Eccrine spiradenoma is exceedingly rare, accounting for roughly 1 of 13,000 specimens submitted to a dermatopathologist, especially at hand\(^1\). It most commonly arises in people aged 15 to 35 years, and the front of the trunk and proximal limbs are the most common sites.

Proper diagnosis of eccrine spiradenoma is important because of the occurrence of potentially life-threatening malignant transformation. From literature review, there is few case of spiradenoma on hand\(^2\). Author present a case of solitary spiradenoma on subcutaneous layer mimicking neurilemmoma.

**CASE REPORT**

A 36-year-old Asian woman with no significant medical past history presented with history of tingling and numbness of little finger in the ulnar nerve sensory territory of the right hand without tenderness, since the past ten years. Physical examination of the right hand showed about 15 mm size oval shape well-defined soft tissue...
mass situated on subcutaneous fat. There is positive Tinel’s sign at her ulnar side of hand. A neurilemmoma was suspected on the basis of findings at clinical examination. Under local anesthesia, excisional biopsy was performed (Fig. 1). The incision line was designed along the long axis of mass for easy approach. Grossly, Specimen was 15 mm size pinkish color oval shape mass with firm encapsulated characteristics (Fig. 2). Histopathologic examination demonstrated well-demarcated tumor with capsulation and cystic change. The tumor cells consisted of small basaloïd cells, paler cells, and lymphocytes, arranged with a ribbon and reticular pattern (Fig. 3). The histopathologic findings of the skin were consistent with spiradenoma. The patient recovered uneventfully from the surgery. There is no evidence of recurrence and metastasis.

**DISCUSSION**

Eccrine sweat glands are simple tubular glands that open directly to the skin and are present in all body areas, with their highest concentration in the palms, soles, and axillae. The etiology of eccrine spiradenoma is not fully understood. It is often misdiagnosed with neuromas, leiomyomas, neurofibroma, hidradenocarcinoma of the sweat glands, glomus tumors, lipoma, angiolipoma, dermatofibroma, hemangioma, and angioleiomyoma. Computed tomography and magnetic resonance imaging can help the diagnosis of this kind of tumor, especially for tumors in the deep dermis or subcutaneous tissues. However, the definitive diagnosis requires skin biopsy. Low incidence of this tumor may lead to a delay in diagnosis and treatment.

Clinically, it presents as numerous nodules and also can present as a painful, slow-growing, solitary nodule that approximately one third of eccrine spiradenoma occur in the head and neck, one third on the trunk, and one fifth on the extremities. Random multiple tumors have been reported to occur on the chest, upper extremities, forehead, and scalp. Noto et al. reported linear spiradenoma arising in medial canthus and cheek. Gupta et al. reported a 23-year-old woman with linear facial spiradenoma and coexistent eyelid spiradenoma. Pain or tenderness are not always suggested as a clinical characteristic. Several study showed non tender eccrine spiradenoma. The generation of pain is thought to be related to small unmyo-
clinated axons permeating the hyalinized stromal mantle. Some researchers believe that the expansion of cysts in the tumor also gives rise to pain. Although the pain of an eccrine spiradenoma is ‘the most frequent and striking symptom’, the mechanism of pain generation has not been elucidated entirely. In 1996, Criton and Aravindan stated that pain might be due to the contraction of myoepithelial cells. However, electron microscopy has not proven the presence of myoepithelial cells in the tumor.

Histopathologically, eccrine spiradenoma presents as intradermal lobules surrounded by a fibrous capsule without connections to the epidermis. On higher magnification view, the epithelial cells within the tumor lobule are arranged in intertwining cords. The epithelial cells within the tumor lobule are arranged in intertwining cords with two types of epithelial cells. There have been some reports of malignant transformation of spiradenomas.

Complete surgical excision with longitudinal incision is the first choice of treatment for tumor clearance. The lesion does not usually recur after excision. Surgery is unnecessary if the lesions are not disfiguring, increasing in size and number, or painful. There are no clear recommendations for margins for standard excision. Radiation, chemotherapy, and hyperthermic limb perfusion chemotherapy have been carried out in malignant spiradenomas.

In conclusion, author misdiagnosed as neurilemmoma due to its neurologic symptom rather than typical spiradenoma’s characteristics like pain symptoms. The use of transverse incisions parallel to the long axis of the mass other than longitudinal incision should be avoided. Failure to identify its morphologic features may lead to a mistaken diagnosis. Considering the anatomic location and clinical history, differential diagnoses and preoperative magnetic resonance imaging (MRI) should be performed. Because of possibility of malignant change, close local surveillance (MRI plus contrast) is essential.

CONFLICTS OF INTEREST

The authors have nothing to disclose.

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신경초종으로 오진된 손의 땀샘종

변제언 · 김준혁
순천향대학교 의과대학 천안병원 성형외과학교실

 المدني종은 양성 종양으로서 땀샘에서 기인한다. 36세의 특이 내과적 과거력 없는 동양인 여성이 새끼손가락의 무감각 증상을 호소하였다. 티넬 징후가 손의 칙골 부위에서 확인되었다. 피하지방에 단발성의 단단하고 둥근 형태의 종괴가 축지되었다. 임상적으로 신경초종을 의심하여 국소 마취하에 절제 생검술을 시행하였다. 병리학적으로 땀샘종의 병리소견을 나타내었다. 이번 증례에서 저자는 땀샘종을 신경초종으로 오진하였는데, 이는 통상적으로 통증을 일으키는 것과 다르게 신경학적 증상을 유발하며 신경초종과 비슷한 형태를 보였기 때문이다. 때때로 평활근종, 섬유종, 지방종, 이물 육아종, 토리종양, 혈관종, 신경섬유종, 방추세포육종, 결절성 흑생종도 위치에 따라 유사한 증상을 나타낼 수 있다. 따라서 처음 진단 과정에서 이들을 감별 진단하는 것이 중요하며, 수술 전 충분한 검사가 중요하다. 땀샘종의 정확한 진단이 중요한 이유는 때때로 생명을 위협하는 악성 종양으로 진행할 수 있기 때문이다.

색인단어: 손, 생증, 신경초종, 티넬 징후

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교신저자 김준혁
31151, 충남 천안시 동남구 순천향6길 31, 순천향대학교 의과대학 천안병원 성형외과학교실
TEL 041-570-2195  FAX 041-574-6133  E-mail psdoctor@schmc.ac.kr