

[CASE REPORT]

Lenvatinib-induced Interstitial Pneumonia in a Patient with Hepatocellular Carcinoma

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Abstract:

Lenvatinib is a multi-targeted tyrosine kinase inhibitor available for the treatment of unresectable hepatocellular carcinoma (HCC). We herein report an 84-year-old-man with interstitial pneumonia caused by lenvatinib. Four months after the start of lenvatinib administration for HCC, chest computed tomography revealed bilateral ground-glass opacity. However, he continued to take lenvatinib for four more months until he complained of dyspnea on exertion. This is a case of lenvatinib-induced interstitial pneumonia that progressed relatively slowly with a long asymptomatic period despite the appearance of pneumonia on image findings.

Key words: lenvatinib, interstitial pneumonia, hepatocellular carcinoma

(Intern Med 61: 1211-1217, 2022) (DOI: 10.2169/internalmedicine.7300-21)

Introduction

Lenvatinib is a multi-targeted tyrosine kinase inhibitor that blocks tumor cell proliferation and angiogenesis mainly by inhibiting vascular endothelial growth factor receptor (VEGFR) 1, 2, and 3; fibroblast growth factor receptor (FGFR) 1, 2, 3, and 4; platelet-derived growth factor receptor (PDGFR) α ; RET; and KIT (1, 2). In Japan, lenvatinib was approved as a treatment for unresectable thyroid cancer (3, 4) and unresectable hepatocellular carcinoma (HCC) (5). Only a few cases among 1,343 patients included in clinical trials presented with lenvatinib-induced interstitial pneumonia (3-5). However, a considerable number of cases of interstitial pneumonia were reported in a post-marketing survey of lenvatinib (6) (Table 1).

The characterization of lenvatinib-related interstitial pneumonia is poorly understood at present, and there are only two case reports of lenvatinib-induced interstitial pneumonia (7, 8). In these previous reports, lenvatinib administration was immediately discontinued after the appearance of interstitial pneumonia.

We herein report a relatively slowly progressive case of lenvatinib-induced interstitial pneumonia. The patient continued to take lenvatinib for a long time due to the absence of suitable alternative medications.

Case Report

An 84-year-old-man was a former smoker of 1 pack of cigarette per day for 24 years and had a history of hypertension and chronic kidney disease. He was referred to the Department of Gastroenterology in our hospital for the examination of a tumor lesion in the right lobe of the liver and was diagnosed with HCC. Drug-eluting bead transcatheter arterial chemoembolization or transcatheter arterial chemoembolization with cisplatin was performed for HCC a total of six times over the next seven months.

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Received: February 15, 2021; Accepted: July 29, 2021; Advance Publication by J-STAGE: September 18, 2021 Correspondence to Dr. Yasuhiko Nishioka, yasuhiko@tokushima-u.ac.jp

Age	Sex	Primary disease	Time from starting lenvatinib to the onset of drug-induced interstitial pneumonia (day)	Outcome	Time from onset to outcome (day)
70s	М	HCC	28	death	3
70s	М	HCC	43	death	16
70s	М	HCC	46	death	27
50s	М	HCC	9	death	3
70s	М	TC	16	death	3
60s	М	HCC	40	recovered	6
70s	F	HCC	6	recovered	unknown
70s	М	HCC	87	recovered	2
80s	М	HCC	109	recovered	17
70s	М	HCC	63	recovered	103
70s	F	HCC	125	recovered	11
60s	М	TC	51	recovered	9
60s	М	TC	30	recovered	167

Table 1. Summary of Post-marketing Survey which was Conducted toHepatocellular Carcinoma (HCC) or Thyroid Cancer (TC) Patients Ad-ministered Lenvatinib Cited from Guide to Proper Use of Lenvatinib.

