

Machine learning-based combined nomogram for predicting the risk of pulmonary invasive fungal infection in severely immunocompromised patients

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Background: Early and accurate diagnosis of invasive fungal infection (IFI) is pivotal for the initiation of effective antifungal therapy for patients with hematologic malignancies.

Methods: This retrospective study involved 235 patients with hematologic malignancies and pulmonary infections diagnosed as IFIs (n=118) or bacterial pneumonia (n=117). Patients were randomly divided into training (n=188) and validation (n=47) datasets. Four feature selection methods with nine classifiers were implemented to select the optimal machine learning (ML) model using five-fold cross-validation. A radiomic signature was constructed using a linear ML algorithm, and a radiomic score (Radscore) was calculated. The combined model was developed with the Radscore, the significant clinical and radiologic factors were selected using multivariable logistic regression, and the results were presented as a clinical radiomic nomogram. A prospective pilot study was also conducted to compare the classification performance of the combined nomogram with practicing radiologists.

Results: Significant differences were found in the Radscore between IFI and bacterial pneumonia patients in the training (0.683 *vs.* –0.724, P<0.001) and validation set (0.353 *vs.* –0.717, P=0.002). The combined model showed good discrimination performance in the validation cohort [area under the curve (AUC) =0.844] and outperformed the clinical (AUC =0.696) and radiomics (AUC =0.767) model alone (both P<0.05).

Conclusions: The clinical radiomic nomogram can serve as a promising predictive tool for IFI in patients with hematologic malignancies.

Keywords: Machine learning (ML); invasive fungal infection (IFI); radiomics; computed tomography (CT); hematologic malignancy

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Introduction

Invasive fungal infection (IFI) in the lung is a common complication in severely immunocompromised patients, such as those with hematologic malignancies who have undergone hematopoietic stem cell transplantation or those receiving high-dose chemotherapy (1). IFI is associated with substantial morbidity, having a mortality rate of 50% to 85% (2); given this, early and accurate diagnosis of IFI is vital to enable optimized treatment and improved patient outcomes. Clinicians now have various commercially available tools for IFI diagnosis, and each tool has its inherent advantages and disadvantages (3). Serum galactomannan testing may help with the diagnosis, but it has limited sensitivity, which ranges from 30% to 46%, and it frequently yields falsepositive results (4,5). Recently, a consensus group of the European Organization for Research and Treatment of Cancer (EORTC) and the Mycoses Study Group (MSG) established diagnostic criteria for IFI (6). These updated guidelines are based on host factors, clinical information, and microbiological criteria, which allow classification of IFIs into various levels of probability as proven, probable, or possible IFIs. However, only a minority of cases are proven using invasive procedures because of the risk of fatal bleeding due to thrombocytopenia.

Computed tomography (CT) is an important imaging modality in the diagnosis and management of high-risk patients with pneumonia. Several studies have focused on describing early predictive CT features of pulmonary IFIs (7-9). Characteristic CT signs of invasive fungal pneumonia consist of multiple nodules with a surrounding "halo sign" and/or wedge-shaped areas of consolidation (8). However, these CT findings of IFIs are usually nonspecific and indistinguishable from other respiratory infections. In particular, when consolidation is present early during pneumonia, differential diagnosis is often difficult and only presumptive.

Radiomics, an emerging quantitative imaging method we proposed in 2012 (10), can extract mineable high-dimensional data from medical images at the voxel or pixel level. It serves as the bridge between imaging biomarkers and personalized medicine to reflect underlying structural and pathophysiologic information (11). A recent radiomics study indicated potential in the discrimination of progressive pulmonary tuberculosis from community-acquired pneumonia (12). More recently, the combination of radiomics and machine learning (ML) models has been demonstrated to potentially offer improved diagnostic,

prognostic, and predictive accuracy (13). The correlation of imaging findings with clinical information is fundamental for rapid, accurate diagnosis (14). However, to the best of our knowledge, an ML-based approach for incorporating imaging biomarkers with clinical characteristics in the prediction of IFIs for clinical decision support has not been developed.

Therefore, the purpose of this study was to investigate whether an ML-based clinical radiomic nomogram integrating CT radiomic features and clinical factors can distinguish IFI from bacterial pneumonia in patients with hematologic malignancies. In addition, the benefit provided by 3-dimensional (3D) radiomic features compared with 2-dimensional (2D) texture analysis was explored. We present the following article in accordance with the TRIPOD reporting checklist (available at https://atm. amegroups.com/article/view/10.21037/atm-21-4980/rc).

Methods

The clinical trial was registered on https://www.chictr. org.cn (registration number: ChiCTR2000038557). The study was conducted in accordance with the Declaration of Helsinki (as revised in 2013). The study was approved by institutional review board of Nanfang Hospital (No. NFEC-2017-087), and individual consent for this retrospective analysis was waived.

Patient selection

Patients from the Department of Hematology who underwent chest CT between June 2017 and March 2021 were identified using our institution's database. The inclusion criteria were as follows: (I) age 18 years and over and with host factors according to the EORTC and/or MSG criteria (6); (II) suspected of having a pulmonary infection and clinical symptoms and signs of persistent cough, pleural pain, or hemoptysis; (III) CT scan with thin section (slice thickness less than 1.5 mm); and (IV) CT findings of a consolidation measuring greater than 5 mm in diameter. The exclusion criteria were as follows: (I) patients who underwent CT scan within more than 5 days after initiation of symptoms (n=16); (II) patients with incomplete clinic and etiological data including basic information, comorbidities, symptoms, and laboratory findings (Table S1) available for re-evaluation (n=47); and (III) patients who received antibiotic therapy (n=23).

