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CASE REPORT

Coexistence of Low-Grade Fetal Adenocarcinoma and Adenocarcinoma in situ of the Lung Harboring Different Genetic Mutations: A Case Report and Review of Literature

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Abstract: Low-grade fetal lung adenocarcinoma (L-FLAC) is an exceptionally rare pulmonary tumor, presenting with unclear histological and molecular features. In particular, the potential driver genes of L-FLAC remain elusive. To date, only five reports have documented genetic aberrations in L-FLAC. In the current study, we describe an unusual case of L-FLAC coexisting with adenocarcinoma in situ (AIS) of the lung, harboring different genetic mutations. A 39-year-old non-smoker female patient was referred to our hospital with the chief complaint of dyspnea for 20 days. Chest computed tomography (CT) revealed a - $2.5 \times 1.5 \times 1.5$  cm nodule in the right middle lobe, with no mediastinal lymph node enlargement or distant metastases. Thoracoscopic surgery was performed to remove the nodules. Histopathological analysis of the tissue sections, based on findings from immunohistochemical staining, confirmed a diagnosis of L-FLAC coexisting with AIS of the lung. Nextgeneration sequencing revealed L-FLAC-based mutations in DICER1 and CTNNB1, and AIS harboring KRAS mutations. Currently, the patient remains recurrence-free 17 months after the initial diagnosis. This report presents the first case demonstrating the coexistence of L-FLAC and AIS in the same pulmonary nodule, harboring different genetic mutations. Based on the literature review, this is the second reported case of L-FLAC bearing *DICER1* mutations. Keywords: low-grade fetal adenocarcinoma, CTNNB1, DICER1, KRAS, NGS

### Introduction

Fetal lung adenocarcinoma is markedly rare and was first reported in 1982.<sup>1</sup> An extremely very rare lung cancer, it accounts for only 0.1–0.5% of all lung neoplasms. In 1999, this tumor was classified as a variant of lung adenocarcinoma by the World Health Organization (WHO) and is currently termed as "fetal adenocarcinoma of the lung" (FLAC).<sup>2</sup> The tumor mass is generally solitary, well-circumscribed, and located peripherally in the lung. Typically, FLAC causes clinical symptoms such as chest pain, dyspnea, cough, and hemoptysis. Additionally, some asymptomatic patients are diagnosed with incidental findings on chest radiographs. According to the histological patterns, the tumor was further classified into two subtypes: low-grade FLAC (L-FLAC) and high-grade FLAC (H-FLAC).<sup>3</sup> Furthermore, these two subtypes present different clinical characteristics and outcomes. Patients with H-FLAC generally present advanced-stage diseases and a poorer prognosis. Nakatani et al have reported that L-FLAC, histologically, is pure in pattern and may present an even better

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#### **Case Report**

A 39-year-old non-smoker female patient, predominantly complaining of dyspnea for 20 days, was referred to our hospital. Hematological, biochemical, and sputum tests showed no abnormalities. Chest computed tomography (CT) revealed a  $2.5 \times 1.5 \times 1.5$ cm nodule in the right middle lobe, with no mediastinal lymph node enlargement or distant metastases (Figure 1). Following lobectomy and mediastinal lymph node dissection, an intraoperative frozen section was examined and diagnosed as lung adenocarcinoma.

The resected tumor presented a smooth, whitish, and firm surface, with small hemorrhagic areas, and appeared to be relatively well-circumscribed. No obvious necrosis or cystic changes were identified. Microscopically, the tumor tissue contained two distinct non-adjacent lesions. One lesion demonstrated complex glandular and tubular structures, with prominent morule formation, resembling the developing fetal lung in its pseudo glandular stage. The tumor cells displayed low nuclear atypia, and no necrosis was detected (Figure 2A and B). Based on these findings, a diagnosis of L-FLAC was proposed. In the second lesion, the tumor cells were distributed alongside the alveolar structures, without stromal, vascular, and alveolar space invasion, presenting a maximum diameter of only 2 mm (Figure 2C and D). Therefore, this lesion was deemed an AIS of the lung. Although these two lesions were located within the same nodule, they were individually present in different tissue sections. After screening all tissue sections, no spindle-shaped tumor cells or sarcomatous structures were observed. Furthermore, no vessel invasion or pleural involvement was detected.

Immunohistochemically, the L-FLAC cells presented diffuse staining for cytokeratin, thyroid transcription factor-1 (TTF-1, Figure 3A), epithelial membrane antigen (EMA), and sal-like protein 4 (SALL4, Figure 3B). Chromogranin A (CgA, Figure 3C) and synaptophysin (Figure 3D) were only focally stained. The tumor cells were negative for cytokeratin 7,  $\alpha$ -fetoprotein, CD56, and glypican 3. Notably,  $\beta$ -catenin showed diffuse staining on the cell membrane, with only sporadic nuclear/cytoplasmic expression (Figure 3E). Additionally, the  $\beta$ -catenin expression pattern was similar in the morule and non-morule structures. The Ki-67 index was approximately 40% (Figure 3F).

