CASE REPORT

Sudden Death in a Patient with Pulmonary Veno-occlusive Disease (PVOD) and Severe Pulmonary Hypertension

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Abstract

A 58-year-old woman was referred to our hospital with a chief complaint of exertional dyspnea. Bronchoscopy failed to establish a diagnosis, and the patient subsequently died suddenly due to respiratory insufficiency because of advanced pulmonary hypertension (PH). The pathological diagnosis at autopsy was pulmonary veno-occlusive disease (PVOD). PVOD is difficult to diagnose antemortem and has a poor prognosis. Lung transplantation is the only curative treatment for PVOD.

Key words: pulmonary veno-occlusive disease, pulmonary hypertension, exertional dyspnea, sudden death

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Introduction

Pulmonary veno-occlusive disease (PVOD) presents with severe and progressive exertional hypoxemia and pulmonary hypertension (PH), resulting in a very poor prognosis.

In the 2015 European Society of Cardiology (ESC)/European Respiratory Society (ERS) guidelines, PVOD/pulmonary capillary hemangiomatosis (PCH) have been given the subgroup designation of group 1, because they differ greatly from the pulmonary arterial hypertension (PAH); the guidelines have proposed the further classification of PVOD/PCH into idiopathic, heritable, drug-, toxins-, and radiation-induced and associated forms (1). When the drugs used in the treatment of PAH are given to PVOD patients, they dilate the pulmonary arteries, increase the capillary hydrostatic pressure, and are likely to worsen pulmonary edema. Thus, lung transplantation is the only curative treatment for PVOD. We herein present the case of a patient with PVOD in whom transbronchial lung biopsy (TBLB) failed to establish a diagnosis. The patient’s PH progressed over a period of several months and did not improve with drug treatment. The diagnosis of PVOD was first made at autopsy.

Case Report

A 58-year-old woman visited her primary physician with a chief complaint of exertional dyspnea that began 2 months before hospitalization. Chest radiography showed a reticulated pattern in both lower lung fields and bilateral hilar lymphadenopathy. She was referred to our hospital and admitted for further evaluation.

She was a current smoker with 60-pack-year smoking history. The findings on admission were as follows: New York Heart Association (NYHA) Class III, body temperature, 36°C; blood pressure, 118/72 mmHg; pulse rate, 78/min; respiratory rate, 28/min; and SpO2, 88% (room air).

The laboratory tests on admission (Table 1) showed mild elevations in the patient’s lactate dehydrogenase (LDH), surfactant protein-D (SP-D), carcinoembryonic antigen (CEA), and rheumatoid factor (RF) levels. The patient was negative for all connective tissue disease (CTD) markers. An arterial blood gas analysis showed hypoxemia and respiratory alkalosis. Pulmonary function tests showed a mild restrictive disorder (vital capacity, 76.9%), whereas the diffusing capacity of the lung for carbon monoxide (DLCO), as determined by the steady state method, was reduced to 33.5% of the predicted value. An electrocardiogram showed peaked p
that the left ventricular systolic function was maintained, waves in leads II, aVF, and V1. Echocardiography showed that the left ventricular systolic function was maintained, while the right ventricular systolic pressure (RVSP) and tricuspid regurgitation pressure gradient (TRPG) were mildly elevated at 52.0 mmHg, 42.0 mmHg, respectively and the right ventricle was enlarged. We did not perform a 6-minute walk test. Chest radiography on admission (Fig. 1) showed a reticular pattern in both lower lung fields, bilateral hilar adenopathy, and right atrial enlargement. Chest computed tomography (CT) on admission (Fig. 2) showed bilateral hilar and mediastinal lymphadenopathy, ground glass opacities with lobular septal thickening, predominantly in both lower lung fields, and pulmonary artery enlargement.