All patients with proven or probable pulmonary IFI were

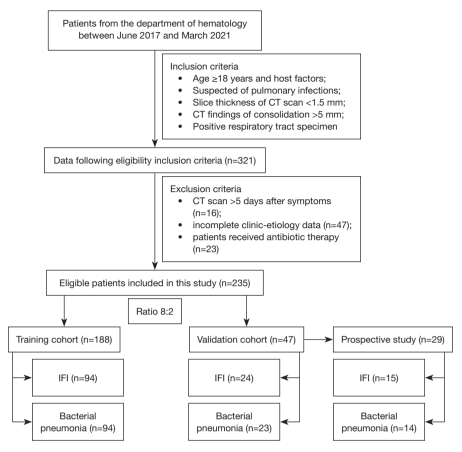


Figure 1 The flowchart of patient recruitment. CT, computed tomography; IFI, invasive fungal infection.

diagnosed according to the consensus EORTC and/or MSG criteria. Proven IFI was defined based on histopathologic evidence following percutaneous needle aspiration biopsy. Probable IFI was defined based on the presence of host factors, an area of consolidation on CT, and mycological evidence of fungal infection from culture analysis of bronchoalveolar lavage fluid or serum or microbiologic evidence of galactomannan positivity (8). Diagnosis of bacterial pneumonia was based on a positive culture of respiratory tract specimen for a bacterial organism, such as sputum or bronchoalveolar lavage fluid and CT evidence of pulmonary consolidation >5 mm in diameter (8). The pathway of patient enrollment is shown in *Figure 1*.

CT image acquisition

All patients were examined with the following CT scanners: Brilliance iCT (Philips Healthcare, Best, Netherlands), GE Discovery CT 750 HD (GE Medical Systems, Chicago, IL, USA), and Somatom Definition and Somatom Emotion

(Siemens Medical Solutions, Erlangen, Germany). The CT scanning parameters included a tube voltage of 100 to 130 kV, automatic tube current modulation, a detector collimation of 192×0.6 mm or 128×0.625 mm, and a slice thickness of 1 to 1.5 mm. All images were reconstructed with a high kernel (b60) and a matrix of 512×512 .

Lesion segmentation

The CT images were independently interpreted by 2 thoracic radiologists (G.W and C.Y, with 6 and 12 years of chest CT experience, respectively) who were blinded to the clinical information. The following semantic radiological features were assessed according to published definitions (8) by consensus: single/multiple lesions, consolidation (patchy, segmental, or wedge-shaped), halo sign, cavitation, nodule, and ground-glass opacity (GGO).

The volumes of interest (VOIs) were manually drawn on CT axial slices by 1 thoracic radiologist (C.Y) using ITK-SNAP (version 3.6.0, www.itksnap.org) and encompassed

the entire lesion. The 2D-labeling regions of interest (ROIs) were manually delineated based on the single-axial image with the largest lesion area. If multiple lesions were present, only the largest cross-sectional lesion was segmented.

Image preprocessing and feature extraction

Image resampling and gray-level normalization were performed before feature extraction. All image data were resampled to a 1×1×1 mm³ voxel size, and the gray level was normalized to 64 levels. Images were filtered and radiomic features were extracted from VOIs and ROIs using Pyradiomics 2.1.2 (https://pyradiomics.readthedocs.io/en/latest/index.html) (15). The 3D (n=835) and 2D (n=815) radiomic features included (I) first-order statistic features; (II) shape-based features; (III) texture features; and (IV) wavelet-based features.

Dimension reduction with reproducibility and collinearity analysis

A high-dimensional feature reduction process was then performed to select the most relevant features for differential diagnosis. To verify the robustness of the radiomic features, CT images of 30 cases were randomly chosen and independently delineated by 2 radiologists (C.Y and S.W). To assess the intraobserver reproducibility, the same procedure was performed twice by 1 radiologist (C.Y) within 2 weeks of the first delineation. The intra- and interclass correlation coefficient (ICC) was calculated to evaluate the reproducibility of radiomic features. Features with good agreement (both ICCs >0.8) were selected for further analyses. Collinearity analysis with Pearson correlation coefficients was performed to evaluate the relevance and redundancy of the features. Features indicated by a Pearson correlation coefficient (r) >0.9 were eliminated.

Feature selection and model building

The ML models were built with different combinations of 4 common feature selection methods [analysis of variance (ANOVA), recursive feature elimination (RFE), Relief, and Kruskal-Wallis rank-sum test (KW)] and 9 classification methods (logistic regression (LR), least absolute shrinkage and selection operator (LASSO), support vector machine (SVM), linear discriminant analysis (LDA), decision tree, random forest, Adaboost, Gaussian process, and naïve Bayes). Patients were randomly divided into the training

and validation group at a ratio of 8:2. All the classifiers were trained with a 5-fold cross-validation. The area under the receiver operator characteristic (ROC) curve (AUC) was used to evaluate the discriminative performance of different models. The model with the highest AUC value in the cross-validation cohort was considered the radiomic model. The radiomic signature (Radscore) was then calculated using a multivariable logistic regression model by linearly combining the most predictive features, which were weighted by their respective coefficients.

