A Rare Case of Chylothorax in a Patient with Schimmelpenning Syndrome

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Abstract
Chylothorax originating in a patient with Schimmelpenning syndrome is rare and poses a problem in diagnosis and treatment. A 22-year-old male was admitted with dyspnea indicative of a large pleural chylous effusion. Besides conservative dietary treatment measures, the chylous effusion was drained (2,000 mL/day). Computed tomography-lymphography after ligation of the thoracic duct and pleurectomy revealed a small collateral flow of chylous fluid toward the chest wall and entering the thorax. Eventually, local radiation therapy with 36 Gy effectively treated the chylothorax. Five months later, an epitheloid angiosarcoma developing from a preexisting cutaneous lesion was detected and treated by surgical resection.

Keywords
► lung
► rare disorders
► pleural disease
► thoracic duct

Introduction
Chylous leaks, such as chylothorax, are uncommon effusions in nontraumatic patients, resulting from leakage of the thoracic duct and its tributaries. Schimmelpenning syndrome is a rare neurocutaneous disorder characterized by one or more sebaceous nevi of the face or scalp, in association with anomalies of the central nervous system, ocular or skeletal pathology. This article describes vascular malformations of Schimmelpenning syndrome and also lymphatic anomalies in very rare cases.

Case Report
A 22-year-old male student, who was formerly diagnosed with Schimmelpenning syndrome, presented to the emergency room with shortness of breath and a feeling of pressure in the chest. Symptoms were aggravated 1 week prior to his admission. He had a 10-year history of right-sided leg lymphedema (►Fig. 1) and his right eye had been enucleated because of a teratoma at the age of 1. The patient’s skin showed multiple eflorescences suggestive of sebaceous nevi. Chest X-ray showed a right-sided pleural effusion (►Fig. 2). The patient was initially treated with a chest tube, which drained 2,000 mL chylous effusion and persisted despite parenteral nutrition. Computed tomography (CT)-lymphography revealed a leakage of the thoracic duct at the mid-thoracic level. The patient underwent thoracotomy with ligation of the thoracic duct and pleurectomy revealed a small collateral flow of chylous fluid toward the chest wall and entering the thorax. Eventually, local radiation therapy with 36 Gy effectively treated the chylothorax. Five months later, an epitheloid angiosarcoma developing from a preexisting cutaneous lesion was detected and treated by surgical resection.

During the course, the patient did well for 4 months but was then admitted to the intensive care department with severe exsiccosis, and dyspnea from chylous thoracic effusion and...
chylous ascites, and cachexia. Despite all supportive efforts, the patient succumbed due to renal insufficiency and septicemia.

**Comment**

Schimmelpenning syndrome is considered a rare disease, and is listed in the US-registry of the National Organization for Rare Disorders. It has been proposed that Schimmelpenning syndrome results from genetic mosaicism, with underlying genetic mutations of the HRAS gene and KRAS gene.\(^3\)

The term Schimmelpenning syndrome has been widely used since 1962 and describes the classic triad of sebaceous nevi, seizures, and mental retardation.\(^4,5\) However, further studies revealed additional anomalies in the affected patients, and thus the diagnostic definition by the three symptoms was relativized.\(^6\)

Also, vascular anomalies, such as hemangiomas or nevus flammeus, have been described, with the term Schimmelpenning syndrome being used nosologically imprecise.\(^7\)

Lymphatic malformations are equally rarely seen and documented within a review of the literature in 3 of 119 patients. To our knowledge, only Greene et al\(^2\) reported two patients with chylothorax in a series of nine patients with Schimmelpenning syndrome.

The basic principles of chylothorax treatment include efficient drainage, complete lung expansion by chest tube drainage, and reduction of the chylous flow by dietary and medical treatment. Surgical options by video-assisted thoracic surgery or thoracotomy include ligature of the thoracic duct and pleurodesis by intrapleural irritants. However, in postoperative chylothorax up to 11% of the patients underwent repeat surgical procedures due to complications, and 9 to 15% experienced recurrences.\(^8,9\)

In recent years, percutaneous embolization\(^10\) and radiotherapy\(^11\) have been suggested as further treatment options to avoid invasive procedures and as an alternative in recurrent chylothoraces to obliterate the thoracic duct and adjacent lymphatic vessels.
In previous years, surgical excision of the sebaceous nevus was considered the treatment of choice to prevent malignant degeneration; at present, such prophylactic measures are not considered necessary, as the estimated rate of malignant transformation seems to be low.\(^1\) In our case, an angiosarcoma developed out of a preexisting cutaneous lesion. This underlines the importance of close clinical observation of the cutaneous lesions in Schimmelpenning syndrome.

In conclusion, ectatic dysplastic lymphovenous malformations with lymphatic collateral flow in Schimmelpenning syndrome made ligation of the thoracic duct for chylothorax difficult. Alternative measures for lymphatic obliteration, such as local radiation therapy, are feasible. Close observation of concomitant symptoms is warranted.

References
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Fig. 3  Computed tomography-lymphography showing the thoracic duct ligated and a leakage at the midthoracic level.

Fig. 4  Angiosarcoma developed at the lower right leg.