A case report of lymphangioma circumscriptum of the vulva

S Murugan, G Srinivasan, M C A Kaleelullah, Lewellyn Rajkumar

Abstract
Lymphangioma circumscriptum of the vulva mimicking condyloma acuminata is reported for its rarity.

Introduction
Lymphangioma circumscriptum is a localised group of thin walled and translucent vesicles resembling frog spawn.1 Axilla, adjacent chest-wall, oral cavity and tongue2 are common sites. It is a form of nevus and recurrence even after surgery is common. The vulva is a rare site for lymphangioma circumscriptum and only four such cases have been reported in the literature.

Case report
An unmarried, 20 year old, female was referred to the department of sexually transmitted diseases (STD) of this hospital in December 1990 as she had a progressive swelling of the genititals for 2 years. She had no sexual contact so far. She had undergone some irregular allopathic and native treatment for the present ailment. On examination, both labia majora and minora were swollen, hypertrophied, and hyperpigmented over which numerous greyish white vesicles were present. Swelling extended up to the right groin and perineal region. Clear fluid exudated from a punctured vesicle. Profuse white purulent discharge per vagina was present. The cervix was healthy. Investigation by wet film for Trichomonas vaginalis and Gram stain for gonococci were negative, but candidal filaments were seen in KOH preparation. Microfilaria were absent in a night peripheral smear. The Mantoux test was negative. Total and differential WBC count were within normal limits. Erythrocyte sedimentation rate was 15 mm/h (Westegren). There was no evidence of pulmonary tuberculosis on a chest radiograph. VDRL slide test was non reactive.

A biopsy specimen from the lesion was stained with H and E and Elastica von Gissen and revealed hyperplasia of the epidermis with pigmentation. The dermis showed numerous dilated lymphatic and angiomatic spaces with proliferation of capillaries and an extensive lymphocytic and plasma cell infiltration. The diagnosis of lymphangioma circumscriptum was confirmed.

She was referred for plastic surgery. Vulvectomy was performed. She returned with a recurrence after 9 months. Once again the lesion was excised.

Discussion
Lymphangioma circumscriptum is a rare form of nevus, usually occurring at an early age but it may become clinically apparent up to 35 years and even later.3
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