Bronchoscopy was performed on hospital Day 5. TBLB of the right lower, middle, and upper lobes was performed. The histopathology was nonspecific, showing only alveolar septal fibrosis and small round cell infiltration, leading to a pathological diagnosis of chronic interstitial pneumonitis. The TBLB findings were again reviewed after the diagnosis was made. Although alveolar septal fibrosis and the fibrous obliteration of the small veins and venules were present (Fig. 3), these changes were slight and could not lead to the correct diagnosis. A thoracoscopic lung biopsy was contraindicated because of the high risk of morbidity and mortality. She was discharged from hospital on Day 17 with home oxygen therapy (1.5 L/min at rest and 3 L/min during exertion) to maintain SpO2 level of >90%. We continued the prescription of furosemide (20 mg/day) and enalapril (2.5 mg/day), which had been prescribed by her primary care doctor before admission.

However, the patient’s symptoms did not improve. At approximately 6 months after her discharge from hospital, her dyspnea worsened due to a cold and she revisited our hospital. Since her SpO2 was 80-85% with 2 L/min of oxygen by cannula, the patient was readmitted to our hospital.

### Table 1. Laboratory Data on the First Admission.

| Hematology | Serology | Arterial blood gas (room air) |
|------------|----------|------------------------------|
| WBC 8,300 μL | CRP 0.78 mg/dL | pH 7.442 |
| Neut 58 % | IgG 2.215 mg/dL | pCO2 29.6 mmHg |
| Lymp 37 % | IgA 407 mg/dL | pO2 48.8 mmHg |
| Eosi 3 % | IgM 91 mg/dL | HCO3^- 19.7 mmol/L |
| RBC 4.47×10⁶/μL | IgE 164 mg/dL | BE -3.1 mmol/L |
| Hb 14 g/dL | Ferritin 154 ng/mL |
| Plt 27.7×10⁹/μL | KL-6 371 U/mL |
| Biochemistry IL2R 410 U/mL |
| TP 7.9 g/dL | ACE 8.7 IU/mL |
| Alb 4 g/dL | βD glucan 11.8 pg/mL |
| LDH 255 IU/L | CEA 8.9 ng/mL |
| AST 26 IU/L | RF 35 IU/mL |
| ALT 33 IU/L | ANA <40 |
| γ-GTP 37 IU/L | Anti-RNP Ab (-) |
| ALP 235 IU/L | Anti-Scl70 Ab (-) |
| Cre 0.77 mg/dL | Anti-centromere Ab (-) |
| Na 142 mEq/L | Anti-SSA Ab (-) |
| K 4.1 mEq/L | Anti-SSB Ab (-) |
| Cl 105 mEq/L | PR3-ANCA (-) |
| MPO-ANCA (-) |

**Note:** LDH: Lactate Dehydrogenase, SP-D: surfactant protein D, IL2R: interleukin-2 receptor, ACE: angiotensin converting enzyme, βD glucan: (1→3)β-D-glucan, CEA: carcinoembryonic antigen, RF: rheumatoid factor, ANA: anti nuclear antibodies, Anti-RNP Ab: anti ribonucleoprotein antibody, Anti-Scl70 Ab: anti scleroderma antibody, Anti-centromere Ab: anti centromere antibody, Anti-SSA Ab: anti SS-A antibody, Anti-SSB Ab: anti SS-B antibody, PR3-ANCA: proteinase-3 anti neutrophil cytoplasmic antibody, MPO-ANCA: myeloperoxidase anti neutrophil cytoplasmic antibody.