To further identify the genetic mutation profile, we performed NGS on both the L-FLAC and AIS regions, using adjacent normal lung tissue as the germline control. Molecular studies have revealed that these two lesions harbored distinct genetic mutations (Supplementary Figure 1).



Figure I Computed tomography (CT) scans of the chest. (A and B) Photographs present a well-circumscribed and lobulated 2.5-cm nodule in the right middle lobe.



Figure 2 Histological manifestations of L-FLAC and AIS. L-FLAC shows complex glandular structures, with prominent morule formation (A, 100×; B, 200×). The AIS presents tumor cells distributed along alveolar structures without stromal, vascular, and alveolar space invasion (C, 40×; D, 100×).

L-FLAC revealed somatic mutations in *DICER1* frameshift mutation in the exon 23 (c.4407\_4410delTTCT (p. S1470Lfs\*19)) with an abundance of 31.9%, and missense mutations in exon 24 (c.5125G>A p.D1709N) with an abundance of 38.3%. Simultaneously, missense mutations were identified in *CTNNB1* in exon 3 (c.98C>G p.S33C) with an abundance of 41.2%. However, AIS demonstrated a *K-RAS* missense mutation in exon 2 (c.35G>C p.G12A). Finally, the diagnosis of L-FLAC coexisting with AIS stage 1A (pT1bN0M0) was confirmed. Postoperatively, no evidence of recurrence was observed for 17 months.

#### Discussion

As a distinct entity according to the WHO classification, FLAC is a malignant tumor with extremely low incidence, accounting for less than 0.5% of all lung tumors.<sup>2</sup> The two FLAC subtypes, L-FLAC and H-FLAC, differ in multiple aspects, including histopathology and clinical characteristics.<sup>5</sup> L-FLAC is commonly detected in female patients in the third-fourth decade of their lives. Typically, the initial diagnosis of L-FLAC occurs during the early stages (I–II).<sup>6</sup> H-FLAC mostly occurs in elderly male smokers, presenting advanced-stage disease and severe symptoms.<sup>7</sup>

L-FLAC showed a pure histological pattern, with prominent morule formation and low nuclear atypia. However, H-FLAC, generally displays fetal morphology (over 50%), together with other common types of lung adenocarcinomas.<sup>8</sup> Moreover, H-FLAC commonly demonstrates prominent nuclear atypia and nucleoli, accompanied by necrosis and a high frequency of mitosis. In the current case, no prominent nuclear atypia, necrosis, or a direct transition to conventional adenocarcinoma was observed. Accordingly, a diagnosis of L-FLAC, rather than H-FLAC, was confirmed.

In addition to histopathological patterns, these two subtypes demonstrate distinct immunohistochemical staining. Owing to the activation of the WNT signaling pathway, typically, but not in all cases, L-FLAC can display aberrant cytoplasm/nuclear distribution of  $\beta$ -catenin.<sup>9</sup> In H-FLAC, the  $\beta$ -catenin expression is usually localized on the cell membranes. L-FLAC presents lower expression of p53, which in contrast, is commonly overexpressed in H-FLAC. Morita et al and Suzuki et al have reported that  $\alpha$ -fetoprotein, glypican 3, and SALL4 are frequently expressed in H-FLAC, especially in non-fetal-like components.<sup>7,8</sup> However, in the current case, SALL4 also demonstrated diffuse expression. Therefore, immune profiles can be distinct in different cases. As few studies have



Figure 3 Immunophenotype of L-FLAC lesions. Immunohistochemically, the L-FLAC cells present diffuse staining of TTF-1 (A, 100×) and SALL4 (B, 100×); focal expression of CgA (C, 100×) and synaptophysin (D, 100×); and predominant membranous  $\beta$ -catenin expression with only sporadic nuclear/cytoplasmic staining (E, 200×); The Ki-67 index is approximately 40% (F, 100×).

documented SALL4 expression in L-FLAC, further research is crucial to elucidate whether SALL4 can be utilized as a useful marker in the differential diagnosis of H-FALC and L-FLAC. We speculated that SALL4 expression was associated with the *DICER1* mutation, as shown in the current study. Several studies have reported that *DICER1*-associated malignancies, especially with blastomaoid components, show SALL4 expression.<sup>10,11</sup> However, owing to the limited data available, this interpretation needs further supportive evidence.