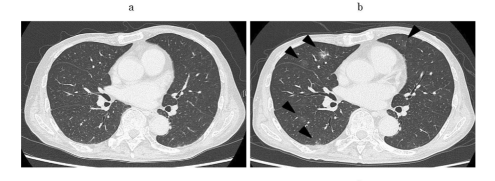
After first-line therapy failed, the treatment for HCC was changed to sorafenib and then regorafenib. Eight months after starting regorafenib, high-resolution computed tomography (HRCT) revealed enlargement of the HCC with worsening of the hepatic function. At this time, HRCT showed no abnormality (Fig. 1a). The chemotherapy was changed to 80 mg/day of lenvatinib. Three months later, the dosage of lenvatinib was decreased to 40 mg/day due to palmar-plantar erythrodysaesthesia. Four months after starting lenvatinib, HRCT showed the appearance of focal ground-glass opacity (GGO) in both lung fields (Fig. 1b). Although the patient had no respiratory symptom, two months later, the spread of GGO on HRCT was observed, with accompanying findings of infiltration shadows. At this point, lenvatinib-induced interstitial pneumonia was assumed, and the pneumonia was considered to be Grade 1 pneumonitis according to the National Cancer Institute Common Terminology Criteria for Adverse Events (CTCAE) v5.0.

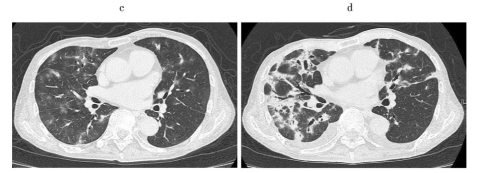
Because there were no other treatments approved for HCC that had never been tried to him, lenvatinib was continued with careful observation under his consent. Two months later, the spread of GGO was observed on HRCT, along with infiltration shadows (Fig. 1c). Nevertheless, because he still had no respiratory symptoms and strongly requested to continue lenvatinib, the administration of lenvatinib was continued under very careful observation. Two months later, he complained of exertional dyspnea. The HRCT findings now showed distinct patchy infiltration shadows in both lung fields (Fig. 1d). He was referred to our department after lenvatinib discontinuation and was admitted.

On admission, he complained dyspnea on exertion, with a Modified British Medical Research Council (mMRC) grade of 2. His initial vital signs were as follows: blood pressure, 103/63 mmHg; body temperature, 36.0 °C; percutaneous oxygen saturation (SpO₂), 96%. Resting-state partial pressure of arterial oxygen (PaO₂) while breathing room air was 83.1 mmHg. Laboratory test showed: Krebs von den Lungen-6 (KL-6), 1,906 U/mL (normal value: <500 U/mL); surfactant protein D (SP-D). 230 ng/mL (normal value: <100 ng/mL); C-reactive protein (CRP), 1.74 mg/dL (Table 2). No elevated levels of autoantibodies were observed (Table 2). He did not meet the criteria for the diagnosis of any connective tissue diseases. Cytomegalovirus antigens were negative (Table 2). Lung sounds were normal on auscultation. Other physical examination findings were also normal.

Bronchoalveolar lavage (BAL) of the left middle lobe showed a mild increase in the lymphocyte rate (lymphocyte: 20.8%) (Table 3). A transbronchial lung biopsy (TBLB) of the left upper lobe with Hematoxylin and Eosin (H & E) staining showed infiltration of lymphocytes into the alveolar interstitium but no thickening of the alveolar wall or Masson's body (Fig. 2a, b). Fibrosis was not evident on Elastica van Gieson (EVG) staining (Fig. 2c, d). DLST was positive (599 cpm, stimulation index 2.5; normal range: lower than 1.8), suggesting that the present case had an allergic reaction to lenvatinib. Given these results, a diagnosis of druginduced interstitial pneumonia caused by lenvatinib was made.