Clinical and radiological model building and evaluation

Considering the potential additional value of clinical characteristics, CT features, and laboratory results (Table S1) to improve the predictive performance, we first identified independent variables (P<0.1) using univariate analysis. In the training cohort, a clinical radiomic model incorporating the radiomic signature and clinical predictors was developed using multivariable logistic regression analysis. A collinearity diagnosis with a variance inflation factor was performed before the multivariate analysis. Backward stepwise selection was employed using the likelihood ratio test with Akaike's information criterion (AIC) as the stopping rule. The model with the lowest AIC value was selected as the final model. A clinical model was also constructed separately using multivariable logistic regression analysis based on statistically significant clinical factors. The ROC curve and AUC were applied to evaluate the predictive accuracy of established models in the validation cohort. Differences in AUC between various prediction models were compared using the DeLong test.

Construction and validation of the radiomic nomogram

The radiomic nomogram was built based on proportionally converting regression coefficients of each predictor in the combined model to a 0- to 100-point scale. Calibration of the clinical radiomic nomogram was plotted using the calibration curve, which measures the consistency between the nomogram-predicted probability and the actual rate. The Hosmer-Lemeshow test was used to assess the goodness of fit of the nomogram. In addition, decision curve analysis (DCA) was conducted to estimate the clinical utility of the radiomic signature, clinical risk factors, and nomogram by quantifying net benefits at different threshold probabilities in the validation dataset. The decision curves of the treat-all strategy and the treat-none strategy were

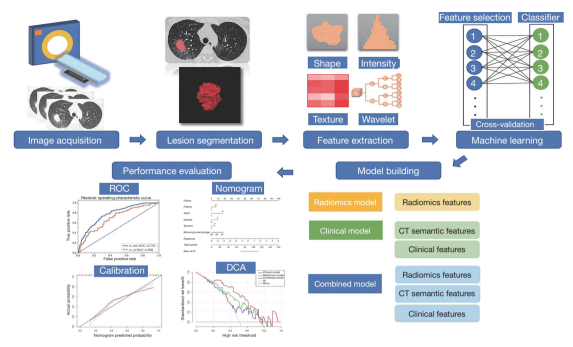


Figure 2 The workflow of data analysis. ROC, receiver operator characteristic; AUC, area under the ROC curve; GGO, ground-glass opacity; IFI, invasive fungal infection; DCA, decision curve analysis; CT, computed tomography.

used as references in the DCA. The workflow of this study is shown in *Figure 2*.

Comparison of the combined nomogram against practicing radiologists

A prospective pilot study was also conducted at our institution from April 2021 to August 2021. The inclusion and exclusion criteria and the definitions for bacterial pneumonia and pulmonary IFI mentioned above were used in the pilot study cohort. A total of 15 patients with IFI and 14 patients with bacterial pneumonia were enrolled to compare the performance of the combined nomogram with practicing radiologists in terms of classification. One junior and one senior radiologist (with 6 and 12 years of experience, respectively) were employed to participate in the study. Ground truth was established based on the enrollment criteria mentioned above.

Statistical analysis

All statistical analyses were performed using R software (version 3.5.2; The R Foundation for Statistical Computing, Vienna, Austria) and SPSS software (version 22.0, IBM Corp., Armonk, NY, USA). Continuous and categorical data

were compared using independent *t*- (or Mann-Whitney U) tests and chi-square (or Fisher's exact) statistics, respectively, as appropriate. The ML algorithms were programmed with Python scikit-learn environment (version 0.19.1). A 2-sided P value <0.05 was considered to be significant.

The "glmnet" package was used to perform LASSO logistic regression, the ROC was plotted using the "pROC" package, nomogram construction and calibration plotting were performed using the "rms" package, the Hosmer-Lemeshow test was performed using the "generalhoslem" package, and the "rmda" package was used to construct the DCA curve.

Results

Clinical and CT imaging characteristics

A total of 235 patients consisting of 118 individuals with IFI and 117 with bacterial pneumonia were enrolled in the study. Clinical characteristics including CT semantic features, clinical variables, and laboratory parameters of patients with IFI and bacterial pneumonia in both the training and validation datasets are shown in *Tables 1,2*. There were no differences in sex, age, or most clinical and CT features between the 2 groups. The halo sign

