Poromatosi s in Pregnancy: A Case of 8 Eruptive Poromas in the Third Trimester

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The poroid family of neoplasms includes hidroacanthoma simplex, eccrine poroma, dermal duct tumor, and poroid hidradenoma. These benign adnexal neoplasms are derived from the eccrine or apocrine sweat ducts or glands. Poroid neoplasms, including poromas, have been reported during pregnancy and have been hypothesized to be hormonally influenced. Poromatosis, the occurrence of multiple poromas, rarely has been reported in association with hidrotic ectodermal dysplasia, prior radiation therapy, and non-Hodgkin lymphoma occurring after chemotherapy. We report a case of eruptive poromatosis in pregnancy with 8 poromas occurring in the third trimester, further supporting the hypothesis of a hormonal association in the etiology of this neoplasm.

Case Report
A 33-year-old woman presented at 35 weeks’ gestation with painful red papules on the arms, hands, and feet of 2 weeks’ duration. These papules all appeared during the third trimester of her pregnancy. Her medical history was notable for melanoma of the leg and Hodgkin lymphoma treated with chemotherapy and an autologous stem cell transplant 15 years prior. Physical examination revealed 8 firm, flesh-colored to red papules measuring 1 to 4 mm: 1 on her left forearm, 2 on her right palm, 1 on her right lateral foot, 1 on her right fifth toe, 1 on her left volar forearm, and 2 on her left plantar foot (Figure, A and B).

Excisional punch biopsies from 2 papules demonstrated well-demarcated neoplasms in the superficial dermis comprised of monomorphic basaloid cells with multiple epidermal connections exhibiting duct formation consistent with the diagnosis of eccrine poromas (Figure, C and D). Scattered mitotic figures were present, but no significant cytologic atypia was noted. The 6 remaining papules were successfully treated with cryotherapy. There was no recurrence at 6 months postpartum.

Comment
Poromas are benign adnexal neoplasms derived from the epithelium of sweat ducts of either eccrine or apocrine origin. There are 4 recognized types of poroid neoplasms: hidroacanthoma simplex, eccrine poroma, dermal duct tumor, and poroid hidradenoma. A malignant poroid neoplasm with metastatic capability also exists. Eccrine poroma originally was described by Pinkus et al and is characterized by collections of monomorphic, cuboidal, basophilic cells (poroid cells) in the superficial dermis with a variable degree of duct formation and broad epidermal connections. Dermal duct tumor and poroid hidradenoma are both nodular dermal neoplasms without epidermal connection, the former consisting of multiple nodules of poroid cells and the latter as a nodule of poroid cells with solid and cystic components. Hidroacanthoma simplex is differentiated from others by its intraepidermal location.

The derivation of poromas (eccrine vs apocrine) is somewhat controversial. For decades, poromas were thought to derive from the acrosyringium (intraepidermal portion of the eccrine sweat duct). However, as more cases have emerged with elements of sebaceous or follicular differentiation, the possibility that poromas also arise from apocrine ducts has gained...
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Poromas typically present as solitary papules, plaques, or nodules, often on acral skin where eccrine ducts are abundant, but they also have been reported on the head, face, trunk, and anogenital regions. Poromas can be flesh colored or red due to a highly vascularized stroma that gives them an appearance similar to a pyogenic granuloma. They occasionally present as pigmented lesions. Although most poromas occur in adults with no sexual or racial predilection, they also can occur in children.

Solitary poromas have been reported in association with radiation therapy, nevus sebaceous, trauma/burns, and basal cell nevus syndrome. Poromatosis, the occurrence of multiple poromas, rarely has been reported in patients with hidrotic ectodermal dysplasia, prior radiation therapy, and non-Hodgkin lymphoma occurring after chemotherapy. A linear variant also has been described.

Solitary poromas also have been reported in pregnancy as well as malignant eccrine poroma. Ban and Kitajima and Guimerá Martin-Neda et al discussed the possibility of hormonal changes during pregnancy, which predisposes patients to the development of poromas; however, according to a PubMed search of articles indexed for MEDLINE using the terms poroma and pregnancy as well as poromatosis, no cases of poromatosis in pregnancy have been reported.

Conclusion
We report a case of eruptive poromatosis in pregnancy with 8 poromas occurring in the third trimester,
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lending further support to a potential hormonal contribution in the pathogenesis of poromas.

REFERENCES


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