Following intravenous pulse steroid therapy with 500 mg of methylprednisolone (mPSL) for 3 days, we started the oral administration of 25 mg/day (0.5 mg/kg) of prednisolone (PSL). After starting corticosteroid treatment, his respiratory condition improved with a reduction in the KL-6 value (Fig. 3). Three months after starting corticosteroid therapy, the radiological findings gradually improved (Fig. 1e). However, after he stopped the administration of lenvatinib, his HCC gradually worsened with an increase in





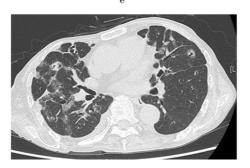


Figure 1. (a) High-resolution chest tomography (HRCT) image at the start of lenvatinib. There was no shadow suggestive of pneumonia in the lung field. (b) HRCT image at four months after starting lenvatinib. Ground-glass opacity (GGO) appeared in both lung fields (arrowheads). (c) HRCT image at six months after starting lenvatinib. The spread of GGO was observed, with the accompaniment of some infiltration shadows. (d) HRCT image at eight months after starting lenvatinib. Patchy infiltration shadow appeared in both lung fields. (e) HRCT image at three months after starting corticosteroid therapy. The infiltration shadow had decreased.

the alpha-fetoprotein (AFP) and protein induced by vitamin K absence or antagonist (PIVKAII) values. He ultimately passed away four months after starting corticosteroid therapy due to exacerbation of HCC.

Discussion

The mechanism underlying lenvatinib-induced interstitial pneumonia has not been clarified yet. In general, direct toxicity of the causative drug can be responsible for druginduced pneumonia (9). Some previous reports have suggested that the induction of alveolar epithelial cell apoptosis by VEGF receptor inhibition might be involved in the development of interstitial pneumonia (10-12). Similarly, one previous case report of lenvatinib-induced interstitial pneumonia stated that it was induced by alveolar epithelial cell apoptosis following VEGF receptor inhibition (8). However, drug-induced pneumonia is often caused by immunemediated reaction (9). The DLST of lenvatinib was positive in both the present case and a previous case (Table 4), suggesting that an allergic reaction to lenvatinib may also cause drug-induced interstitial pneumonia.

The typical imaging findings of lenvatinib-induced interstitial pneumonia are unclear. In two previous reports, HRCT showed GGO (Table 2) (7, 8). In the present case as well, the shadow of GGO was initially the main component. However, the HRCT findings changed to distinctive patchy alveolar opacities resembling a fibrosing organizing pneumonia pattern when the lenvatinib-induced interstitial pneumonia worsened. After the corticosteroid therapy, abnormal interstitial thickening remained despite the bilateral alveolar opacities gradually disappearing, suggesting that progressive

WBC	6,500 /µL	Na	140 mEq/dL
Neut	71.6 %	Κ	4.8 mEq/dL
Eo	3 %	TP	5.5 g/dL
Baso	0.3 %	Alb	2.5 g/dL
Mono	9.8 %	CRP	5.33 mg/dL
Lymp	12.4 %	β D-Glucan	<6.0 pg/mL
RBC	3.27×10 ⁴ /μL	KL-6	1,906 U/mL
Hb	9.5 g/dL	SP-D	230 ng/mL
Plt	25.9×10 ⁴ /μL	RF	<10 IU/mL
AST	46 U/L	ANA	<40 times
ALT	18 U/L	anti-SS-A antibody	(-)
ALP	659 U/L	PR3-ANCA	<1.0 IU/mL
LDH	189 U/L	MPO-ANCA	<1.0 IU/mL
γ-GTP	104 U/L	CMV anitigen*	0 cells/2 slides
BUN	22 mg/dL	T-SPOT	(-)
Cre	1.29 mg/dL		

Table 2. The Serum Test Results at the Time of Admission toOur Department.

