# Correlation between clinical, CT features and EGFR gene in minimally invasive adenocarcinoma

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### **ABSTRACT**

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Background: To analyze the correlation between epidermal growth factor receptor (EGFR) gene mutations and clinical as well as computed tomography (CT) signs in minimally invasive adenocarcinoma (MIA). Materials and Methods: A total of 118 patients with MIA treated at the Department of Thoracic Surgery, Tangshan Gongren Hospital, from December 2018 to December 2021 were included in this study. Patients were categorized into two groups based on their Epidermal Growth Factor Receptor (EGFR) mutation status: a mutation group and a wild-type group. Differences in clinical and CT features between the two groups were recorded and statistically analyzed. Results: Among the 118 patients, 69 were in the mutation group, while 49 were in the wild-type group. No significant differences were observed in gender, smoking history, age, mean nodule diameter, peripheral emphysema, air bronchogram, lobulation sign, pleural indentation sign, morphology, tumor-lung interface, peripheral fibrosis, and density between the two groups (P>0.05). However, significant differences were noted in the spiculation sign, vacuole sign, and vascular convergence sign between the groups (P<0.05). Logistic regression analysis identified the vacuole sign with an odds ratio (OR) of 0.410 (P>0.05) and the spiculation sign and vascular convergence sign with ORs of 1.498 and 2.262, respectively (P<0.05), as independent risk factors for predicting EGFR mutations. The areas under the ROC curves (AUC) for the spiculation sign and vascular convergence sign were 0.634 and 0.735, respectively, in distinguishing EGFR gene mutations in MIA. Conclusion: CT imaging demonstrated that MIA patients presenting with spiculation, vacuole, and vascular convergence signs had a higher likelihood of EGFR gene mutations. Among these, the spiculation and vascular convergence signs are critical, independent risk factors for predicting EGFR gene mutations in MIA patients.

#### **INTRODUCTION**

Lung cancer ranks second in about 18.4 percent of the causes of cancer deaths worldwide, according to the latest data <sup>(1)</sup>. In the pathological classification of NSCLC, adenocarcinoma has the highest incidence rate <sup>(2)</sup>. With the rapid development of pathology and molecular biology, the study of lung adenocarcinoma has gradually deepened. The growth of lung adenocarcinoma follows the evolution pattern from in situ adenocarcinoma (AIS), MIA to gradually progressing to invasive adenocarcinoma (IAC) <sup>(3, 4)</sup>. Different subtypes of lung adenocarcinoma subtypes need different surgical methods with different clinical outcomes <sup>(5)</sup>. Therefore, accurate preoperative pathological type identification of lesions is crucial to the selection of surgical methods.

At present, most patients with lung adenocarcinoma have hidden clinical symptoms, and the pathological type of lesions cannot be determined when some lesions are detected. Due to a variety of reasons, patients cannot be treated surgically, mainly

with radiotherapy and chemotherapy, but the side effects of chemotherapy drugs cannot be tolerated by some patients, and the curative effect is not good <sup>(6)</sup>.

With the continuous development of biological research, more and more scholars have transformed the traditional view of lung cancer treatment. The main clinical applications of biological therapy are also gradually becoming mature. Targeted drugs have major breakthroughs in the field of cancer, among which the EGFR epidermal growth factor receptor is the most studied and most documented molecular target compared with other driver genes (7). Studies show that is a critical receptor on the surface of various normal and mutant cells (8). As the EGFR gene decisive effect in promoting the growth and proliferation of tumor cells, increased levels of EGFR receptors can lead to disease progression and poor prognosis (9). Therefore, confirm the mutation status of EGFR is important for the molecular biological treatment lung adenocarcinoma because it allows the development of targeted therapeutic strategies to improve prognosis (10). Gefitinib is the most common

