SHORT REPORT

Neuroenteric cyst of the optic nerve: case report with immunohistochemical study

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Abstract
A 36 year old man developed slowly progressive unilateral visual loss due to a cystic lesion of the intraorbital optic nerve. Pathologically the lesion was an epithelial lined cyst entirely within the atrophic nerve. The cyst lining consisted of columnar epithelium partly pseudotratified and ciliated with evidence of mucin secretion, and was immunoreactive for cytokeratin but not glial fibrillary acidic protein. This lesion is considered to be similar to neuroenteric cysts that have been reported elsewhere in the nervous system, and the immunocytochemical results are consistent with a heterotopia derived from the primitive ectoderm of the stomatodeum.

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Cystic lesions of the optic nerve are rare. They can occur in association with gliomas or meningiomas, from extension of a craniopharyngioma or from the arachnoid. A variety of cysts with an epithelial lining have been described in the nervous system in many locations1–7,12–20,24–27 but there has been only one report of a cyst of this type in the optic nerve.10 The classification of these cysts, and hypotheses about their origin have depended on their histology and their location. This case is of an epithelial cyst within the optic nerve. The cyst was removed in its entirety, extensively studied histologically and characterised by immunocytochemical staining.

Case report
The patient presented at the age of 19 when he was noted to have a visual acuity (VA) of 6/9 in the left eye and a pale optic disc. He attended irregularly for follow up. Fourteen years later the VA had fallen to 6/12. He declined investigation and was seen again two years later, now at 36 years, at which point his vision had fallen to 6/36. At this time he complained of progressive blurred vision over the previous 2 years and left frontal headaches. His past medical history and family history were unremarkable.

On examination the right eye was entirely normal. The visual acuity had decreased to 6/36 in the left eye, with a small central area of preserved field. The optic disc on the left was pale. The remainder of the neurological examination was normal. CT (fig 1) showed a well defined intrinsic mass enlarging the posterior intraorbital portion of the optic nerve which was felt to represent either a glioma or meningioma. T2 weighted MRI scans confirmed this. An orbito-cranial exploration was performed and the optic nerve inspected. The posterior orbital optic nerve was distended and cystic, partly extending into the optic canal and, on incision, a milky fluid was released. The affected optic nerve was totally excised leaving the dura intact. Postoperatively the patient recovered uneventfully.

Pathological examination
The surgical specimen consisted of 2 cm of optic nerve measuring 2.5 mm in diameter. This appeared normal, presumably due to decompression of the intraneural cyst surgically, and the gross appearance was unremarkable. The tissue was embedded in paraffin and transverse and longitudinal sections were stained with routine methods. Immunocytochemical staining for glial fibrillary acidic protein (GFAP, Dako Ltd, UK) carcinoembryonic antigen (CEA, Dako Ltd, UK), cytokeratin (CAM 5.2 Becton Dickinson, USA), epithelial membrane antigen (EMA, Dako Ltd, UK), neuron specific enolase (NSE), (BAKO Ltd, UK) and synaptophysin (Dako Ltd, UK) was performed using the avidin-biotin method. CAM 5.2 recognizes keratin subgroup 8,18 and 19.11 On microscopic examination an irregular slitlike cavity extended to much of the length of the nerve (fig 2a); the cyst occupied almost the whole width of the nerve, without reaching its edges. The lining of the cyst varied from columnar to pseudostratified epithelium, the latter with cilia in most areas, which covered a thin but well developed matrix of collagen fibrils. These were highlighted by reticulin stains, but were PAS negative. With PAS staining it was also apparent that there were a few scattered mucin secreting goblet cells. No evidence of squamous differentiation suggestive of a dermoid or epidermoid cyst and no mitotic figures were seen. The epithelium stained weakly with CEA, and intensely with cytokeratin (fig 2b) but did not react with EMA, NSE, synaptophysin or GFAP, although gliotic tissue surrounding the cyst was immunoreactive for the latter. The surrounding nerve contained only a few myelinated axons along one edge, and was for the rest largely devoid of myelin sheaths.
Discussion

