Von Recklinghausen’s Disease with a Typical Features

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Abstract
A 42-year-old man consulted complaining of an irritated skin-colored nodule to the left side of his head that was identified as a neurofibroma. Numerous similar lesions were found to his torso and neck. He had no dysmorphic features or bone abnormalities, but he was found to have left axilla and right inguinal freckling, long soft-brown asymmetric skin patches of irregular borders on his torso that crossed the midline with no fullness under the skin. His left iris had some orange-brown colored specks. He had normal blood pressure and no neurological abnormalities. MRI of his body showed no plexiform neurofibromas. Nobody else in his family had similar findings. Based on his clinical findings, a diagnosis of a mosaic form of Neurofibromatosis type 1 (von Recklinghausen’s disease) was made, as a de novo mutation.

Case Presentation
A 42-year-old man consulted complaining of a longstanding soft flesh-colored nodule on the left side of his head that was irritated and sometimes bled after combing his hair (Figure 1). He remembered having a similar lesion removed from his chest area 15 years prior in another country that was not sent for pathologic study. His physical examination revealed numerous similar skin-colored nodules on his torso, behind both ears and neck (Figures 2 and 3). His exam demonstrated neither dysmorphic features, nor scoliosis or tibial bowing. Freckling was found to his left axilla
Neurofibromatosis affects 1 in 3500 people, is a multisystem autosomal dominant condition where tumors are developed along the course of peripheral nerves, and soft-tissue and bone deformities can also be present [1,2]. The majority of patients are asymptomatic but some can present with neurological or bone complaints. The expression of the disease is highly variable among family members with the same mutation [3]. The diagnosis is made on a clinical basis although mutational analysis and molecular testing are now available for specific cases [1].

The described nodular lesions are neurofibromas, which are benign tumors of Schwann cells, perineural cells and fibroblasts. The superficial cutaneous manifestations are hallmarks of this disease. In some cases, firm subcutaneous neurofibromas can be found. Diffuse subcutaneous masses (diffuse plexiform neurofibromas) can produce disfiguring deformities, and lesions involving spinal nerve roots (nodular plexiform neurofibromas) can produce neurological symptoms. Neurofibromas have the potential to become malignant. The patients suffering from this disease also present with skin macules ranging from soft-brown (café-au-lait spots) to freckling patches that can be appreciated in the trunk, pelvis or extremities [4]. Skin fold freckling (Crowe sign) is the most specific finding for this condition [1]. The mentioned eye findings (orange-brown colored specks) are Lisch nodules which are benign hamartomas that can be seen without magnification. Slit lamp examination can differentiate them from nevi on the iris by demonstrating elevated lesion instead of flat ones. Other associated ophthalmologic findings are optic gliomas [2]. Lisch nodules occur in 90% of adults with neurofibromatosis 1. Optic gliomas that can alter color vision and can produce progressive sight loss [1]. Skin findings are usually bilateral but asymmetrical presentation can be seen in mosaic forms of the disease (when not all of the patient’s cells carry the genetic change in the NF-1 gene) as is the case of the presented patient. In 50% of cases the genetic mutation is inherited from a parent and 50% of the time the mutation is de novo [6,7].

Cognitive problems can be present in 60% of cases including difficulties at school, learning disabilities or attention deficits. Possible associated neurological symptoms include weakness, numbness and paresthesias in any extremities or parts of the body, as well as headaches in 20% of patients and seizures in up to 10% [2]. Some cardiovascular manifestations include hypertension, which suggests renal artery stenosis (1% of patients), and vascular dysplasias most frequently affecting the aorta and carotid arteries [1]. Bone abnormalities include scoliosis in 10% of patients [5], pseudoarthrosis (mainly of the lower third of tibia and fibula), bowing of legs, sphenoid wing dysplasia, fibrous dysplasia and subperiosteal bone cysts [1,4]. If the patient presents with neurological symptoms, an MRI or CT scan of the affected area can determine the presence of underlying plexiform neurofibromas. Treatment is required if the neurofibromas
cause severe symptoms, disfiguration or if they are irritated by their external location. Surveillance for malignancy is recommended.

**Authors’ Contributions**

JAGR is the primary health provider for the patient, conceived, designed, compiled the data for the article and wrote the article.

**Authors’ Information**

JAGR is the primary health provider for the patient, who practices Family Medicine and Sports Medicine. He is an Assistant Professor at the Family Medicine Department of the University of Calgary.

**References**


<p>| Table 1: Differential Diagnosis: Diagnostic Features of Neurofibromatosis type 1 and 2. |</p>
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<th>Condition</th>
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| Neurofibromatosis type 1  
(Two or more are required to make the diagnosis) | 1. Six café-au-lait spots of 5 mm or more in longest diameter in prepubertal patients and 15 mm in the longest diameter in postpubertal patients.  
2. Two or more neurofibromas of any type or 1 plexiform neurofibroma.  
3. Freckling in the axilla or inguinal regions.  
4. Optic glioma (optic pathway glioma).  
5. Two or more Lisch nodules (iris hamartomas).  
6. Distinct osseous lesions, such as sphenoid wing dysplasia or cortical thinning of the cortex of long bones with or without pseudoarthrosis.  
7. First-degree relative (parent, sibling, or child) with NF1 according to the above-listed criteria. |
| Neurofibromatosis type 2 | 1. Bilateral vestibular schwannomas  
2. Unilateral schwannoma by 30 years plus:  
A. A – A parent, child or sibling suffering NFM type 2, or  
B. B – At least two of the following: Meningioma, glioma, schwannomas, or juvenile posterior sub capsular lenticular opacity/juvenile cortical cataract. |

*Information from reference [7]*