Table 1 CT sematic features of patients with pulmonary infections

OT factories	Training set (n=188), n (%)			Validation set (n=47), n (%)			
CT features	IFI (n=94)	Bacterial pneumonia (n=94)	P value	IFI (n=24)	Bacterial pneumonia (n=23)	P value	
Number of lesions			0.631			0.238	
Single	29 (30.9)	26 (27.7)		9 (37.5)	5 (21.7)		
Multiple	65 (69.1)	68 (72.3)		15 (62.5)	18 (78.3)		
Consolidation							
Patchy	71 (75.5)	82 (87.2)	0.039*	16 (66.7)	19 (82.6)	0.210	
Segmental	46 (48.9)	34 (36.2)	0.077	10 (41.7)	9 (39.1)	0.859	
Wedge-shaped	10 (10.6)	12 (12.8)	0.650	2 (8.3)	3 (13.0)	0.601	
Halo sign			0.013*			0.172	
Absence	33 (35.1)	50 (53.2)		12 (50.0)	16 (69.6)		
Presence	61 (64.9)	44 (46.8)		12 (50.0)	7 (30.4)		
Cavitation			0.470			0.157	
Absence	89 (94.7)	91 (96.8)		22 (91.7)	23 (100.0)		
Presence	5 (5.3)	3 (3.2)		2 (8.3)	0 (0.0)		
Nodule			0.011*			0.298	
Absence	72 (76.6)	85 (90.4)		18 (75.0)	20 (87.0)		
Presence	22 (23.4)	9 (9.6)		6 (25.0)	3 (13.0)		
GGO			0.005*			0.955	
Absence	91 (96.8)	80 (85.1)		21 (87.5)	20 (87.0)		
Presence	3 (3.2)	14 (14.9)		3 (12.5)	3 (13.0)		

^{*,} P<0.05. CT, computed tomography; IFI, invasive fungal infection; GGO, ground-glass opacity.

and nodule were more common in patients with IFI than in those with bacterial pneumonia, while patchy consolidation and GGO were more frequent in those with bacterial pneumonia (each P<0.05) in the training set. Significant differences between the 2 groups were observed in the incidence of sputum, neutrophil percentage, and lymphocyte percentage.

Model assessment

A total of 36 predictive radiomic models were built through the combination of 4 selection methods and 9 classifiers. The heatmaps of mean AUC values from various ML models in the training, cross-validation, and validation cohorts are shown in *Figure 3* and Figure S1. Generally, the 3D-based models represented better performance than did the 2D-based models. Moreover, LR classifiers

outperformed other classification methods in terms of discrimination. Among models using parameters from 3D and 2D images, the selection method KW with LR classifier yielded the highest AUC of 0.644 and 0.641, respectively in the cross-validation group.

The selected radiomic features to discriminate IFI and bacterial pneumonia in the training cohort are summarized in *Table 3*, including 1 shape feature, 2 texture features, and 9 wavelet features. A radiomic signature (Radscore) was then constructed using the features and their respective coefficients. There was a significant difference in Radscore between the IFI and bacterial pneumonia patients in the training cohort (0.683 *vs.* -0.724; P<0.001), which was confirmed in the validation cohort (0.353 *vs.* -0.717, P=0.002). The AUC of the radiomic signature performance was 0.803 (95% CI: 0.739–0.866) in the training cohort and 0.767 (95% CI: 0.624–0.910) in the validation cohort.

Table 2 Demographic and clinical characteristics of patients with pulmonary infections

Clinical factors		Training set (n=188)	Validation set (n=47)			
Clinical factors	IFI (n=94)	Bacterial pneumonia (n=94)	P value	IFI (n=24)	Bacterial pneumonia (n=23)	P value
Age (years)	43.3±15.6	40.5±13.7	0.188	39.3±11.9	43.7±16.2	0.293
Sex			0.766			0.671
Male	57 (60.6)	55 (58.5)		14 (58.3)	12 (52.2)	
Female	37 (39.4)	39 (41.5)		10 (41.7)	11 (47.8)	
Hematologic malignancy			0.379			0.859
AML	55 (58.5)	49 (52.1)		14 (58.3)	14 (60.9)	
Non-AML	39 (41.5)	45 (47.9)		10 (41.7)	9 (39.1)	
Infection history	43 (45.7)	35 (37.2)	0.236	11 (45.8)	6 (26.1)	0.024*
Chemotherapy	75 (79.8)	65 (69.1)	0.094	20 (83.3)	11 (47.8)	0.010*
Immunosuppressant	5 (5.3)	7 (7.4)	0.551	2 (8.3)	1 (4.3)	0.576
HSCT	30 (31.9)	27 (28.7)	0.634	8 (33.3)	8 (34.8)	0.917
GVHD	13 (13.8)	8 (8.5)	0.247	4 (16.7)	4 (17.4)	0.947
Remission	7 (7.4)	6 (6.4)	0.774	4 (16.7)	0	0.041*
Fever	56 (59.6)	56 (59.6)	1.000	13 (54.2)	18 (78.3)	0.081
Body temperature (°C)	37.4±4.1	37.9±1.3	0.248	37.31.3	38.21.4	0.233
Cough	34 (36.2)	44 (46.8)	0.139	10 (41.7)	14 (60.9)	0.188
Sputum	24 (25.5)	38 (40.4)	0.030*	6 (25.0)	13 (56.5)	0.028*
Chest pain	6 (6.4)	7 (7.4)	0.774	1 (4.2)	2 (8.7)	0.525
Hemoptysis	2 (2.1)	3 (3.2)	0.65	0	0	1.000
Dyspnea	2 (2.1)	4 (4.3)	0.407	0	0	1.000
Chill	10 (10.6)	7 (7.4)	0.446	4 (16.7)	4 (17.4)	0.947
Headache	12 (12.8)	12 (12.8)	1.000	6 (25.0)	6 (26.1)	0.932
WBC (×10 ⁹ /L)	10.0±46.8	8.8±28.2	0.831	7.8±10.4	8.3±15.8	0.893
Neutrophil (×10°/L)	3.5±10.9	2.6±5.1	0.505	7.6±15.6	2.5±3.2	0.137
Neutrophil percentage (%)	48.5±29.2	40.1±27.0	0.043*	55.9±26.6	42.1±26.4	0.088
Lymphocyte (×10 ⁹ /L)	5.1±31.1	2.6±7.3	0.449	1.4±1.6	1.7±2.0	0.543
Lymphocyte percentage (%)	36.9±27.9	44.9±27.2	0.049*	31.0±24.4	34.5±23.5	0.635
Monocyte (×10 ⁹ /L)	1.3±5.6	3.5±17.0	0.251	1.3±3.3	4.1±11.3	0.257
Monocyte percentage (%)	10.6±11.3	14.8±17.1	0.052	11.9±14.3	21.3±24.7	0.127
CRP (mg/dL)	72.8±65.5	72.2±70.8	0.954	62.1±67.5	70.4±65.5	0.690
G test (%)	12/31 (38.7)	8/41 (19.5)	0.072	2/9 (22.2)	3/9 (33.3)	0.599
PCT (pg/mL)	1.2±3.9	1.5±5.7	0.746	0.6±1.2	0.3±0.3	0.405