targeted therapy drug, which not only has better therapeutic effect, but also has the advantages of oral administration, good compliance and low toxicity (11). However, assessment targeting EGFR mutation status relies mainly on biopsy tissue. In practice, EGFR genetic testing poses challenges, especially because due to the high cost, many patients cannot afford the test and therefore cannot biopsy the tissue for molecular analysis. Moreover, there is obvious heterogeneity of tumor. The tumor tissue obtained by puncture cannot totally represent lesion, and duplicated puncture will increase the cost and secondary injury (12). Although serum circulating tumor cell DNA (ctDNA) can be used as an addist method for congcention DNA testing, it is still impossible to accurately understand the EGFR gene mutation (13). Accurate analysis of EGFR mutation status by invasive biopsy or ctDNA testing is sometimes not feasible. Therefore, researchers have been looking for more straightforward practical, cost-effective and approaches.

At present, CT, as the most commonly used test for lung cancer, is important in the diagnosis and efficacy evaluation of lung cancer. The biological behavior of the lesion determines its different growth patterns, and the differences in the growth patterns between the lesions can manifest by different imaging features (14). This makes it possible to combine molecular biology with modern medical imaging, laving the theoretical foundation for molecular imaging. Imaging physicians are trying to indirectly obtain genetic mutation information from imaging features with lung cancer to help ensure driver genes. At present, there are many related studies in many directions and in the number, including more studies on morphology (15, 16). Several studies have evaluated the characteristics of lung adenocarcinoma using computed tomography (CT) and concluded that CT signs can predict molecular subtypes based on genetic mutations (17). However, these findings remain controversial (18).

If we can apply CT to diagnose lesions early, and apply CT to extract multidimensional imaging features from the lesion area non-invasively analyze the tumor heterogeneity and find its internal connection with EGFR gene mutation, it can help clinical decision of treatment plan, choose surgery type or avoid unnecessary surgery. Most studies have focused on the general cases of invasive lung adenocarcinoma, which is highly innovative based on the study of clinical features, correlation of CT signs and *EGFR* gene mutations in this pathological type.

# **MATERIALS AND METHODS**

## Subjects

This study was approved by the Tangshan Gongren Hospital Ethics Committee (No. GRYY-LL-2018-48). A retrospective analysis inducted 118 patients that

underwent thoracic surgery at Tangshan Gongren Hospital between December 2018 and March 2021, all of whom had been diagnosed with pathological minimally invasive adenocarcinoma (MIA). Each patient underwent a CT examination, and EGFR gene detection performed post-surgery. Among the 118 patients with MIA, 69 categorized into the mutant group, and 49 classified as the wild-type group. The cohort included 67 females and 51 males, aged 31 to 81 years. These cases further split into two teams, 60 years as the threshold: 48 patients were 60 years or younger, and 70 patients were older than 60. Additionally, the cohort was divided based on smoking history, with 63 smokers and 65 non-smokers. The inclusion criteria: (1) postoperative pathology verify MIA following pulmonary lobectomy segmentectomy, (2) complete thin-slice CT images available preoperatively, (3) EGFR gene detection conducted postoperatively, (4) absence of other spaceoccupying lesions before surgery, and (5) no antitumor treatment before surgery. The exclusion criteria: (1) multiple lung lesions, (2) patients who had received treatment before surgery, and (3) patients whose pathology revealed other types of spaceoccupying lesions.

## **Equipment and reagents**

Philips 64-slice spiral CT machine (Nederlands Philips); Auantitative real-time PCR (Switzerland Roche Cobas Z 480); The EGFR gene detection kit (Switzerland Roche).

## CT examination method

Preoperative CT scans were using a Philips Brilliance 64-slice spiral CT scanner, 5 mm slice thickness, with reconstruction settings adjusted to a slice thickness of 1 mm. Set 120 kV of the tube voltage and set current between 120 and 380 mA.

#### CT image analysis

Two radiologists own five years of experience, independent analysis, diagnosed, and recorded the various CT features of pulmonary nodules without knowledge of the patients' gene mutation statuses. The average of the two radiologists' readings used as the final value for measured values. In cases of disagreement regarding a lesion, a consensus reached through discussion. The CT features analyzed included the average diameter of nodules, pleural indentation sign, peripheral fibrosis, spiculation sign, tumor-lung interface, lobulation sign, vacuole sign, vascular convergence sign, density, air bronchogram, peripheral emphysema, and morphology. Partial CT signs of MIA illustrated in figure 1.