WBC: white blood cell, Neut: neutrophil, Eo: eosinophil, Baso: basophil, Mono: monocyte, Lymp: lymphocyte, RBC: red blood cell, Hb: hemoglobin, Plt: platelet count, AST: aspartate aminotransferase, ALT: alanine aminotransferase, ALP: alkaline phosphatase, LDH: lactate dehydrogenase, γ -GTP: γ -glutamyl transpeptidase, BUN: blood urea nitrogen, Cre: creatinine, TP: total protein, Alb: albumin, CRP: C-reactive protein, β D-Glucan: beta-d-glucan , KL-6: krebs von den lugen-6, SP-D: surfactant protein D, RF: rheumatoid factor, ANA: antinuclear antibody, anti-SS-A antibody: anti-Sjögren's-syndrome-related antigen A autoantibodies, PR3-ANCA: proteinase3 antineutrophil cytoplasmic antibody, MPO-ANCA: myeroperoxidase antineutrophil cytoplasmic antibody, CMV antigen: cytomegalovirus antigen, T-SPOT: T-SPOT®-TB

* cytomegalovirus, viral antigen (pp65. C10, C11)

Table 3.The Results of Bronchoalveo-lar Lavage Fluid (BALF).

Total cell counts	22.5×10 ⁴ /mL
cell fractionation	
alveolar macrophages	74.3 %
lymphocytes	20.8 %
neutrophils	4.1 %
eosinophils	0.8 %
basophils	0 %
CD4/8 ratio*	0.59
Cytodiagnosis	Class II
BALF cultures	negative

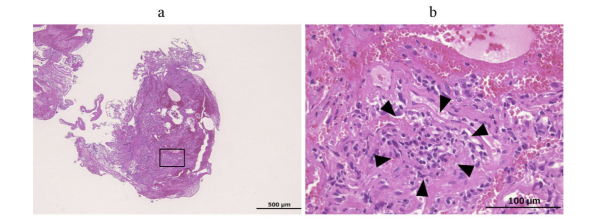
*CD4+/CD8+T-lymphocyte ratio

fibrosis might have occurred in the lungs (Fig. 1e). Although the fibrosis was not evident on EVG staining, this might have been due to the small size of specimen collected by the TBLB. Therefore, if lenvatinib had been discontinued earlier, the abnormal shadow might have disappeared without leaving scars in the lung.

However, in the present case, lenvatinib was continued when drug-induced pneumonitis was suspected six months after the initiation of therapy. Consequently, the progressionfree survival (PFS) of lenvatinib for HCC was about 8 months, and the overall survival (OS) for HCC was about 12 months. These data are compatible with the results in a phase 3 study of lenvatinib for unresectable HCC, showing that the PFS and OS for first-line chemotherapy were 7.4 and 12.5 months, respectively (5). However, in the guide to the proper use of lenvatinib, drug discontinuation and a close examination, including laboratory and imaging tests, are recommended when drug-induced pneumonia is suspected (6). Therefore, in the present case, the discontinuation of lenvatinib at least six months after starting its therapy should have been selected.

In 9 out of 13 cases reported in the post-marketing survey, lenvatinib-induced interstitial pneumonia was observed within 2 months after starting the administration of lenvatinib (Table 1). Among these nine acute or subacute progressive cases, five died within one month after the onset of pneumonia. In contrast, the other four cases that took more than two months to develop pneumonia eventually recovered (Table 1). In the present case, fortunately, the lenvatinib-induced interstitial pneumonia in our patient recovered with corticosteroid treatment. This good responsiveness to treatment may be related to the slow progression of pneumonia in the present case.

In summary, we described a case of slowly progressive interstitial pneumonia induced by lenvatinib. The Japanese Respiratory Society (JRS) guidelines for the management of drug-induced lung disease recommends the suspected drug



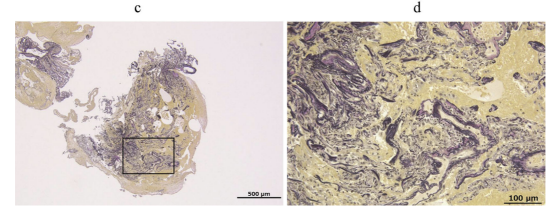


Figure 2. Histological findings of the transbronchial lung biopsy specimen. A histological examination was performed with Hematoxylin and Eosin staining (a, b) and the Elastica van Gieson staining (c, d). The square area of the low-magnification photomicrograph (a, c) is shown as a high-magnification photomicrograph (b, d) with each stain. The arrowheads indicate the infiltration of lymphocyte into the alveolar interstitium. A scale bar is shown in each figure.