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**Figure 1.** Chest radiography on initial admission. Reticular patterns were observed in both lower lung fields, along with bilateral hilar adenopathy, and right atrial enlargement.
Laboratory tests showed an elevated LDH level, severe hypoxemia and the deterioration of her respiratory alkalemia (Table 2). An electrocardiogram showed peaked p waves in leads II, aVF, and V1, a deep S wave in lead I, and a negative T wave in lead III, all of which were consistent with right heart overload. Echocardiography showed a decreased left ventricular systolic function with an ejection fraction (EF) of 50%. Her RVSP and TRPG values were further elevated to 72.0 mmHg and 60.0 mmHg, respectively, and right ventricular enlargement and ventricular septal flattening were observed. In addition to a reticular pattern in both lower lung fields and bilateral hilar lymphadenopathy, chest radiography also showed right atrial enlargement and a right pleural effusion (Fig. 4). Chest CT revealed an increase in both lower lobes and bilateral hilar lymphadenopathy, a small amount of right pleural effusion, a severely enlarged pulmonary artery, increased pulmonary vascular markings, and a reticular granular pattern, predominantly in the lower lung fields (Fig. 5). We could not perform pulmonary function tests or a 6-minute walk test.

Based on the marked hypoxemia, the imaging findings of a reticular pattern with lobular septal thickening, and the TBLB findings during her previous hospitalization, an acute exacerbation of interstitial pneumonitis was initially considered. Steroid pulse therapy (methylprednisolone [1 g for 3 days]) was started on admission. Her SpO2 was maintained at about 95% with an oxygen cannula (2 L/min), and there was a slight temporary improvement in her exertional dyspnea. However, on hospital Day 14, the patient’s dyspnea and oxygenation worsened, and 12 L/min of 50% oxygen by venturi mask was required to achieve an SpO2 of 95%. Thus, a second course of steroid pulse therapy was administered. In addition, her brain natriuretic peptide (BNP) was elevated to 827 pg/mL, pleural effusion developed, and echocardiography showed an increase in TRPG to 62.1 mmHg (Fig. 6a). Diuretics were added to treat the patient’s worsening right heart failure.

Her respiratory status improved with an SpO2 level of ≥ 90% on 6 L/min of oxygen by cannula. However, the patient developed severe lower leg edema, and echocardiography on hospital Day 42 showed a further increase in TRPG to 72.9 mmHg (Fig. 6b). For the PH with right heart failure, beraprost sodium (120 μg/day) was started on hospital Day 43. On hospital Day 48, however, her respiratory status worsened again and required 8 L/min of oxygen by reservoir mask. Chest radiography showed an increased reticular pattern in both lungs, again suggesting an acute exacerbation of interstitial pneumonitis. A third course of steroid pulse therapy was given, and then prednisolone (30 mg) was initiated as an after-treatment. Nevertheless, the respiratory insufficiency progressed, and the patient died suddenly on hospital Day 59 (Diagram: Fig. 7).

Consent was obtained from the patient’s family for an autopsy. The left lung weighed 485 g, and the right lung weighed 445 g. A histopathological examination showed hyaline fibrosis of the alveolar septa and fibrous obliteration and concentric intimal thickening of the venules (Fig. 8a and b). Heath-Edwards grade II (2) medial hypertrophy of the pulmonary artery was present, but plexiform lesions were not observed (Fig. 8c). There were also dilated, multi-layered capillaries, hemorrhages in the alveolar spaces and alveolar septum, and areas of hemosiderosis (Fig. 8d). The mediastinal lymph nodes were enlarged to 40 mm.
PVOD was first reported by Hora et al. (3) in 1934, and it was recognized as a separate disease entity by Heath et al. (4) in 1966. The exact prevalence of PVOD is unknown, but it is estimated to be 0.1-0.2 per one million population per year (5). Pathologically, the venules and small septal veins in PVOD are most often involved with the arterial and capillary components of the pulmonary vasculature, whereas plexiform lesions, a characteristic of PAH, are not found. In this case, the pathological features were compatible with PVOD. In terms of the clinical course, PVOD typically demonstrates a poor response to the vasodilator drugs used in PAH, resulting in severe pulmonary edema (6).

