

Myxopapillary Ependymoma Revealed by Dysphagia (A Case Report)

Dr. Touati Lina, MD¹; Pr Jabbouri B., PhD²; Pr El Kacemi I., PhD ; Pr Hakkou M. PhD ; Pr Oudrhiri Y. PhD ; Pr Boutarbouch M. PhD ; Pr Melhaoui A. PhD ; Pr. Arkha Y. PhD

Department of Neurosurgery , Hopital des Spécialités, Rabat, Morocco

Abstract :

Myxopapillary Ependymomas are slow growing tumors of the cauda equina with a high incidence in young adult. We report the case of a 35-year-old male, who consulted our emergency for a difficulty swallowing food . Upon further investigation, the patient complained of a gradually worsening heaviness of the lower left limb which has since spread to all the other limbs, making it difficult to walk. In addition , the patient had reported notions of drip urination and chronic constipation. Physical examination revealed numbness of the 4 limbs with normal and symmetrical reflexes , as well as thermoalgnesia and hypoesthesia on the left of the abdomen . During cranial nerve examination, a vertigo and dysphagia were noted .

A T1-Weighted MRI showed a hyposignal intradural tumor from L2 to L4 with a strong enhancement after gadolinium injection.

A T2-Weighted MRI showed a hypersignal tumor.

Additionally , there was a syringomyelic cavity clearly visible which extended from the medulla oblongata to the conus medullaris.

An L2 L3 laminectomy was performed on the patient with a gross total resection of the tumor that was attached to the cauda equina roots.

Upon good clinical recovery , the patient was discharged after satisfactory clinical recovery.

An MRI performed 6 months later showed a total resection of the tumor.

Ependymoma of the filum terminale associated with holocord syringomyelia are a very rare entity with a still uncertain pathogenesis.

Introduction :

Intramedullary spinal tumors are well known to be associated with secondary syringomyelia. Ependymomas and hemangioblastomas are the most common tumors associated with syringomyelia. However, holocord syringomyelia secondary to intradural extramedullary spinal tumors is extremely rare. We present one such rare case.

Case Report :

We present the case of a 35 year old male patient whose medical history includes a birth related left brachial plexus lesion which required physiotherapy . The patient suffered a 6 year long gradually worsening of his already existing heaviness of the left upper limb . The heaviness started affecting his lower left limb before spreading to the opposite side . In addition , the patient had reported notions of drip urination and chronic constipation .

A week before his admission to our institution , the patient consulted our emergency team for a recent difficulty swallowing food .

Upon examination , the patient was conscious with a tetraparesis making it difficult to walk , as well as thermoalgesia and hypoesthesia on the left side of the thorax and the abdomen . During cranial nerve examination, a vertigo and dysphagia were noted .

A T1-Weighted MRI of the spine(figure 1) showed a hyposignal intradural tumor from L2 to L4 with a strong enhancement after gadolinium injection .

A T2-Weighted MRI (figure 2) showed a hypersignal tumor .

Additionally , there was a syringomyelic cavity clearly visible which extended from the medulla oblongata to the conus medullaris . (Figure 3)

Contrast-enhanced MRI revealed a well-defined intradural extramedullary mass at the level of L2/L4 vertebrae with inhomogeneous postcontrast enhancement with associated holocord syrinx .

An L2 L3 laminectomy was performed on the patient with a gross total resection of the tumor that was attached to the cauda equina roots . (Figure 4)

Intraoperatively, the tumor was greyish-white, moderately vascular, and encapsulated with a good plane of cleavage between the tumor and the cauda equina roots(Figure 5) . Histopathological examination confirmed the diagnosis of ependymoma. He was doing well at the last follow-up six months after surgery and his symptoms had improved significantly with no sign of dysphagia . MRI of the spine done 6 months later did not show any evidence of residual tumor (figure 6) .



Figure 2: A T2-Weighted MRI

Figure 1: T1-Weighted MRI of the spine showing a hyposignal intradural tumor showing a hypersignal intradural tumor



