INTRODUCTION

Tuberous sclerosis complex (TSC) is a hereditary neurocutaneous syndrome with a classic triad of seizures, mental retardation, and adenoma sebaceum. It is inherited by an autosomal dominant pattern and is caused by a mutation in the TSC1 and TSC2 genes encoding hamartin and tuberin respectively, both of which are thought to act as tumor suppressor genes (1). Lack of functional hamartin or tuberin may explain the development of hamartomas and various tumors in TSC. Almost all systems or organs of the body can be involved in TSC. Clinical manifestations are unique in many cases but in some cases, characteristic findings from imaging are the first step to raise a possibility of TSC.

Hamartomas in TSC patients are most commonly found in the skin, kidneys, brain, and heart. Less frequently, they can involve the retina, gingiva, bones, gastrointestinal tract, and lungs. Lymphangioleiomyomatosis (LAM) is a well known and major feature of the pulmonary manifestation of TSC. Pulmonary manifestations are known to occur in approximately 1–2.3% of TSC patients, but the incidence turned out to be higher based on radiologic findings in up to 26–39% of female TSC patients (2). More recently, multifocal micronodular pneumocyte hyperplasia (MMPH) has been reported to occur in association with TSC (3, 4). MMPH is a hamartomatous process of the lung with multiple small nodules which are composed of type II pneumocytes. Plain radiography and chest CT in MMPH may demonstrate numerous small nodules measuring 1–10 mm in diameters, distributed randomly throughout both lungs. If MMPH is an initial presentation of TSC, and unless we are familiar with this lung manifestation of TSC, radiologic findings can mimic miliary tuberculosis or metastatic disease. We report a teenage girl with TSC and histologically confirmed MMPH which mimicked miliary tuberculosis at the initial presentation.
CASE REPORT

A 15-year-old girl was referred to our institute because of abnormal findings on chest radiographs. An initial chest radiograph for student screening a year ago showed numerous fine nodular opacities, measuring a few millimeters, and evenly scattered throughout both lungs (Fig. 1A). With an initial diagnosis of miliary tuberculosis, anti-tuberculosis medications were given at the local public health center. However, standard anti-tuberculosis treatment for more than 9 months failed to show radiological improvement. Subsequently, she was referred to our hospital for further evaluation.

On admission, the patient had no respiratory symptoms. Physical examination revealed wheezing over both lung fields and skin lesions including hypopigmented macules and shagreen patches. She was mentally retarded (Full Scale Intelligence Quotient 55). The laboratory test did not show any abnormal results at the time of her first visit. She had a history of temporary seizure-like events during infancy. About ten months ago, she had multiple events of decreased responsiveness and automatic hand movements. Video-electroencephalography showed a mild cerebral dysfunction but interictal epileptiform discharge was not observed. Antiepileptic medication was not given. She had no family history of tuberous sclerosis.

The high-resolution CT (HRCT) showed miliary nodules and multiple small nodules, varying in diameter from 3 to 12 mm, and randomly distributed throughout both lung fields (Fig. 1B). No cysts, pleural effusions, or significant mediastinal or hilar lymphadenopathy were noted. Bronchoscopy revealed no endobronchial lesions. Brain MRI demonstrated multiple subependymal nodules along the ventricular margins, and tubers in the cerebral cortex and subcortical white matter of both hemispheres (Fig. 2). Subsequent ultrasonography and CT of the abdomen showed small cysts and presumed angiomyolipomas in the both kidneys.

A biopsy was performed on the right upper and middle lobes by means of video-assisted thoracoscopic surgery. Histologically, the parenchymal tissue showed multifocal areas of increased cell density, and consisted of increased septal fibrosis and papillary growths of hyperplastic type II pneumocytes (Fig. 3A). No caseating necrosis or granulomatous inflammation was identifiable. Immunohistochemically, most of the proliferating epithelial cells of the lesion were positive for anti-pan-cytokeratin (Fig. 3B) and anti-thyroid transcription factor-1. However, they were negative for anti-melanocyte marker (HMB-45).

DISCUSSION

TSC is an autosomal dominant disorder characterized by the triad of seizures, mental retardation, and multiple hamartoma lesions. Our patient could be diagnosed as having definite TSC since she had all the features of the classic triad and two major fea-
tures (subependymal nodules and renal angiomyolipoma) of the clinical diagnostic criteria. In our patient, there was no family history of TSC, supporting this case was sporadic, although gene study was not performed.

LAM and MMPH are two major lung manifestations of TSC. LAM is a rare lung disease characterized by diffuse proliferation of abnormal smooth muscle-like cells and cystic destruction of the lung. It is more common in young women who present with dyspnea and pneumothorax (3).

MMPH is another lung manifestation of TSC. The precise prevalence of MMPH in patients with TSC is not known, but may be as high as 40–58% (6). There is no gender restriction, and MMPH may occur in association with LAM in TSC patients (3). In our case, there was no associated feature of LAM on CT.