^{*,} P<0.05. IFI, invasive fungal infection; AML, acute myelogenous leukemia; HSCT, hematopoietic stem cell transplant; GVHD, graft-versus-host disease; WBC, white blood count; CRP, C-reactive protein; PCT, procalcitonin.

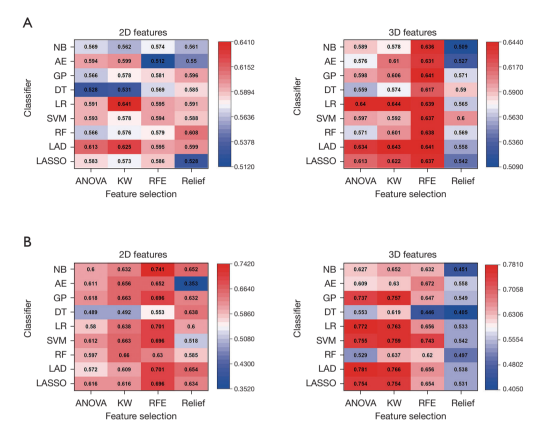


Figure 3 Heatmaps of the AUCs from different combinations of feature selection methods (columns) and classification algorithms (rows) for 2D and 3D radiomics features in the 5-fold cross-validation (A) and validation (B) dataset. ANOVA, analysis of variance; KW, Kruskal-Wallis rank-sum test; RFE, recursive feature elimination; NB, naïve Bayes; AE, Adaboost ensemble; GP, Gaussian process; DT, decision tree; LR, logistic regression; SVM, support vector machine; RF, random forest; LDA, linear discriminant analysis; LASSO, least absolute shrinkage and selection operator; AUC, area under the receiver operator characteristic curve.

Effect of combining clinical parameters and radiomics in prediction

In multivariable logistic regression analysis, significant risk factors for IFI included a higher Radscore, lower monocyte percentage, the absence of sputum, and the presence of halo sign, nodule, patchy consolidation, or GGO (all P<0.05). There was no multicollinearity between the significant factors and the Radscore. The list of the selected factors and their associated coefficients in the clinical and combined model are illustrated in *Table 3*. Our combined model integrating clinical and radiological variables with a radiomic signature achieved an AUC of 0.844 in the validation set, which indicated improved prediction performance compared to the clinical model (AUC =0.696) and radiomic model (AUC =0.767) according to the Delong test (both P<0.01). The predictive performance results of

different models are presented in Figure 4 and Table 4.

Nomogram apparent performance and clinical use of DCA

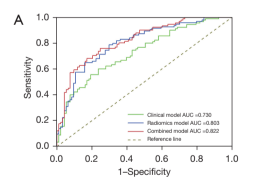
The clinical radiomic nomogram was constructed based on the combined model (*Figure 5A*). The calibration curves demonstrated good consistency between the prediction and actual observation of IFI and bacterial pneumonia in both cohorts (*Figure 5B*, *5C*). The Hosmer-Lemeshow test yielded nonsignificant statistics in the training cohort (P=0.619) and validation cohort (P=0.446), suggesting a perfect fit of the nomogram.

The clinical DCA was used to demonstrate clinical decision utility of the combined nomogram (*Figure 6*). The nomogram showed higher net benefit than did that using the radiomic signature or clinical features alone, which were better than the treat-all or treat-none strategies.

Table 3 Significant variables selected by multivariate logistic regression analysis

Radiomics model	Coefficient	Clinical model	Coefficient	Combined model	Coefficient
original_shape_Sphericity	4.327	Halo sign	0.726	Radscore	0.959
original_glcm_lmc1	2.942	Patchy consolidation	-1.112	Patchy consolidation	-0.763
original_glcm_lmc2	-1.747	GGO	-1.748	GGO	-1.087
wavelet-LLH_firstorder_Kurtosis	4.45	Nodule	1.035	Nodule	1.087
wavelet-LLL_firstorder_Robust Mean Absolute Deviation	2.004	Lymphocyte percentage	-0.013	Sputum	-0.599
wavelet-LLL_glcm_Correlation	3.919	Monocyte percentage	-0.029	Monocyte percentage	-0.024
wavelet-LLL_glcm_Joint Energy	0.501	intercept	1.352	intercept	1.096
wavelet-LLL_glcm_lmc1	-0.978				
wavelet-LLL_glcm_lmc2	3.403				
wavelet-LLL_glcm_Maximum Probability	1.908				
wavelet-LLL_glszm_Large Area High Gray Level Emphasis	-2.48				
wavelet-LLL_gldm_Large Dependence High Gray Level Emphasis	0.109				
Intercept	-0.021				

GGO, ground-glass opacity.