Figure 3 : syringomyelic cavity
Laminectomy L2 L3

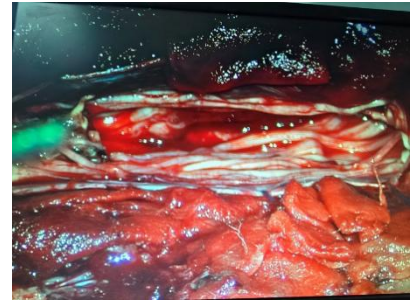


Figure 4 : Intraoperative picture showing
Gross total removal of the tumor after

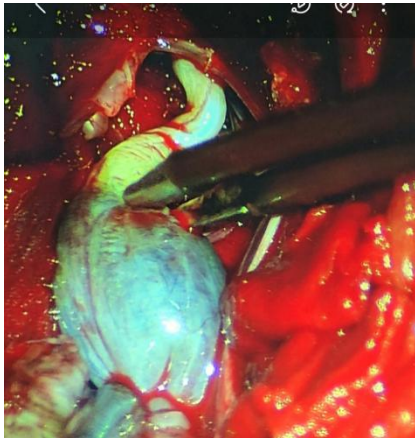


Figure 5 : the tumor was greyish-white, moderately vascular

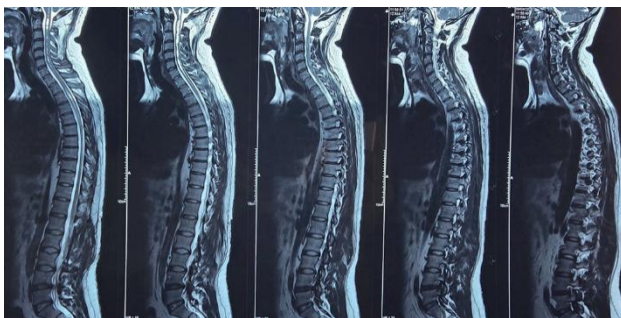


Figure 6 : post op MRI of the spine done 6 months later

Discussion :

Syringomyelia associated with spinal tumors is commonly reported with hemangioblastomas and ependymomas (1) . In most cases , tumors are intramedullary located in the cervicothoracic region. However, tumors which are located in the distal end of the spinal cord rarely cause syringomyelia (2) . In our case , the tumor was purely extramedullary and arose from the cauda equina with extensive holocord syringomyelia from the medulla oblongata to the conus medullaris .

The presentation of syringomyelia depends on the extent of spinal involvement , going from motor and sensory weakness to cranial nerves involvement in case of syringobulbia (3) , such as with our case .

The pathogenesis of syringomyelia associated with spinal cord tumors is still not well understood . It appears to be most likely due to transudation of fluid from the pathological tumor vessels(4). The other possible mechanisms include obstruction to CSF flow in the central canal and extracellular perimedullary fluid flow (5). However, the reason is still uncertain .

Our case was unique as to the presentation of the syringomyelia with initial weakness of the upper limbs associated with dysphagia that led the patient to consult . As in our case , the initial presentation of ependymoma may be a complication of syringomyelia . Initial presentation with only bulbar palsy is also possible and was prior reported in a 16 year old female patient(6) . Likewise , in all other similar cases that we came across , the removal of the tumor was sufficient for the resolution of syringomyelia without any additional procedure (2) .

Conclusion

A unique patient with extramedullary myxopapillary ependymoma associated with extensive syringomyelia has been reported. Serial MRI demonstrated the total removal of the lesion and clinical recovery has been promising . However, the reason for the occurrence of syringomyelia associated with the tumor is still uncertain. Furthermore , the removal of the tumor was sufficient for the resolution of symptoms .

References

- (1)Borkar, Sachin A.; Satyarthee, G. D.; Sharma, B. S Department of Neurosurgery, All India Institute of Medical Sciences, New Delhi, India
- (2)Yavuz Samanci, Suat Erol Celik Neurosurgery Department, Ministry of Health Okmeydani Education and Research Hospital, Istanbul, Turkey
- (3)Nagahiro, Shinji M.D.; Matsukado, Yasuhiko M.D.; Kuratsu, Jun-ichi M.D.; Saito, Yoshiki M.D.; Takamura, Seishi M.D.

(4)Paul N.M. Lohle^a, Hans A.L. Wurzer^b, Peter H. Hoogland^a, Petrus J. Seelen^c, K.Gwan Department of Radiology, Westeinde Hospital, The Hague, The Netherlands Department of Neurosurgery, Westeinde Hospital, The Hague, The Netherlands Department of Clinical Chemistry, Westeinde Hospital, The Hague, The Netherlands Department of Neurosurgery, University of Groningen and University Hospital, Oostersingel 59, 9713 EZ Groningen, The Netherlands

5) A. Leclerc^{a b}, L. Matveeff^{a b}, E. Emery^{a b c}CHU Caen, Department of Neurosurgery, Avenue de la Côte-de-Nacre, 14000 Caen, France Université Caen Normandie, Medical School, 14000 Caen, France INSERM, UMR-S U1237, Physiopathology and Imaging of Neurological Disorders (PhIND), GIP Cyceron, 14000 Caen, France

(6)Smriti Sinha, Prathibha Shankar Ashwini, Pelala Nayan Baba, Rathika Damodar Shenoy
Department of Pediatrics, K. S. Hegde Medical Academy (KSHEMA), Mangaluru, Karnataka, India