The etiology of PVOD is not completely understood, but associations with genetic factors, anticancer drugs such as mitomycin and bleomycin, bone marrow transplantation, radiation therapy, smoking, CTDs such as systemic lupus erythematosus (SLE) and rheumatoid arthritis (RA), and chronic hepatitis have been reported (6). The French PH network reported the genealogy of 13 PVOD families with heritable disease and demonstrated that biallelic mutations were present in eukaryotic translation initiation factor 2 alpha kinase 4 (EIF2AK4) (7). The French PH networks stated that these mutations were also identified in 9% of apparently sporadic cases of PVOD, when they offered genetic counselling and EIF2AK4 mutation screening to all patients with or without a family history (8). The present patient had no family history of PVOD, but these mutations might have been associated. In this case, the screening tests for CTD were all negative, and although smoking has been men-
Figure 5. a: Chest CT on readmission (lung fields). An increase in the reticular granular pattern was observed, predominantly in both lower lobes, along with increased pulmonary vascular markings. b: Chest CT on readmission (mediastinum). Further enlargement of the mediastinal lymph nodes was observed.

Figure 6. a: An echocardiographic image on day 19 of the second hospitalization. b: An echocardiographic image on day 41 of the second hospitalization.

The clinical course after readmission in a diagram

|                   | Day0 | Day14 | Day22 | Day47 | Day59 |
|-------------------|------|-------|-------|-------|-------|
| Diuretics         |      |       |       |       |       |
| Steroid           |      |       |       |       |       |
| Vasodilator drug  |      |       |       |       |       |
| Amount of oxygenation |     |       |       |       |       |
| KL-6 (U/mL)       | 380  |       |       |       |       |
| BNP (pg/mL)       | 827  | 1,110 |       |       |       |
| TRPG (mmHg)       | 62.1 | 72.9  |       |       |       |

Figure 7. A diagram of the clinical course of the present case.
reviewed the TBLB findings again, we found that we had
missed slight pathological findings. These results indicate
the difficulty of making an antemortem diagnosis of PVOD.

The present patient had various imaging findings, includ-
ing ground glass opacities with lobular septal thickening,
mediastinal lymphadenopathy, and centrilobular granular
shadows, and these findings initially suggested sarcoidosis
or hypersensitivity pneumonitis. Resten et al. (10) compared
the chest CT findings in PVOD with those in idiopathic
PAH. They reported that ground glass opacities, lobular septal
thickening, and mediastinal lymphadenopathy were signifi-
cantly more frequent in PVOD, and that the ground glass
opacities more often had a centrilobular pattern than a pan-
lobular pattern. These findings reflect interlobular septal
edema followed by fibrosis and the lymphatic dilation asso-
ciated with pulmonary venous occlusion (11). The imaging
findings in the present patient was a characteristic of PVOD.

The prognosis of PVOD is very poor, with a one-year
mortality rate of 72% (12). Most patients die within 2 years
after being diagnosed, and the prognosis is even poor in
comparison to PAH (13). Lung transplantation is the only
curative treatment (14), but the long average waiting time in
Japan (889.6 days) (15), means that most patients with
PVOD progress before they have the opportunity to undergo
transplantation. With regard to drug treatment, when the

tioned as a risk factor, there is no firm evidence.

An evaluation by right heart catheterization (RHC) is es-
sential for the diagnosis of PH. However, in this case, be-
cause our institution had not yet completely arranged an
RHC system, we could not perform a hemodynamic assess-
ment.

A definitive diagnosis of PVOD requires a surgical lung
biopsy, but because of the invasive risk of this procedure,
few patients, including the present patient, are diagnosed
with PVOD antemortem (6). Rabiller et al. (9) found that
hemosiderin-laden macrophages were significantly increased
in the bronchoalveolar lavage (BAL) fluid of PVOD pa-
tients, which meant that PVOD is associated with occult al-
veolar hemorrhage due to the presence of post-capillary
block. The 2015/ESC/ERS Guidelines of PH (1) state that
BAL may be a useful diagnostic strategy for PVOD unless
severe hypoxemia is present, while TBLB is no longer rec-
ommended in patients with severe PH due to the high risk
of bleeding. However, in this case, we decided to give prior-
ity to TBLB. Because the initial diagnosis at the first hospi-
talization was interstitial pneumonia, we considered the se-
verity of her PH to be mild, and thought that her respiratory
status gradually deteriorated during bronchoscopy. When we
reviewed the TBLB findings again, we found that we had