## Pathological examination method

Surgically excised specimens were sent to the pathology laboratory, where experienced technicians soak the specimens in a 10% neutral formalin solution

(approval number: Jiangsu Drug Administration (quasi) word 2016 No. 20160001 China) for fixation, dehydration, transparency, embedding, sectioning, and processing with the use of hematoxylin-eosin (HE) staining (Beijing Yili Fine Chemicals Co., LTD. China). The diagnosis of MIA was verified by two pathologists after consultation, on the basis of the 2021 pathological criteria for lung adenocarcinoma (19). Pathological images of MIA illustrated in figure 2.

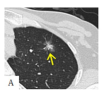
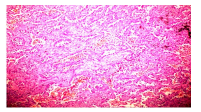






Figure 1. Legend of CT signs of minimally invasive adenocarcinama (MIA) with EGFR gene mutation. A: CT scan of a 54-year-old woman showing a mixed ground-glass opacity (GGO) nodule in the upper lobe of the left lung, measuring approximately 1.5 cm in diameter, with a visible spiculation sign (indicated by the arrow). B: CT scan of a 48-year-old man depicting a GGO nodule in the upper lobe of the left lung, measuring approximately 1.0 cm in diameter, with a visible vacuole sign (indicated by the arrow). C: CT scan of a 68-year-old woman showing a GGO nodule in the upper lobe of the left lung, measuring approximately 1.1 cm in diameter, with a visible vascular convergence sign (indicated by the arrow).

Figure 2. Pathological image of minimally invasive lung adenocarcinoma, stained with hematoxylin and eosin (HE, ×10).



#### Gene detection methods

DNA was extracted from formaldehyde-fixed, paraffin-embedded tissue sections using Roche FFPE Tissue Kit (Switzerland Roche) according to the manufacturer's instructions. PCR analysis of The EGFR gene detection kit (Switzerland Roche) was performed using Auantitative real-time PCR (Switzerland Roche Cobas Z 480) and the results were interpreted according to the manufacturer's instructions. With reference to the CAP / IASLC / AMP guideline  $^{(20)}$ , molecular mutations in the EGFR gene were defined as mutations in EGFR exons 18,19,20,21, and other types of EGFR mutations were defined as wild-type EGFR.

The detection steps are as follows: (1) Making amplification agent: the reaction mixture thawed at room temperature, and the thawed reagent, mixed enzyme solution and purified water, followed by centrifugation for 10s, the mixture was made and packaged in each PCR reaction tube, and then added the sample. (2) Sample addition:  $2\mu L$  DNA sample, EGFR negative and positive quality control were successively added to the PCR reaction tube, centrifuged for 10s, and then amplified. (3) PCR amplification: First, the reaction was carried out at

95°C for 5min, cycle once. Second, the reaction was carried out at 95°C, 60°C and 72°C with 30s, and the cycle was 32 times. In the third stage, the cells were incubated at 95°C, 60°C and 72°C for 25s and 10 cycles, respectively. (4) Read the results.

Because this study is a retrospective analysis, the classification of cases is compared according to whether there is the EGFR gene mutation, without detailed analysis of which mutation site, and the gene detection in this study is selected to use the kit provided by Roche. It gives the test results (mutation type) directly by itself. These cases were not verified by gene sequencing. Sequencing maps could not be provided.

#### Statistical analysis

Data were used SPSS version 22.0. The clinical data and CT features of the cases categorized into a mutation group and a wild-type group. The Chi-square with Fisher's exact tests were used to assess statistical significance. Binary logistic regression analysis was guided on statistically significant variables to develop a predictive model for *EGFR* gene mutations in MIA.