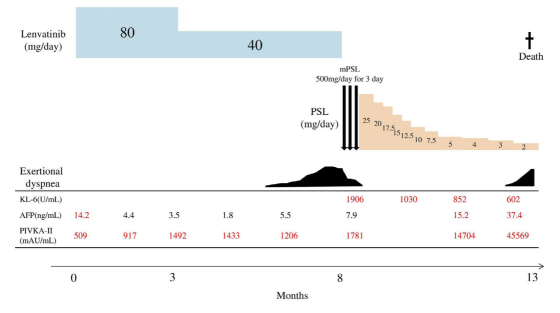


Figure 3. Clinical course of the patient after starting lenvatinib. AFP: alpha-fetoprotein, KL-6: Krebs von den Lugen-6, mPSL: methylprednisolone, PIVKA-II: protein induced by vitamin K absence or antagonist II, PSL: prednisolone

Table 4. Su	umma	ıry of	the Patient	t with	the Three	e Lenvatinib-induc	Table 4. Summary of the Patient with the Three Lenvatinib-induced Interstitial Pneumonia Reported in the Literature.	nonia Reporte	d in the I	iterature.				
Case of Study [referene]		Age Sex	Cancer type	PS	Smoking history (pack year)	Complication of interstitial pneumonia before the start of lenvatinib	Time from first administration to the onset of drug-induced interstitial pneumonitis (month)	Pattern of HRCT	(U/mL) KL-6	Lymphocyte ration of BAL(%)	TBLB	DLST	treatment	outcome
Case Rep Oncol 11:75-80, 2018 [7].	67	67 M	Cncer of unknown primary (Squamous cell carcinoma)	0	15	no complication	-	bilateral GGO	582	61.10%	no data	positive	positive discontinuation recoverd of lenvatinib	recoverd
Clin J Gastroenterol 12: 355-360, 2019 [8].	59	Μ	нсс	0	45	complication of mild interstitial pneumonitis induced by regoratenib	-	bilateral GGO	1283	no data	no data	negative	negative discontinuation recoverd of lenvatinib steroid therapy	recoverd
present case	84	М	нсс	-	24	no complication	4	bilateral GGO 1906 ⇒ bilateral infiltration shadow	1906	20.80%	infiltration of positive lymphocyte into alveolar interstitium	positive	discontinuation recoverd of lenvatinib steroid therapy	recoverd
BAL: bronchoal	veolar	lavage,	, DLST: drug	induce	ed lymphocy	te stimulation test, GGC	BAL: bronchoalveolar lavage, DLST: drug induced lymphocyte stimulation test, GGO: ground-glass opacity, HCC: hepatic cell carcinoma, HRCT: high-resolution chest tomography, KL-6: Krebs von den Lugen-6,	ICC: hepatic cell	carcinoma,	HRCT: high-re	solution chest to	mography, H	KL-6: Krebs von d	en Lugen-6,

be discontinued soon after the pneumonia is determined to have been drug-induced, as a principle (13). However, according to the characteristics of interstitial pneumonia, several drugs, such as sirolimus, everolimus, pembrolizumab, atezolizumab, and durvalumab, can be continued when the grade of drug-induced pneumonia is Grade 1 (asymptomatic) (14-18). In the case of lenvatinib, although the discontinuation is generally recommended when drug-induced interstitial pneumonia is suspected, further information on more cases would enable the development of a precise management strategy depending on each case.

The authors state that they have no Conflict of Interest (COI).

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