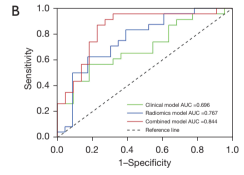


Figure 4 ROC curves of the clinical, radiomics, and combined models in the training (A) and validation (B) dataset. AUC, area under the receiver operator characteristic curve; ROC, receiver operator characteristic.

Comparison of the combined nomogram with practicing radiologists

The performances on differentiating IFI from bacterial pneumonia of the developed nomogram and radiologists were then compared. The diagnostic system achieved an accuracy of 79.3%, a sensitivity of 80.0%, and a specificity of 84.6%, which was overall superior to the values of the junior and senior radiologist (*Figure 7*). Representative cases are shown in *Figure 8*.

Discussion

In the current study, we developed and validated an ML-based nomogram that incorporates the radiomic signature and clinical characteristics of noninvasive, individualized diagnoses of IFI in patients with hematologic malignancies. The highest performance was achieved with the combination of the selection method KW with LR classifier. The proposed clinical radiomic nomogram demonstrated favorable discrimination in the validation

Table 4 Diagnostic performance of each model in the training and validation cohorts

Model	Training cohort			Validation cohort			
	Clinical model	Radiomics model	Combined model	Clinical model	Radiomics model	Combined model	
AUC (95% CI)	0.730 (0.658, 0.802)	0.803 (0.739, 0.866)	0.822 (0.762, 0.881)	0.696 (0.542, 0.850)	0.767 (0.624, 0.910)	0.844 (0.723, 0.965)	
Accuracy	0.681	0.748	0.755	0.689	0.734	0.822	
Sensitivity	0.556	0.656	0.756	0.565	0.609	0.913	
Specificity	0.806	0.839	0.753	0.818	0.864	0.727	
Positive predictive value	0.557	0.657	0.756	0.592	0.638	0.952	
Negative predictive value	0.804	0.837	0.753	0.781	0.825	0.697	

AUC, area under the receiver operator characteristic curve; CI, confidence interval.

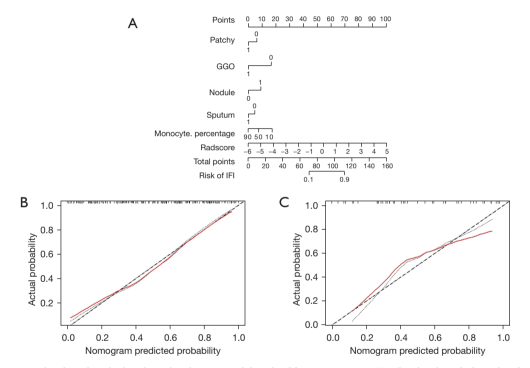


Figure 5 Nomogram developed with the clinical radiomic model and calibration curves. (A) The developed clinical radiomic nomogram for predicting the probability of IFIs. (B,C) Calibration curves for predicting IFIs in the training and validation cohorts. Calibration curves indicate the goodness of fit of the constructed nomogram. The predictive performance of the nomogram (red line) closer to the ideal prediction line (45° gray line) represents a higher predictive accuracy of the nomogram. GGO, ground-glass opacity; IFI, invasive fungal infection.

cohort (AUC =0.844), outperforming the radiomic-only and clinical prediction model. The diagnostic performance of the combined model was demonstrated to be superior to that of practicing radiologists in a prospective fashion. ML-based radiomic analysis could potentially serve as a quantitative imaging biomarker to aid discrimination of IFI from bacterial pneumonia.

Pulmonary IFI is life-threatening condition with a considerable rate of mortality in immunocompromised patients (16). Patients with hematologic malignancies have an elevated risk of fungal infection due to hypofunction of the immune system and frequent use of immunosuppressive agents in clinical treatment, and symptoms of pulmonary IFI are often nonspecific (17). In clinical practice, the

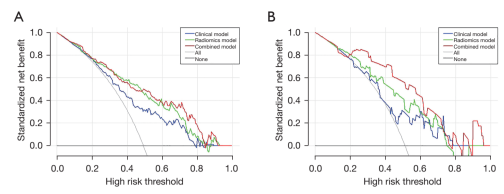


Figure 6 The decision curve analysis for the clinical, radiomic, and combined models in the training and validation cohorts. The y-axis indicates the net benefit; the x-axis indicates threshold probability. The gray line represents the decision curve of the assumption that all patients have IFIs ("treat all"); the black line represents the decision curve of the assumption that no patients have an IFI ("treat none"). The farther the decision curve is from the 2 extreme lines, the higher the clinical decision net benefit of the model. IFI, invasive fungal infection.