#### RESULTS

Clinical Data and EGFR Gene Mutation Status

Among the 118 cases of minimally invasive lung adenocarcinoma, 69 by mutation group, 49 by the wild -type group. This cohort consisted have 67 females (62.3%) and 51 males (37.7%), with no significant in gender (P>0.05). Although the mutation rate was relatively high (66.7%) among the 70 patients more than 60 years, the difference was not significant (P>0.05). Patients were also divided into smokers (n=63) and non-smokers (n=65), no significant difference with smoking history (P>0.05) (table 1).

**Table 1.** Comparison of clinical data between EGFR mutation group and wild-type group in patients with (MIA).

Analytical factor	Cases (n=118)	Mutation Wild group group (n=69) (n=49)		χ2	Р
Age (year)					
< 60	48	23(33.3%)	25(51%)	3.714	0.054
≥60	70	46(66.7%)	24(49%)	3.714	0.054
Gender					
Man	51	26(37.7%)	25(51%)	2.078	0.149
Woman	67	43(62.3%)	24(49%)	2.078	0.149
Smoking					
history					
Yes	63	35(50.7%)	32(65.3%)	2.482	0.115
No	65	34(49.3%)	17(34.7%)		

NOTE: MIA: Minimally Invasive Adenocarcinoma.

#### CT features and EGFR mutation status

The analysis of CT features with the *EGFR* gene mutation status in MIA revealed no statistically significant differences in the average diameter for nodules, density, morphology, lobulation sign, pleural indentation sign, air bronchogram, tumor-lung interface, peripheral emphysema, and peripheral fibrosis (P>0.05). Between the two groups regarding

spiculation sign, vacuole and vascular convergence sign had statistically significant (P<0.05). Partial CT signs with pathological images of MIA illustrated in table 2.

Table 2. Comparison of CT characteristics and EGFR gene mutation status in MIA

mutation status in MIA.								
	All	The EGFR ger						
CT features	nationts		Inspection	p-				
3	(n=118)	Mutation	Wild group	value	Value			
A	,	group (n=69)	(n=49)					
Average								
diameter	26	21 (20 40/)	1F /20 C0/\		0.549 <sup>#</sup>			
≤10mm	36 68	21 (30.4%) 38 (55.1%)	15 (30.6%) 30 (61.2%)	-	u.549"			
≤20mm	14	_ , _ ,						
≤30mm Spiculation sign	14	10 (14.5%)	4 (8.2%)					
	6E	4E /GE 20/\	21 (42.9%)	5.812*	0.016			
Yes No	65 53	45 (65.2%) 24 (34.8%)	28 (57.1%)	3.812	0.016			
Vacuola sign	55	24 (34.8%)	28 (57.1%)					
Yes	62	42 (60 00/\	20 (40.8%)	4.621*	0.032			
Yes No	_	42 (60.9%) 27 (39.1%)	20 (40.8%)	4.021	0.032			
No Density	56	27 (39.1%)	29 (59.2%)					
GGO	60	2E /EO 70/\	22 (67 20/\	3.242*	0.073			
Mixed GGO	68 50	35 (50.7%)	33 (67.3%)	3.242*	0.072			
	50	34 (49.3%)	16 (32.7%)					
Lobulation sign	20	10 /26 10/\	21 //2 00/\	6.642*	0.056			
Yes No	39 79	18 (26.1%)	21 (42.9%)	0.042	บ.บวิธ			
	79	51 (73.9%)	28 (57.1%)					
Morphology Regular	82	46 (66.7%)	36 (73.5%)	0.625*	0.429			
Irregular	36	, ,		0.625*	0.429			
Pleural	30	23 (33.3%)	13 (26.5%)					
indentation sign								
Yes	47	26(37.7%)	21(42.9%)	0.320*	0.571			
No Yes	71	43 (62.3%)	28 (57.1%)					
Air	/1	43 (02.3%)	20 (37.1%)					
bronchial sign					0.202			
Yes	45	23 (33.3%)	22 (44.9%)	1.624*	0.202			
No	73	46 (66.7%)	27 (55.1%)					
Tumor-lung	/3	+0 (00.770)	21 (33.170)					
interface								
distinct	64	36 (52.2%)	28 (57.1%)	0.285*	0.593			
indistinct	54	33 (47.8%)	21 (42.9%)					
Vascular	J-F	33 (47.070)	(-2.3/0)					
convergence sign								
Yes	78	40(58%)	38(77.6%)	4.902*	0.027			
No	40	29 (42%)	11 (22.4%)					
Peripheral		23 (12/3)	( 170)					
emphysema								
Yes	10	3 (4.3%)	7 (14.3%)	-	0.091#			
No	98	66 (95.7%)	42 (85.7%)					
Peripheral	- 50	23 (33.770)	(03.770)					
fibrosis								
Yes	30	16 (23.2%)	14 (28.6%)	0.438	0.508#			
No	88	53 (76.8%)	35 (71.4%)					
.40		33 (70.070)	JJ (11.470)					

