Clinical findings of the phakomatoses: Neurofibromatosis

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The two common types of neurofibromatosis (NF) are genetically and clinically distinct diseases, each of which promotes the growth of neurofibromas and schwannomas that arise in the myelin nerve sheaths of peripheral or cranial nerves. Other tumors may arise from a variety of tissues including meningiomas, optic gliomas, and ependymomas. Both NF1 (chromosome 17q) and NF2 (chromosome 22q) are dominantly transmitted, with half of cases arising from new mutations. Diagnostic criteria for NF1 and NF2 have been developed by consensus. Physical features and neuroimaging of NF are shown in figures 1 and 2.

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References

Figure 1. (A) Café au lait (CAL) macules on the back of a six-year-old boy; six or more that are >5 mm in size in prepubertal children support a diagnosis of NF1, being present in 80% of cases by two years and 100% by ten years. (B) This adult has characteristic “axillary freckles” (1–4 mm) as well as larger CALs (found in 70% of NF1 by ten years). The inguinal region is also a common site for freckles. In some individuals, these small CALs become more confluent. (C) Multiple cutaneous neurofibromas may be found in NF1 (5%–10% by three years). They may arise anywhere in the skin. When small, invagination of these soft lesions with a finger discloses a “button-hole”-like orifice at the base. Two or more support a diagnosis of NF1. (D) Plexiform neurofibromas, present in 15%–30% of NF1, arise in a diffuse manner that involve multiple nerves, typically but not limited to the inguinal, or in this example, the shoulder region, and may be associated with cutaneous hyperpigmentation and skin laxity. (E) Patients with periorbital neurofibromas may develop visual changes resulting from diplopia, glaucoma, and amblyopia. (F) Lisch nodules (arrows), yellow-brown hamartomas within the iris, are found in 80% of NF1 cases by ten years.2

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Figure 2. (A and B) Right optic nerve gliomas (arrows) that involve the optic chiasm (arrowhead) on axial T1-weighted MRIs with contrast in an asymptomatic adult with NF1. These occur about in 10%–20% of patients with NF1 and are usually discovered in symptomatic children. They are rarely symptomatic if discovered after age ten. (C) Bilateral cranial nerve VIII schwannomas on axial T1-weighted MRI with contrast are the hallmark of NF2. Unilateral acoustic schwannomas may occur in NF1 but when bilateral are diagnostic of NF2. (D) Photomicrograph of a resected acoustic schwannoma shows characteristic “Verocay bodies”—palisades or whorls of cylindrical or rod-shaped nuclei. (E) Multiple meningiomas, as seen in this adult, on contrasted T1-weighted axial images, occur more commonly in NF2, as do astrocytomas or ependymomas.
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