Cerebriform intradermal nevus as a cause of cutis verticis gyrata

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Cutis verticis gyrata (CVG) is a rare disease that is characterized by excess scalp skin, producing thick folds that form creases and ridges similar in appearance to the whorls of the cerebral cortex. It was first described in the literature by Robert in 1843 and the name cutis verticis gyrata was proposed by Unna in 1907 and remains the accepted name to date.1,2 It has an estimated prevalence of one case in every 100,000 men and 0.026 cases for every 100,000 women.3 It can be an isolated manifestation that is present from birth or it can be part of other syndromes. Cutis verticis gyrata is classified as primary (subdivided into essential and non essential) or secondary. The essential primary form is not associated with neurological or ophthalmological disorders, and only the folds on the scalp mimicking the cerebral whorls appear. It appears in puberty and affects more men than women. The non essential primary form affects 0.5% of patients with mental retardation. Cerebral palsy, epilepsy, cataracts and blindness may also be present.2,4,5,6

The secondary forms of CVG generally occur as a result of inflammatory or neoplastic processes that cause changes to the structure of the scalp.2,6 One of the rarest forms of secondary CVG is the nevoid form, in which it is melanocytic intradermal nevi that cause the cutaneous hypertrophy. This condition is known as cerebriform intradermal nevus (CIN) and was first linked with CVG in 1937 by Hammond and Ransom.7 Patients affected by CIN have normal intelligence and are free from other local or systemic diseases.

The objective of this paper is to describe a rare case of cerebriform intradermal nevus, discussing its histopathological and clinical features.

Case
A white female patient, 43 years old, presented at the Instituto de Dermatologia Professor Azulay with large dimension lesions on the scalp. She stated that at birth she had had a small normal colored macula on her scalp and that it had gradually developed, reaching its largest size during puberty. On physical examination, multiple patches of alopecia were observed in addition to a normal-colored convoluted mass covering the right parietal, right temporal and right occipital regions and small areas of foul-smelling seborrheic dermatitis (Figures 1A and 1B). Neurological and ophthalmological test results were normal. Family history included no relevant details.

X-rays and computerized tomography of the head did not reveal damage to the calvarium. Blood tests and liver, kidney and thyroid function were normal. A cutaneous biopsy revealed melanocytic intradermal nevi with progressive maturation inwards towards the depth of the tissues. Nests and bundles reached the deep dermis and parts of the subcutaneous zone, thereby indicating the intradermal cerebriform subtype (Figures 2 and 3).

Discussion
Cutis verticis gyrata is a rare disease that is characterized by excess scalp skin, producing folds that are reminiscent of the...
whorls of the cerebral cortex. The different forms of CVG are classified as follows:  

1. **Primary**
   A. Essential: extremely rare form in which only the cutaneous manifestations are observed.
   B. Non essential: associated with mental deficiency, schizophrenia, epilepsy, cerebral palsy, cranial abnormalities (especially microcephaly) and cataracts.

2. **Secondary**
   A. Inflammatory acute or chronic: dermatosis (eczema, psoriasis, impetigo, erysipelas, keloid folliculitis and pemphigus).
   B. Proliferative: cerebriform intradermal nevus, lipoma, neurofibroma, dermatofibroma.
   C. Miscellaneous: acromegaly, pachydermoperiostosis, acromegaly, syphilis, myxedema, cretinism, Ehlers-Danlos syndrome, acanthosis nigricans, tuberous sclerosis, amyloidosis and diabetes mellitus.

**D. Trauma.**

Therefore, CIN is a rare form of secondary CVG, characterized by well-delimited lesions, that affects the parietal region of the calvarium and prefers females. Lesions are normally present at birth as normal-colored or over-pigmented macules that increase in size and stand further proud as the years pass. The lesions gradually begin to take on their brain-like appearance which leads to alopecia in addition to the possibility of itching, bleeding, secondary infections and foul smell, due to bacterial proliferation in the folds. The social and esthetic repercussions of CIN make it extremely important.

With the patient described here, the lesions located in the parietal, temporal and occipital areas are in line with the majority of reports in the literature. Furthermore, although rare, the brain-like lesions can affect other areas and cases have been described where the neck and scrotum were involved as a case where a malignant peripheral melanoma was associated with a giant melanocytic intradermal nevus. With relation to size, CIN can vary from 2 x 3 cm to 22.5 x 25 cm. Diagnosis of CIN is based on clinical findings and confirmed by histopathology. Characteristics of a positive histopathological result are: nevocytes arranged intradermally with a minimal junctional component, possibly affecting superficial parts of the hypodermis; adjacent structures enveloped by nevocytes; loss of the typical arrangement of nests in stretches; a prevalence of type C nevus cells with large areas of neuroid differentiation; irregular arrangement and distribution of intracellular melanin pigment and perivascular nevocyte aggregation.

O differential diagnosis is to rule out primary CVG, the other forms of secondary CVG and other pathologies such as: cerebriform nevus sebaceous and aplasia cutis congenita. Treatment consists of surgical excision with plastic reconstruction where necessary. Small lesions can be removed by simple excision and suture. Where lesions are larger, flap rotation or serial surgeries should be used, with or without an expander or other techniques, minimizing esthetic deformities.

The possibility that these lesions may undergo considerable growth during puberty means that early excision and suture should be considered in order to avoid extensive surgery later in life. Another argument in favor of removal of CIN is the possibility of melanomas, although they are rare.

**References**
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