Note: 1 # represents the Fisher exact probability test; 2 \* represents the Chi-square test; GGO: Ground Glass Opacity; MIA: Minimally Invasive Adenocarcinoma.

Chi-square tests revealed that the spiculation sign and vascular convergence sign in MIA were statistically (P<0.05). These signs were included in a binary logistic regression analysis to make up a predictive model for *EGFR* in MIA. This analysis demonstrated that vascular convergence sign and spiculation sign are independent risk factors for predicting *EGFR* gene mutations. ROC curves were generated based on the spiculation and vascular convergence signs. AUC for spiculation sign is 0.634, with sensitivity of 85.02% and a specificity of 35.21%. The AUC for the vascular convergence sign was 0.735, with a sensitivity of 84.48% and a specificity of 37.56% (table 3, figure 3).

**Table 3.** Results of Logistic regression analysis for minimally invasive adenocarcinoma (MIA).

Variable	В	S.E.	Wald	P	OR	95%CI	
vascular	1 160	0.729	4.353	0.026	2.262	1.030, 4.547	
convergence sign						· ·	
vacuole sign	-0.443	0.965	0.529	0.521	0.410	0.339, 4.212	
spiculation sign	1.423	0.995	2.092	0.017	1.498	0.247, 4.903	

NOTE: MIA: Minimally Invasive Adenocarcinoma. B: The regression coefficient reflects the influence of the independent variable on the dependent variable; S.E. The standard error reflects the estimation accuracy of the regression coefficient, and the smaller the standard error, the more accurate the estimation of the regression coefficient; Wald: The Wald statistic was used to test whether the regression coefficients were significantly not 0; P: Reflecting the significant level of this test, with p <0.05, this independent variable was considered to have a significant influence on the dependent variable; OR: The image degree index reflecting the variable for the classification outcome.

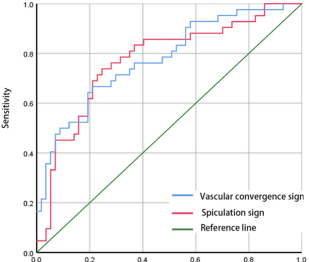


Figure 3. Ensure that the receiver operating characteristic curve (ROC) analysis of the spiculation sign and vascular convergence sign in predicting minimally invasive adenocarcinoma (MIA) is of high quality.

#### **DISCUSSION**

The pathogenesis of lung adenocarcinoma is complex, involving many factors such as genetics, chronic lung infection and environmental pollution. Its malignant degree is high. If timely screening and treatment cannot be done, tumor invasion and distant metastasis can appear with the progression of the disease, and the mortality rate is high (21). The rapid advancements and widespread application of imaging technologies have greatly improved the detection rate of early-stage lung cancer. Imaging techniques can partially reflect the molecular characteristics of tumors, and predicting EGFR gene mutation status through imaging examinations has become an important research field (22). However, the relationship between imaging characteristics with the EGFR gene mutations of lung adenocarcinoma remains in the early stages of research, leading to inconsistent conclusions across studies (15, 17, 18).