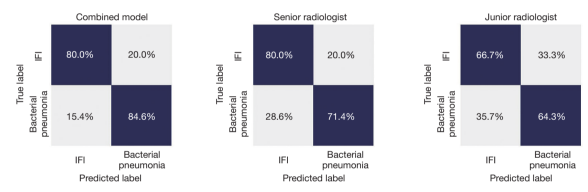


Figure 7 Comparison of diagnostic performance between the combined nomogram and practicing radiologists in a prospective fashion. IFI, invasive fungal infection.

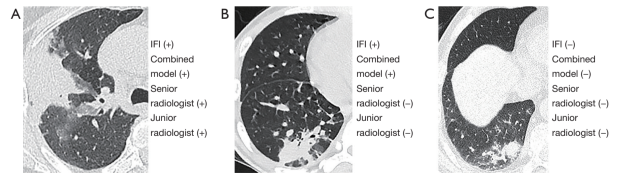


Figure 8 Representative cases to show diagnostic ability of the combined model and practicing radiologists for IFIs. (A) A 69-year-old woman with acute myelogenous leukemia and IFI. The mass was correctly diagnosed as an IFI by the combined model and practicing radiologists. (B) A 23-year-old man with acute lymphatic leukemia and IFI. The lesion was correctly diagnosed as an IFI by the combined model but was misdiagnosed by the 2 radiologists. (C) A 67-year-old man with acute myelogenous leukemia and bacterial pneumonia. The lesion was correctly diagnosed by the combined model and the senior radiologist but was misdiagnosed by the junior radiologist. IFI, invasive fungal infection; +, positive; -, negative.

most challenging task is to distinguish IFI from bacterial pneumonia. Similarly, the more conventional bacterial pathogens are present in opportunistic infections in immunocompetent individuals, with fever, chest pain, hemoptysis, focal consolidation, and rapid increases in inflammatory markers (18). As outlined by the EORTC and MSG in 2002 and updated in 2008, the classification of proven, probable, or possible IFIs is a helpful tool for diagnostic purposes (6). However, the number of proven cases is quite low because fine-needle aspiration is invasive and limited to thrombocytopenia patients.

CT plays a crucial role in clinical practice for the noninvasive diagnosis and management of pulmonary infections (7). The updated guidelines have enlarged the diagnostic role of CT scans, but imaging signs such as macronodules or patchy consolidation with or without the halo sign are still not specific, particularly in the early stages of pulmonary infections. Thus, in most cases, differentiation of fungal and other pneumonias using CT with the traditional practice of visual interpretation remains challenging. Consolidation is the most common CT finding in both diseases. Chen et al. (8) reported that multifocal nonsegmental consolidation was found significantly more often in patients with IFI (48.0%) than in those with bacterial pneumonia (22.6%). By contrast, we found a significantly higher frequency of focal patchy consolidation in patients with bacterial pneumonia than in those with IFI (87.2% vs. 75.5%; P<0.05), although similar definitions were used for bacterial pneumonia and pulmonary IFI. This can be attributed to the distribution bias relating to different patient populations. Moreover, the presence of nodules has been proven to be a significant risk predictor, which has long been considered to increase the risk of IFI (19).

Radiomics is a relatively new technique and has received much attention in cancer research (20). This approach allows high-throughput quantitative extraction of numerous invisible features (21). Radiomic signatures have succeeded in a variety of tumor-related analyses (22-26) but have rarely been used to investigate infectious diseases. It is known that IFI and bacterial pneumonia are completely different in terms of angiogenesis and inflammation. The heterogeneity, therefore, can be revealed by radiomic features. In this study, we found that the radiomic features of pulmonary lesions could be used to facilitate IFI diagnosis, with an accuracy of 0.767 in the validation set. Similar to our results, a recent study used texture analysis from 970 CT imaging features and demonstrated that radiomic features could be used as noninvasive biomarkers for the differential

diagnosis of primary progressive pulmonary tuberculosis from community-acquired pneumonia in children (12). This study also demonstrated that 3D radiomic features showed better reproducibility and discrimination efficacy than did the 2D features, although 3D features are more time-consuming and require heavy-load computation. This is probably because the VOIs derived from 3D segmentation can provide the entire volumetric information of the lesions and be less vulnerable to hand-related artifacts (27).

Recent research has suggested that radiomics combined with ML algorithms showed promising potential in various fields, such as the differential diagnosis of pulmonary lesions (28) as well as the prediction of hospital stays in patients with SARS-CoV-2 infection (29) and diseasespecific survival outcomes (30). Compared with previous studies, more selection methods and classifiers were assessed in this work to identify the optimal model with the best diagnostic performance. Among the classifiers showing feasible discriminative ability, LR is a representative of the linear classifiers, which enables the combination of selected radiomic features into a radiomic signature (31). By contrast, SVM is a nonlinear classifier, which uses support vectors to develop a polynomic hyperplane to separate classes. In this study, LR-based models achieved satisfactory performance with the highest AUC in the cross-validation group. Nonetheless, the suitable ML methods may vary in different clinical applications. For example, Zhang et al. (32) reported that RFE with SVM showed the best performance in molecular subtype stratification of lower-grade glioma.

