Abstract

Filum terminale hemangioblastoma is an extremely rare tumour with only a few case reports in the literature. The natural history is unknown but benign as well as more aggressive presentations have been reported. The authors present the first such lesion discovered incidentally and discuss the available literature.

Key words: Hemangioblastoma; filum terminale; cauda equina; incidental.

Introduction

Hemangioblastomas of the filum terminale are exceptionally rare tumors. Previous clinical presentations have included low back pain, radiculopathy and even partial cauda equina syndrome (Wolbers et al., 1985, Nadkarni et al., 2006). We report the first case of such a lesion discovered incidentally.

Case report

A 75 year old man presented to his family doctor complaining of general malaise and lethargy for 6 months. He had no specific symptoms and clinical examination was normal. He had no back pain. MRI of his cranio-spinal axis revealed a solitary homogeneously enhancing lesion with well demarcated margins within the cauda equina behind the body of L3 (Fig. 1a).

It was felt that the lesion should be surgically removed, largely because of its uncertain significance. A laminectomy was performed from L2-4 and a classical ‘cherry-red’ lesion was found adherent to the filum terminale (Fig. 1b). A feeding artery and vein were seen entering the tumor along the filum terminale from above. The tumor was removed ‘en bloc’ by dividing the filum above and below. The tumor measured $1.4 \times 1.3 \times 1.1$ cm and demonstrated the typical microscopic pattern of numerous plexiform anastomosing capillary channels on reticulin staining. The patient suffered no complications.
Discussion

The term ‘hemangioblastoma’ was coined by Harvey Cushing and Percival Bailey in 1928 in the book ‘Tumors Arising from the Blood-Vessels of the Brain’. The first report of such a lesion in the filum terminale was in 1943 by Wyburn-Mason (Wyburn-Mason, 1943) in a 25 year-old woman with radicular pain. Since then there have been only 6 other reported cases in the literature, all presenting with back pain or nerve root compression. Hemangioblastomas are notoriously vascular structures, a quality which means they may enlarge (Ortega-Martinez M. et al., 2007) or bleed profusely during surgical resection. As a consequence previous authors have performed pre-surgical embolisation via the anterior spinal artery when these lesions have been discovered in the cauda equina (Biondi et al., 2005; Nadkarni et al., 2006). All reported cases of hemangioblastoma of the filum terminale have resulted in uncomplicated total surgical excision without neurological deficit whether embolised pre-operatively or not. Some have reported ‘easy’ removal of such tumors with minimal blood loss (Tibbs 1999, Biondi 2005). Surgical excision in our case was similarly straightforward. Asymptomatic lesions present a dilemma. The rarity of this condition precludes a comprehensive understanding of its natural history. Tibbs et al. reported a case of filum terminale hemangioblastoma which had partially eroded the superior articulating processes of the L2 vertebra bilaterally causing back pain, suggesting the lesion had been present for a very long time (Tibbs et al., 1999). Other authors have described a more malevolent history with partial cauda equina syndrome (Wolbers et al., 1985; Nadkarni et al., 2006). The avid enhancement on MRI should raise suspicion of this pathology and alert the surgeon to be prepared for a vascular lesion.

Conclusion

The authors advocate surgical excision for this lesion in patients fit for surgery. Pre-operative embolisation is not necessary, but may be useful for intramedullary hemangioblastomas of the conus or spinal cord.

REFERENCES


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