Regarding clinical characteristics, some studies suggest that models predicting *EGFR* mutations based on age, gender, and smoking history alone are less accurate <sup>(23, 24)</sup>. Consequently, there are limitations to

using these factors exclusively to predict *EGFR* gene mutations, which aligns with the findings of this study. However, many studies have demonstrated that young, non-smoking women are more likely own the EGFR mutable gene in lung adenocarcinoma <sup>(16, 25)</sup>. These discrepancies could be attributed to differences in clinical characteristics, regional variations, and racial differences among the study populations.

The study pointed out that the size of the lesion was connected with the pathological process. Most solid lesions greater than 3cm were invasive lung adenocarcinoma, while pure ground glass nodules less than 1.5cm in diameter had a low incidence of invasive lung adenocarcinoma (26). In terms of lesion diameter from CT quantitative analysis, this study is consistent with most reports, showing no correlation between lesion size and EGFR gene mutation status (27, 28). However, Zhang et al. observed that as the average lesion diameter increases, indicating a positive correlation between lesion size and EGFR mutations  $(29)_{.}$ This study had microinvasive adenocarcinoma lesions less than 3cm long, which differs from studies focusing on invasive lung adenocarcinoma, where most lesions exceed 3cm. This difference in case selection may account for the divergent conclusions. Regarding the indentation sign on CT, this study agrees with Song's findings, which concluded that there are no relationship EGFR mutations with pleural indentation (30). Pleural indentation is primarily caused by reactive fibroplasia or scar formation within the lesion, pulling the local pleura and forming a cable-like shadow. It is commonly associated with pleural invasion, indicating a more aggressive tumor with a poorer prognosis (31, <sup>32)</sup>. In contrast, Nie *et al.* found that pleural indentation was more prevalent in patients with EGFR mutations, suggesting it may have some predictive value (33). However, their study included more cases of invasive lung adenocarcinoma, which could explain the difference in findings, as this study focused on microinvasive lung adenocarcinoma with a lower malignancy rate. Most studies suggest that signs such as vacuolation, spiculation, and vascular convergence correlated with EGFR gene mutations (34, 35). Huo liwen et al. identified spiculation and vascular convergence as independent predictors of EGFR mutations in the study of 803 cases, consistent with these results (34). However, Juna et al. found no correlation between these CT signs and EGFR gene expression (36). Their study, which included 354 lung adenocarcinoma lesions, showed that invasive lung adenocarcinoma was more prone to EGFR mutations, further supporting the relationship between EGFR mutations and tumor infiltration (37). These differences in pathological results and case selections between studies may contribute to the varying findings. This study did not identify correlations between other CT characteristics and EGFR mutations, possibly due to subjective differences, staging variations, ethnic

differences, and researchers' judgment in evaluating morphological signs.

The main limitations of this study were the insufficient quantitative analysis of lesions. Future research should involve larger sample sizes, incorporate additional CT features and imaging omics data, and utilize digital image processing technology to analyze the morphology of lung adenocarcinoma lesions quantitatively. This approach will further explore the relationship between gene mutations and imaging features.

## **CONCLUSION**

CT imaging of microinvasive lung adenocarcinoma revealed a high rate of *EGFR* gene mutations in cases with spiculation, vacuolation, and vascular convergence signs. Spiculation and vascular convergence signs identified as independent risk factors for predicting *EGFR* gene mutations.

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**Conflicts of Interest:** All authors unanimously declare no conflict of interest in this study.

**Ethical Consideration:** This retrospective study was approved by the research Ethics Committee of Tangshan Gongren Hospital with the number GRYY-LL -2018-48 at 26th Nov 2018.

**Author Contribution:** L.P. Z. and L.M.C.; conceived and designed the study. H.Z. and J.T.W.; conducted the literature search and data collection. L.L.S.; analyzed the data. L.P.Z. and L.M.C.; wrote the paper. L.P.Z. and X.Y.Y.; reviewed and edited the manuscript. All authors read and approved the final manuscript.

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