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Acrochordon-like basal cell carcinomas in patients with basal cell nevus syndrome

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Abstract

Basal cell nevus syndrome is an autosomal dominant disorder characterized by multiple basal cell carcinomas, along with numerous other documented clinical features. Acrochordons (or skin tags) are common benign neoplasms that are appropriately left untreated in most patients. We describe two patients with known BCNS who were found to have multiple BCCs that clinically resembled acrochordons. Our findings support the biopsy of acrochordon-like growths in patients with basal cell nevus syndrome to rule out basal cell carcinoma.

Basal cell nevus syndrome (BCNS or Gorlin syndrome) is an autosomal dominant disorder usually characterized by the early appearance of multiple basal cell carcinomas (BCCs), as well as by other variable clinical features such as odontogenic keratocysts, palmoplantar pits, calcification of the falx cerebri, and a coarse facies [1, 2].

An acrochordon (skin tag or fibroepithelial polyp) is a small, soft, usually pedunculated outgrowth of epidermal and dermal tissue most commonly seen in adults on areas of friction such as the neck, eyelid, axilla, and groin. In clinical practice acrochordons are usually not treated, and it has been written that in most instances they need not be submitted for microscopic examination [3].

Clinical synopsis

Patient 1

A 46-year-old woman with known BCNS was referred to our dermatologic surgery center for Mohs excision of BCCs of the scalp, glabella, and mandible. On the day of surgery, a full body exam revealed five additional suspicious lesions. Two of the lesions were pearly, telangiectatic, pedunculated papules of 2 and 4 mm, respectively, in her right antecubital fossa (Fig. 1). The other three were approximately 1 mm, skin-colored, pedunculated papules on the left side of the patient's neck (Fig. 2).

![Figure 1A](image1.jpg)  
**Figure 1A.** Two pearly, pedunculated, telangiectatic papules of 2 and 4 mm, respectively, in the right antecubital fossa of a 46-year-old woman. Both were BCCs by histology.

![Figure 1B](image2.jpg)  
**Figure 1B.** Typical nodular BCC with some cystic areas. 4x.
Acrochordon-like basal cell carcinomas in patients with basal cell nevus syndrome (BCNS) may present as lesions similar to true acrochordons. These neoplasms were snip-excised utilizing forceps and iris scissors, and the base of each lesion was gently cauterized. Histopathological examination showed both of the antecubital lesions—as well as two of the three neck lesions—to be BCCs. The third neck lesion was a true acrochordon by histology.

### Patient 2

A 12-year-old girl with a new diagnosis of BCNS was referred to our dermatologic surgery center for evaluation of pearly papules on her face, neck, chest, and back. A full body examination revealed nine additional papules in the bilateral axillae ranging in size from 1 mm to 4 mm. These lesions were soft, skin-colored to slightly hyperpigmented, and minimally pedunculated (Fig. 3). They were not telangiectatic or pearly in appearance. Two of the larger lesions were shave-excised and found to be BCCs. Based on the results of these two biopsies, the seven smaller lesions were also excised and found to be BCCs.

On follow-up exam the following month, approximately sixteen additional 1-2 mm growths were noted on the girl’s neck, chest, back, and buttocks. These were skin-colored and inconspicuous (Fig. 4). Each was removed by snip- or shave excision, and—
surprisingly—the majority were BCCs by histology, although a few were acrochordons.

Discussion

Basal cell nevus syndrome is an autosomal dominant disorder usually characterized by early-onset and multiple basal cell carcinomas, along with other variable clinical features such as odontogenic keratocysts, palmoplantar pits, bifid ribs, calcification of the falx cerebri, medulloblastomas, hydrocephalus, mental retardation, spine abnormalities, cataracts, and a coarse facies [1, 2]. About 40 percent of cases represent new germline mutations [4], and a study in the north-west of England found the prevalence of BCNS to be 1 in 55,600 persons [5]. The disorder has been traced to a mutation on chromosome 9 in the PTCH gene (human homologue of the Drosophila “patched” gene) [6], which functions as a tumor suppressor in the hedgehog signaling network [7]. In addition to its role in the cell cycle, PTCH helps control embryogenesis and the development of normal tissue [8].

An acrochordon or skin tag is a benign outgrowth of epidermal and dermal tissue that is most commonly seen in adults on areas of friction such as the neck, eyelids, axillae, and groin. Acrochordons have a reported incidence of 46 percent in the general population [9], and a 1996 study found only 5 of 1335 (~0.4%) clinically diagnosed acrochordons to be malignant (4 BCCs and 1 squamous cell carcinoma in situ) [3]. Because of their high incidence and their low probability of malignancy, acrochordon-like lesions are appropriately left untreated in most patients.

We have reported the cases of two previously diagnosed BCNS patients—one adult and one child—with multiple acrochordon-like papules that proved histologically to be BCCs. These acrochordon-like BCCs had a variety of appearances as shown above, and many of them could have been mistaken for benign growths requiring no treatment. We performed snip- or shave-excisions with 1 to 2 mm margins followed by gentle electrocautery of the base as a quick and clinically effective removal method; no recurrence at the removal sites had been seen clinically as of 10 months post excision. We will continue to follow these patients every 3-4 months throughout their lifetime.

In light of these cases, we recommend the excisional biopsy or destruction of all acrochordon-like lesions on patients with a known history of BCNS. Additionally, we support the conclusion by Chiritescu and Maloney that acrochordon-like lesions found in young patients should be biopsied [10]. Acrochordons are distinctly uncommon in childhood [9], and the discovery of an acrochordon-like BCC in a child can lead to the early diagnosis of BCNS [10]. A patient suspected to have BCNS should follow a regimen of increased sun protection and should see a medical geneticist, in addition to receiving regular dermatologic care.

References

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