Bilateral optic nerve sheath meningiomas in a patient with neurofibromatosis type 2

I A Cunliffe, D A Moffat, D G Hardy, A T Moore

Abstract
A 34-year-old woman who presented with hearing loss and tinnitus was found to have reduced vision bilaterally. Computed tomography scan revealed bilateral acoustic neuromas and bilateral optic nerve sheath meningiomas. The presence of bilateral acoustic neuromas fulfils the criteria for the diagnosis of central neurofibromatosis (neurofibromatosis type 2). Although this is the first report of bilateral optic nerve sheath meningioma in neurofibromatosis type 2, meningiomas are commoner in this dominantly inherited disorder, than in its absence and both forms of central nervous system tumour may be caused by loss of tumour suppressor genes on chromosome 22.

Optic nerve sheath meningiomas are uncommon, and though they may present at any age they are predominantly seen in middle aged females. They are unilateral in most cases. The usual presentation is with slowly progressive visual loss, and most patients have evidence of field loss and optic atrophy or disc swelling at presentation. Opticociliary shunt vessels are found on the optic disc in about 20% of cases. Other findings include mild axial proptosis, limited ocular movements, and gaze evoked amaurosis. Optic nerve meningiomas are thought to be more common in patients with neurofibromatosis than in general, when they are usually unilateral. Bilateral optic nerve sheath meningiomas are rare and in none of the reported cases has there been any evidence of neurofibromatosis.

We report on a patient with neurofibromatosis (type 2) who had evidence of bilateral acoustic neuromas and bilateral optic nerve sheath meningiomas, causing a dual sensory impairment.

Case report
A 34-year-old woman presented with an 18-month history of tinnitus, increasing deafness in the right ear, and numbness of the right cheek. She had been otherwise well and the only other relevant history was that she had a longstanding right convergent squint and associated amblyopia. She had undergone surgery for right squint at age 7 years.

On examination she had reduced hearing in the right ear, reduced sensation in the distribution of the second division of the right trigeminal nerve, and evidence of mild ataxia. The visual acuity was counting fingers in the right eye and 6/6 in the left eye. Colour vision testing with Ishihara plates was normal in the left eye, but she was unable to recognise any plates with the right. There was a 3 mm right proptosis, and a cover test revealed a small right divergent squint. Ocular movements of the right eye were full, but she had limitation of elevation,

Figure 1 Photographs of optic discs showing bilateral pallor and mild swelling of nerve fibre layer. A: Right disc. B: Left disc.
abduction, and adduction of the left eye. There was bilateral gaze-evoked nystagmus. Corneal sensation was reduced on the right but normal on the left. The lenses were clear, and intraocular pressures were normal. Both pupils reacted to light, and there was no relative afferent pupillary defect.

Slit-lamp examination of the iris was normal, with no evidence of Lisch nodules. Both optic discs were pale and slightly swollen but without any opticociliary shunt vessels (Figs 1A, B). In each eye there was a small slightly raised pale subretinal lesion, with associated fibrosis, lying temporal to the macula, probably representing a choroidal hamartoma. Goldmann perimetry revealed a dense right central scotoma and generalised constriction of the left visual field. Visual evoked potentials were unrecordable to both flash and pattern stimuli in the right eye. In the left eye the only abnormality was a mildly prolonged latency of the pattern VEP.

Computed tomography (CT) scan showed a large right acoustic neuroma and a small left acoustic neuroma. In addition there were bilateral optic nerve lesions. The left optic nerve showed a large calcified mass, and there was an enlarged irregular thickening of the right optic nerve, consistent with bilateral optic nerve sheath meningiomas (Fig 2). Systemic examination revealed no other evidence of neurofibromatosis, and there was no family history of hearing loss or central nervous system (CNS) tumours.

The right cerebellopontine angle tumour was excised via a translabyrinthine approach, and histological examination confirmed it to be a benign schwannoma. A postoperative magnetic resonance imaging scan showed the right tumour to have been completely excised and confirmed the presence of bilateral optic nerve tumours with no evidence of chiasmal involvement.

Discussion

The neurofibromatoses are divided into two distinct disorders: neurofibromatosis 1 (Recklinghausen’s disease, NF1) and neurofibromatosis 2 (bilateral acoustic neurofibromatosis, NF2). Both are inherited as autosomal dominant traits with a high degree of penetrance. The genes responsible for each type have been mapped, to chromosome 17 for NF1 and chromosome 22 for NF2. The diagnostic criteria for each disorder are given in the Neurofibromatosis Conference Statement. The criteria for NF2 are met if the patient has (1) bilateral eighth nerve masses, or (2) a first degree relative with neurofibromatosis 2 and either a unilateral eighth nerve mass, or two of the following: neurofibroma, meningioma, glioma, schwannoma, or juvenile posterior subcapsular lens opacity. The patient reported here fulfills the criteria for the diagnosis of NF2.

Patients with NF2 typically present with either unilateral or bilateral hearing loss, and most have evidence of disturbed vestibular function. Large tumours may involve lower cranial nerves or result in raised intracranial pressure. The ophthalmic manifestations include juvenile posterior subcapsular lens opacities, corneal abnormalities associated with loss of corneal sensation or seventh nerve palsies, papilloedema from raised intracranial pressure, nystagmus, sixth nerve palsy, and less commonly visual pathway tumours as in this case. In contrast to NF1 Lisch nodules are rarely present. In NF2 there is an increased incidence of other central nervous system tumours, which may be of schwann cell, glial, or meningeal origin.

Although the mechanism of tumour formation is unclear, similarities with the development of non-ocular tumours in the hereditary form of retinoblastoma suggest that there may be loss of tumour suppressor genes in NF2. Acoustic neuromas (like retinoblastoma), occur in a hereditary and non-hereditary form. Patients with the non-hereditary form have unilateral tumours which present later, and do not develop other CNS tumours. In patients with NF2 there is often a family history, the tumours are bilateral and present earlier and there is an increased risk of developing other CNS tumours, particularly meningioma. Both meningiomas and acoustic neuromas show loss of chromosome 22 material in the tumour tissue, suggesting that in patients with NF2 (who have a germlinal mutation of chromosome 22) tumour formation may result from the loss of the normal allele as a result of a somatic cell mutation. It is unclear, however, whether the mutations responsible for acoustic neuromas and meningiomas are at the same or different loci on chromosome 22. Although unilateral optic nerve meningiomas have been reported in neurofibromatosis, most reports predate the recognition of the two forms of the disorder, and it is not clear whether these optic nerve tumours are seen predominantly in peripheral or central NF. In the family with NF2 reported by Dellemann et al one affected member had a unilateral optic nerve sheath meningioma but without acoustic neuroma. The patient reported here is further evidence that optic nerve sheath meningiomas may develop in NF2 and may result from the loss of normal genes responsible for regulating growth of neural tissues.

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doi: 10.1136/bjo.76.5.310

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