current prediction methods, which use long-term serial CT findings to assess growth risk in patients with persistent pGGNs.

In this study, the prediction accuracy of subgroup D0–1 was superior to that of the other two subgroups (D0–1 vs. D1–2, D2+; AUROC, 0.855 vs. 0.723, 0.715). The pGGNs can be stable for a long time, and their identifying characteristics may be most obvious when they become unstable or about to become unstable within one year of their identification (4,13,25-29). The CT image characteristics of pGGNs that remain stable over long periods before eventually growing probably differ considerably from the CT characteristics of those that grow soon after detection (5,11,30). This may explain why prediction is more accurate in pGGN growth within a short

follow-up than within a long follow-up.

The diagnostic performance of DLCM was lower in the external validation cohort than in the inner validation cohort (AUROC, 0.70 vs. 0.79) but was nonetheless encouraging. Explanations for this discrepancy were different follow-up times and different image parameters between the two cohorts. The average follow-up time of growth pGGNs for the external validation cohort (3.46 years) was longer than for the inner validation cohort (2.19 years), and the diagnostic performance of DLCM is better in pGGN growth within short follow-ups. Differences in image conditions, such as window width and window level, were observed between the inner and external validation cohorts. The window width and level were mostly 1,800 and 800 HU, respectively, in the inner cohort, whereas they were 1,500 and 800 HU, respectively, in the external cohort. The different parameters for CT imaging cause ineradicable differences in baselines for all images inputted into the DLCM. What's more, the follow-up protocol is not rigid and it is determined by each physician according to the preference.

In this study, we chose 2 years of follow-up as the cutoff for defining stable pGGNs, which is one of the limitations of this study. Despite this, most growth of pGGNs occurs within 2 years (25.5–51.9%) or 3 years (41.0–86.0%) of its detection reported in previous studies (11,27,29-35). In the current study, 157 (62.1%) stable pGGNs were followed up for more than 3 years.

In previous studies, the growth of pGGNs had been defined as at least a 20% increase in volume or ≥ 2 mm increase in diameter (26,36-38) between follow-ups. However, it is inaccurate in measurement because of the partial volume effect, especially for small pGGNs, which would cause classification errors. In this study, we minimized classification error by defining growth based on independent assessments by two radiologists rather than based on simple increases in volume or diameter. There was 0.99 agreement between the two radiologists, demonstrating high consistency and low classification error.

Most previous studies have excluded patients with multiple pGGNs or chose the largest pGGN to represent all pGGNs in a single individual (5,13,39,40). In the present study, we included multiple pGGNs and evaluated them independently for two reasons. First, multiple pGGNs comprise approximately 35% of newly discovered pGGNs (27,29,41,42). It would therefore result in a large selection bias if we excluded multiple pGGNs. Second, in patients with multiple pGGNs, the largest lesions and those with a



Figure 6 Representative CT images overlaid with heatmaps for model interpretation from the primary cohort. (A) A CT image of pGGN and (B) a heatmap of pGGN. Pulmonary nodules and their periphery were activated by the CT-based DLCM. CT, computed tomography; pGGN, pure ground-glass nodule; DLCM, deep learning classification model.

high risk of growth are recommended to be simultaneously resected. Therefore, it is important to identify each growing pGGN among multiple pGGNs.

We acknowledge that this study had limitations. First, the CT scan protocol and follow-up times of enrolled patients were heterogeneous because of the nature of the retrospective study. Second, 24.48% of all patients underwent surgical resection, and only 29.8% with growing pGGNs patients were underwent survival resection, which means that most pGGNs were not pathologically diagnosed. What's more, the rate of surgical procedure is different between inner cohort and external validation cohort. Third, the follow-up period for stable pGGNs was not long enough, which may have resulted in bias.

Conclusions

In conclusion, a DLCM that directly analyzes pGGN identified on chest CT images could facilitate the prediction of the growth of pGGNs. Considering all the limitations and the potential sources of bias, we believe further research and a larger database are required to validate and optimize our model.

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Footnote

Reporting Checklist: The authors have completed the TRIPOD reporting checklist. Available at https://tlcr. amegroups.com/article/view/10.21037/tlcr-23-666/rc

Data Sharing Statement: Available at https://tlcr.amegroups. com/article/view/10.21037/tlcr-23-666/dss

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Conflicts of Interest: All authors have completed the ICMJE uniform disclosure form (available at https://tlcr.amegroups. com/article/view/10.21037/tlcr-23-666/coif). The authors have no conflicts of interest to declare.

Ethical Statement: The authors are accountable for all aspects of the work in ensuring that questions related to the accuracy or integrity of any part of the work are appropriately investigated and resolved. The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013). This two-institution, retrospective cohort study was approved by the institutional review board of Beijing Chaoyang Hospital (IRB No. 2021-ke-135), and written informed consent for this retrospective analysis was waived. The People's Liberation Army General Hospital

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was informed and agreed with this study.

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