![Fig. 2](image1.png)

**Fig. 2.** Brain MR imaging of multifocal micronodular pneumocyte hyperplasia with tuberous sclerosis complex in a 15-year old girl.  
A. Axial FLAIR image shows a subependymal nodule near the right foramen Monro and multiple high signal intensities at the cerebral cortex and subcortical white matter suggesting cortical and subcortical tubers.  
B. Axial gradient echo image shows calcified subependymal nodules along the lateral ventricular margins.  
FLAIR = fluid attenuated inversion recovery, MR = magnetic resonance

![Fig. 3](image2.png)

**Fig. 3.** Microscopic findings of VATS biopsy specimens of multifocal micronodular pneumocyte hyperplasia with tuberous sclerosis complex in a 15-year old girl.  
A. Close-up view of the nodule show increased septal thickness and pneumocyte hyperplasia (arrows) (hematoxylin and eosin stain, × 100).  
B. Immunohistochemically, most of the proliferating epithelial cells of the lesion are positive for pan-cytokeratin (brown color) (pan-cytokeratin stain, × 40).  
VATS = video-assisted thoracoscopic surgery
MMPH, multiple pulmonary nodules composed of benign alveolar type II cells are found throughout the lung. These lesions stain with cytokeratin and surfactant proteins A and B, but not with HMB-45, or hormonal receptors which show strong positivity in LAM (4).

Chest CT in patients with MMPH demonstrates small nodules measuring 1–10 mm in diameter, which are randomly distributed throughout the lung with regard to the secondary lobules (4). The differential diagnosis should include a miliary granulomatous infection, hematogenous metastases, Langerhans’ cell histiocytosis (LCH), or atypical adenomatous hyperplasia (AAH). Miliary tuberculosis shows a very fine nodular or reticulonodular pattern with an even distribution on HRCT. A review of HRCT findings in 25 patients who had proven miliary tuberculosis, demonstrated miliary nodules measuring 1 to 3 mm in diameter (7). Additional radiological findings, such as intra- and interlobular septal thickening, intrathoracic lymphadenopathy or pleural effusion on HRCT favors miliary tuberculosis. Hematogenous metastases typically appear as small discrete nodules that have a peripheral and basal predominance on CT when limited in number. Metastatic lung disease is much less common in children than in the adult population. In addition, hematogenous metastases and MMHP are quite different in clinical features. In LCH, there are nodules as well as cysts that predominantly involve upper lung zones with relative sparing of the costophrenic angles (8). Relative sparing of the lung bases is a useful feature of LCH to differentiate it from other lung diseases (8). In MMHP, the nodules are randomly distributed and the costophrenic angles can be involved (3). Kushihashi et al. (9) described CT findings of AAH that demonstrated small nodules with ground-glass opacity. Ground-glass opacity in MMHP is correlated with decreased alveolar air spaces due to hyperplastic proliferation of type II pneumocytes and infiltration of macrophages into the alveoli (10). However, multicentric lesions seen in MMHP are not common in AAH.

Clinically, patients with isolated MMHP usually have no respiratory symptoms as seen in our case (10). Some patients with MMHP may present with dyspnea, cough, and mild to moderate hypoxemia. Unlike pulmonary LAM, treatment is usually not necessary because MMHP does not appear to be fatal or progressive. MMHP has not shown any malignant potential although there is a potential loss of tumor suppressor function in TSC gene mutations (10).

The incidence of MMHP seems to be underestimated in children or adolescents because there is usually no respiratory symptom requiring imaging studies, and chest CT is less frequently performed in pediatric patients than in adult patients. However, it is not practical to perform chest CT for screening in all TSC patients.

It is difficult to differentiate MMHP or benign hamartomatous lung disease from other diseases requiring further evaluation or treatment on the basis of radiographic findings only. A definitive diagnosis of MMHP can only be made by a lung biopsy. Most MMHP occurrences are in patients with TSC, and its clinical course is usually benign. Therefore it is important to find the stigmata of TSC and do clinical and radiological follow-ups before undergoing histological confirmation in MMHP patients.

In summary, we report a teenage girl with MMHP and TSC which mimicked miliary tuberculosis at the initial presentation. MMHP is a benign hamartomatous process of the lung with multiple small nodules which are composed of type II pneumocytes. When randomly distributed tiny nodules are observed on chest radiographs or chest CT in patients suspected of TSC, MMHP, a recently recognized entity in TSC, should be considered.

REFERENCES

결절성경화증 환아에서 발생된 다발성 미세결절 폐세포증식증: 증례 보고

하승미 1 · 윤혜경 1* · 이승구 2

다발성 미세결절 폐세포증식증은 결절성경화증에 동반될 수 있는 폐질환의 하나로서 비교적 드문 것으로 알려져 있으며, 소아 결절성경화증 환자에서 발생한 다발성 미세결절 폐세포증식증에 대한 보고는 더욱 드물다. 다발성 미세결절 폐세포 증식증은 본질적으로 과오종 결절으로서, 제2형 폐세포로 구성된 다발성 작은 결절을 형성하는 것이 특징이다. 단순흉부 촬영과 전산화단층촬영에서 폐야 전체에 걸쳐 분포하는 1~10 mm 크기의 다발성 폐결절을 형성하는 것이 특징이다. 이러한 영상의 학적 소견이 결절성경화증 진단 전에 보이는 경우 속립성 폐결핵 및 전이성 폐암과의 감별을 어렵게 할 수 있다. 이에 우리는 본 증례를 의심하여 치료를 받았던 소아에서 결절성경화증과 미세결절 폐세포증식증으로 확진되었던 증례를 보고 하고자 한다.

강원대학교병원 1영상의학과, 2병리과

8. Webb WR, Muller NL, Naidich DP. High-resolution CT of the lung. Philadelphia: Lippincott Williams & Wilkins, 2014: 368-381