In the current case, the  $\beta$ -catenin demonstrated only sporadic cytoplasmic/nuclear localization in tumor cells and was mostly present on cell membranes. Similar expression patterns of  $\beta$ -catenin have been reported in L-FLAC in

a 16-year-old female patient by both Wu et al<sup>12</sup> and de Kock et al.<sup>13</sup> It worth noticing that, in that case, the L-FLAC also harbored *DICER1* mutations in addition to *CTNNB1* mutations. L-FLAC is considered as a putative precursor lesion of pulmonary blastoma (PB), which is a biphasic tumor consisting of FLAC (typically low-grade) and primitive mesenchymal stroma. Both L-FLAC and PB are frequently associated with mutations of *CTNNB1*.<sup>9</sup> Interestingly, till date, mutations in both *DICER1* and *CTNNB1* have only been identified in two cases of PB.<sup>14</sup> In these two cases, βcatenin also demonstrated predominantly membranous expression with multifocal cytoplasmic/nuclear localization. Collectively, these cases further support that L-FLAC and PB shared partially overlapping genetic abnormalities, and

Author	Age/ Sex	β-Catenin Expression	Size	Genetic Mutation
Nakatani et al, 2002 <sup>4</sup>	N/A N/A	Nuclear/cytoplasmic expression nuclear/cytoplasmic expression	N/A N/A	CTNNB1 (the exon 3, c.101G>T, p.G34V) CTNNB1 (the exon 3, c.110C>G, p.S37C)
Sekine et al, 2003 <sup>15</sup>	32/F 22/F 23/M	Nuclear/cytoplasmic expression Nuclear/cytoplasmic expression Nuclear/cytoplasmic expression	3.2cm 2.8cm 3.0cm	CTNNB1 (the exon 3, c.94G>T, p.A32T) CTNNB1 (the exon 3, c.98C>G, p.S33C) CTNNB1 (the exon 3, c.110C>G, p.S37C)
Wu et al, 2014; <sup>12</sup> de Kock et al, 2016 <sup>13</sup>	16/F	Mostly membranous, with multifocal nuclear/cytoplasmic expression	6.7cm	CTNNB1 (the exon 3, c.98C>G, p.S33C) DICER1 (the exon 21, c.3540C>A, p.Tyr1180; the exon 24, c.5127T>A, p.A1709G)
Fu et al, 2018 <sup>16</sup>	31/M 21/M	Nuclear/cytoplasmic expression Nuclear/cytoplasmic expression	3.0cm 5.8cm	CTNNB1 (the exon 3, c.110C > A, p.S37Y) BRCA2 (the exon 10, c.1871C>T, p.C624V) TSC2 (the exon 12, c.1222T>G, p.F408V) CTNNB1 (the exon 3, c.110C > G, p.S37C) DDR2 (p.E499)
Present case	39/F	Diffusive membranous, with sporadic nuclear/cytoplasmic expression	2.5cm	CTNNB1 (the exon 3, c.98C>G, p.S33C) DICER1 (the exon 23, c.4407_4410delTTCT, p. S1470Lfs*19; the exon 24, c.5125G>A, p.D1709N)

Table I Clinical Review of 6 Reports of L-FLAC with Genetic Analysis

mutations in *DICER1* may be associated with aberrant  $\beta$ -catenin expression patterns.

Besides mutations in CTNNB1, genomic mutation profiles in L-FLAC remain elusive owing to extremely low incidences and reports. To date, only six reports have documented the genetic mutations in L-FLAC, including the current study, and these findings are summarized in Table 1.4,12,13,15,16 By reviewing the literature, although uncommon mutations in BRAC2, TSC2, and DDR2 have been identified, mutations in CTNNB1 demonstrated the highest frequency in L-FLAC. In the current case, although AIS was co-existed with L-FLAC in the same pulmonary nodule, these two lesions harbored different mutations, indicating that both tumors were primary lesions, instead of intrapulmonary metastasis. We speculated that AIS within the same nodule of L-FLAC can be interpreted as intra-tumor spatial heterogeneity owing to different driver genes. If the lesion was not identified and resected during the early stages, the AIS region may progress to invasive adenocarcinoma. Therefore, this case might elucidate an early event of the development of H-FLAC, in which the fetal component presents with a transition to conventional invasive adenocarcinoma.

In conclusion, we report an extremely rare case of a 39-year old woman with L-FLAC and AIS within the same pulmonary nodule. The two lesions demonstrated distinct genetic mutations. This is the second reported case of L-FLAC bearing *DICER1* mutations. Currently, the patient remains asymptomatic and disease-free following a 17-months follow-up period.

# **Data Sharing Statement**

The datasets supporting the conclusions of this article are included within the article.

# Ethical Approval and Consent to Participate

Ethical approval for this study was obtained from the institutional ethic review boards of the First Affiliated Hospital of China Medical University. Writing consent to participate was provided by the patients for the present research.

# **Patient Consent for Publication**

The informed consent was obtained from the patient for the publication of her case and any accompanying images. A copy of the written consent is available for review by the Editor-in-Chief of this journal.

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### **Author Contributions**

All authors made substantial contributions to conception and design, acquisition of data, or analysis and interpretation of data; took part in drafting the article or revising it critically for important intellectual content; gave final approval of the version to be published; and agree to be accountable for all aspects of the work.

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## Disclosure

The authors declare that they have no competing interests.

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