Cystic lesions of the optic nerve are infrequently described in the literature. Examples have been reported in conjunction with neoplasms, particularly craniopharyngiomas involving the chiasm, and gliomas. Instances in which the cystic lesion is clearly an extension or part of the tumour. Meningiomas may produce cystic dilatation of the subarachnoid space around the optic nerve presumably by trapping CSF. Cystic lesions have also been described arising from the coverings of the optic nerve, the so-called arachnoid cysts which are extrinsic to the nerve. Recently a single case, similar to the present one was described as a neuroepithelial cyst of the optic nerve and studied with routine histological methods.

Cysts lined by a mucus secreting ciliated columnar epithelium have been found throughout the nervous system and named according to the location. Thus lesions essentially indistinguishable on histological grounds have been referred to as neuroepithelial cysts in the spinal cord and posterior fossa, neuroepithelial cysts in the hemispheres, colloid cysts of the third ventricle and Rathke cleft cysts in the sella. The histological features at each of these locations cover a similar spectrum but are variable and non-specific. Typical cysts are lined by a columnar epithelium with transition to a low cuboidal and beamed up to a pseudostratiﬁed pattern in other areas covering a collagenous basement membrane. Cilia may be present and, when abundant, the appearances of the epithelium are similar to those of respiratory mucosa; this similarity is further enhanced by the presence of mucus secreting PAS positive goblet cells. Ultrastructural studies have not revealed any features which reliably distinguish cysts at one location from any other.

The pathological features of the cyst in this report ﬁt the general pattern. The possible origin of such cystic lesions has been variously attributed in relationship to the partcular location in which each occurs. It has been suggested that they are trapped embryonic remnants—thus enteric remnants in the spinal cord, ependymal remnants in the forebrain and stomodeal remnants in the hypophysis. Results from a retrospective unpublished study of 7 "neurenteric cysts" of the spinal cord and posterior fossa and 3 hemispheric colloid cysts indicate a pattern of staining similar to that seen in the present lesion, that is, reactivity for either cytokeratin or EMA, and uniform absence of GFAP.

The occurrence of an epithelial lined cyst in the embryological point of view. Arguments based on embryonic connections, between the neural tube and mesoderm or endoderm do not apply at this site. The location seems an implausible one for remnants of the gut, the choroid plexus, the stomodeum, or the paraphysis. Since the eye and the anlage of the optic nerve develop as an outpouching of the diencephalic neural tube, the entrapment

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of primordial ependyma or neuroepithelium seems most likely. Although NSE antibody is not regarded as absolutely reliable, the negative results obtained with it as well as with synaptophysin, rules out the neuroepithelial nature of the epithelium of the cyst. Moreover, there are also obvious differences between the mature ependymal lining and the lining of this cyst. The mature ependyma is usually a simple non-ciliated cuboidal epithelium and, like the epithelium in this specimen, does not have a basement membrane. Immature and reactive ependyma as well as ependymal neoplasms are reactive for GFAP but normal mature ependyma is not.\(^6\) Immunocytochemical staining was performed with this question in mind. The cyst epithelium in the present case was non-reactive for GFAP, which is difficult to reconcile with a trapped remnant of the ependymal lining; furthermore, the reactivity for CAM 5.2 suggests a relationship to an endodermal epithelium, although the reactivity of ependyma with this marker has not been fully explored.\(^6\)

It is clear that in our case there are significant differences between cells derived from the ependyma and the lining of the cyst. The question remains whether or not these can be explained by the degree of maturation and influence of the surrounding tissues or whether the origin of the cells in this case comes from ectopic stomatodeum of Rathke's pouch. Whilst the epithelial characteristics of the cyst would certainly be compatible with an endodermal origin, the method of origin within the optic nerve remains to be explained.

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1 Anderes JP. Neuroenteric cysts of the spinal cord and brainstem, MD Thesis, Lausanne, 1984.