Figure 8. a: The histopathology at autopsy (EVG staining ×40). Hyaline fibrosis of the alveolar
septum and fibrous obliteration of the small veins and venules were observed. b: The histopathology
at autopsy (EVG staining ×100). Fibrous obliteration and narrowing due to concentric intimal thick-
ening of the small veins and venules were observed. c: Histopathology at autopsy (EVG staining
×100). The histopathological examination revealed Heath-Edwards grade II and medial hypertrophy
of the pulmonary artery without plexiform lesions. d: The histopathology at autopsy [Hematoxylin
and Eosin (H&E) staining ×40]. Dilated, multilayered capillaries, hemorrhages in the alveolar spaces
and alveolar septum, and areas of hemosiderosis were observed. e: The histopathology at autopsy
(H&E staining ×40). Enlarged mediastinal lymph nodes with enlarged lymph sinuses and histiocytic
hemophagocytosis were observed.
vasodilator drugs that are used to treat idiopathic PAH are given to patients with PVOD, who have high pulmonary venous resistance, the pulmonary artery dilates, the capillary hydrostatic pressure rises, and pulmonary edema is likely to occur. Thus, the use of these drugs in PVOD is controversial (14).

Increasing the dose of diuretics could not control the rapidly progressing right heart failure in the present patient. Ultimately, beraprost sodium, a prostaglandin I2 (PGI2) analog, was started. However, within a few days after starting the drug, the interstitial opacities increased and the patient’s respiratory status worsened. Thus, the PGI2 drug may have actually worsened the patient’s pulmonary edema.

Ogawa et al. (16) reported the results of initiating treatment with low-dose epoprostenol in PVOD patients who were already using diuretics or vasopressors. With careful management to achieve venodilatation and reduce the hydrostatic pressure, it could eventually be increased to a mean maximum dose of 55.3±10.7 ng/kg/min. The NYHA class, 6-minute walking distance, and BNP levels improved. However, even though epoprostenol improved the exercise tolerance and increased the cardiac output, the pulmonary artery pressure and right atrial pressure were not reduced. Thus, the effectiveness is temporary and may be limited to short-term treatment as a bridge until lung transplantation.

The present patient was not eligible for lung transplantation because she was over 55 years of age. However, we think we should have started beraprost sodium treatment at a low dose of 60 μg/day, or chosen epoprostenol and slowly increased the dose because we would have been able to easily adjust the amount as it is administered by continuous infusion.

The 2015/ESC/ERS Guidelines of PH (1), note that a highly probable diagnosis of PVOD can be made based on the combination of clinical suspicion, a physical examination, bronchoscopy and radiological findings. In the present case, the initial diagnosis was interstitial pneumonitis such as sarcoidosis, hypersensitivity pneumonia, or idiopathic interstitial pneumonia because of the radiological findings and TBLB findings during the prior hospitalization. However, several points were not compatible; her clinical course was very aggressive, granulomas were not detected by TBLB, there was no uptake on Gallium scintigraphy, the patient was negative for Trichosporon asahii antibodies, and there was no history of exposure. On the other hand, the severe desaturation on exercise, the very low DLCO level, the characteristic radiological findings and the rapid progression of right heart failure were consistent with PVOD. When looking back, we should have suspected PVOD before the patient’s second admission and have at least made an appropriate diagnostic approach to progressive right heart failure, especially RHC. It is very important to perform RHC to confirm PH. For an early diagnosis, PVOD should always be considered in patients with these clinical features and imaging findings. PVOD is such a rare case and the clinical course of it is so progressive that it is very important for physicians to suspect it in the early phase by a noninvasive approach and to immediately refer patients who are eligible for lung transplantation.

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

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