According to multiple logistic regression analysis, this study identified lower monocyte percentage as an independent clinical variable associated with IFI. In particular, patients with impaired cellular function of phagocytes are susceptible to fungal infection, with an incidence of up to 25% being reported in the literature (33). The clinical radiomic model achieved better predictive efficacy than did the radiomic signature or clinical features alone, with a higher AUC and net benefit in DCA. Based on our results in this prospective pilot study, the combined model performed equally well (12/15, 80.0%) in the diagnosis of pulmonary IFI to the senior radiologist and outperformed a junior radiologist in discriminating IFI and bacterial pneumonia. The constructed nomogram in our study was conveniently used to predict the individualized probability of IFI.

This study has several limitations. First, the patients were assessed retrospectively in the training and validation phase, and thus selection bias was inevitable. Second, the sample size was relatively small, which partly led to the similar

diagnostic performance across most of the tested models. More samples from different independent centers are needed to validate the robustness and generalization of the prediction model. Third, the boundary delineation of the 3D lesions is time-consuming and complicated. Potential subjectivity during the process of manual segmentation may hamper reproducibility of the results. Fourth, although multifold cross-validation was performed when constructing the ML classifiers, overfitting still occurred. This computational approach requires improvement in further studies. Finally, the gold standard for investigating IFI was based on the respiratory tract specimen and not on the histologic examination undertaken in this study.

Conclusions

A noninvasive ML-based radiomic nomogram was developed through integrating CT radiomic signatures and clinical risk factors, which exhibited favorable accuracy for the differentiation of IFI and bacterial pneumonia in patients with hematologic malignancies. This quantitative radiomic diagnostic model may be useful for precision medicine and improving medical decision support.

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Footnote

Reporting Checklist: The authors have completed the

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Conflicts of Interest: All authors have completed the ICMJE uniform disclosure form (available at https://atm. amegroups.com/article/view/10.21037/atm-21-4980/coif). TZ and HL have associations with Philips Healthcare, and each provided technical support for data analysis. HCW owns (minority) shares in Oncoradiomics. PL reports, within and outside the submitted work, grants/ sponsored research agreements from Varian Medical, Oncoradiomics, ptTheragnostic, Health Innovation Ventures, and DualTpharma. He received an advisor/ presenter fee and/or reimbursement of travel costs/external grant writing fee and/or in-kind manpower contribution from Oncoradiomics, BHV, Merck, and Convert Pharmaceuticals. He owns shares in Oncoradiomics SA and Convert Pharmaceuticals SA. He is a coinventor of 2 issued patents with royalties on radiomics (PCT/NL2014/050248, PCT/NL2014/050728) licensed to Oncoradiomics, 1 issued patent on mtDNA (PCT/EP2014/059089) licensed to ptTheragnostic/DNAmito, and 3 nonpatentable inventions (software) licensed to ptTheragnostic/DNAmito, Oncoradiomics, and Health Innovation Ventures. None of the authors had control of the data in a manner that would present a conflict of interest for the other employees or consultant authors. The other 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). The study was approved by institutional review board of Nanfang Hospital (No. NFEC-2017-087), and individual consent for this retrospective analysis was waived.

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Supplementary

Table S1 Potential clinical characteristics, computed tomography features, and laboratory factors of patients with pulmonary infections

	Laboratory findings
Basic information	
Age	Blood leukocyte count, median (IQR), ×10 ⁹ /L
Sex	Neutrophil count, median (IQR), $\times 10^9/L$ lasting >10 days (within 60 days)
Time of onset of illness, days	Neutrophil percentage, median (IQR), %
IFI history	Lymphocyte count, median (IQR), ×10 ⁹ /L
Comorbidities	Lymphocyte percentage, median (IQR), %
AML/non-AML	Monocyte count, median (IQR), ×10 ⁹ /L
Chemotherapy	Monocyte percentage, median (IQR), %
Immunosuppressant	C-reaction protein mg/dL
HSCT	
GVHD	
Relapse or Remission	
Symptoms	
Fever, No. (%)	
Maximum body temperature, °C	
Cough, No. (%)	
Sputum	
Chest pain	
Dyspnea	
Hemoptysis	
Chill	
Headache or dizziness	

IFI, invasive fungal infection; AML, acute myelogenous leukemia; HSCT, hematopoietic stem cell transplant; GVHD, graft-versus-host disease; IQR, interquartile range.

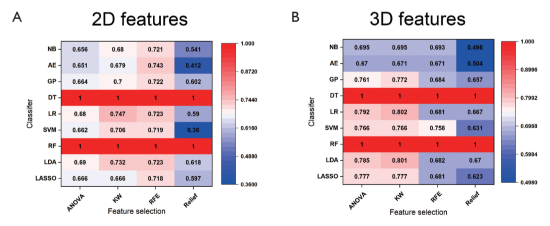


Figure S1 Heatmaps of area under the receiver operator characteristic curves from different combinations of feature selection methods (columns) and classification algorithms (rows) for 2D and 3D radiomics features in the training dataset. ANOVA, analysis of variance; RFE, recursive feature elimination; KW, Kruskal-Wallis rank-sum test; LR, logistic regression; LASSO, least absolute shrinkage and selection operator; SVM, support vector machine; LDA, linear discriminant analysis; DT, decision tree; RF, random forest; GP, Gaussian process; NB, naïve Bayes.