Calcified Pseudoneoplasm of the Neuraxis (CAPNON): A Lesson Learnt from a Rare Entity

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Abstract

We present the case of a 39-year-old female with an exceptional rare entity of the neural axis located in the perisylvian region. We discuss the unique features and the subsequent management of this interesting case.

Keywords: Calcified pseudoneoplasm of the neuraxis; CAPNON; Oedema; Brain stone

Case Report

A 39-year-old, right hand dominant, female presented with tingling sensation in her right arm followed by a first-time, self-terminating generalized tonic-clonic seizure. There was no previous medical history and no regular medications. Detailed neurological examination was unremarkable and there were no signs elicited from cardiorespiratory or abdominal examination. Computed tomography (CT) of the brain revealed a hyper dense calcified mass in the left perisylvian region (Figure 1). Magnetic Resonance Imaging (MRI) showed hypointense signal on T1 and T2-weighted images in the region of calcification as seen on CT. Extensive vasogenic oedema was also noted in adjacent lobes and extended into the external capsule and lentiform nucleus. Prominent vessels were seen surrounding the lesion but no vascular anomaly was noted on Magnetic Resonance Angiography (MRA). A Digital Subtraction Angiogram (DSA) ruled out aneurysm or arterio-venous malformation (AVM). The differential diagnosis remained low-grade astrocytoma versus oligodendroglioma.

Following multidisciplinary discussion surgical intervention was recommended. The patient underwent a left fronto-temporal craniotomy and gross total resection of the calcified mass. The sylvian fissure was noted to be wide open and surrounded by prominent vessels. The calcified mass was removed in three pieces and abnormal looking surrounding brain was biopsied; it had a rubbery consistency with grey-yellow appearance.

Post-operatively the patient was noted to be significantly weak in the right upper and lower limbs. MRI confirmed restricted diffusion in the parietal lobe. Histological diagnosis confirmed a calcified pseudoneoplasm of the neural axis (Figure 2).
At one month follow up the patient remained densely monoplegic in the right upper limb but now able to move anti-gravity in the right lower limb. She remained on anti-seizure medication and continued intensive Neuro-rehabilitation.

Discussion

Calcified pseudoneoplasm of the neuraxis (CAPNON) is a rare entity and is considered benign. Review of previous reports suggests similar incidence in both sexes occurring in different locations intra-and extra-axially [1]. To our knowledge none have previously been reported in the perisylvian region. Presentation varies depending on location but can involve, as seen in this case, sensorimotor deficit or seizures [2]. CAPNON is typically mistaken for more common neoplastic or non-neoplastic lesions and CT/MRI helps to distinguish between the different types. On CT, CAPNON is classically hyper dense given the calcification while on MRI imaging lesions tend to show hypo-intense signal on T1 and T2 weighted sequences [1]. Surrounding vasogenic oedema is not commonly reported and would not typically be expected from a benign lesion. Grabowski et al. reported vasogenic oedema in their case report and suggest seizures as the possible cause [2]. Similarly our patient presented with seizures although the cerebral oedema was more extensive.

Histologically CAPNON contains variable portions of fibrous stroma, bone, palisading spindle and epithelioid cells and a chondromyxoid-like matrix [1]. The etiology and natural history of CAPNON is unclear and thought to result from a reactive process. This is suggested by associated granulomatous inflammation typically surrounding the calcified mass [3]. Rodriguez et al. support this theory in their description of an unusual case of co-existing CAPNON and a low-grade cerebellar ependymoma associated with marked piloid gliosis on histology. The authors conclude that CAPNON is a result of tumefactive reaction rather than metaplasia [3]. The histological descriptions from our case are typical of those for CAPNON and are shown in Figure 2.

There is widespread acceptance that while surgical resection may provide symptom relief, histological diagnosis and prevention of recurrence, it does not come without risk. In our case there was significant postoperative right upper limb weakness as a result of infarction-postoperative MRI demonstrates an area of restricted diffusion extending postero-laterally from the surgical resection site to the left posterior parietal lobe.

Conclusion

CAPNON must be considered in the differential diagnosis of calcified lesions in the brain. The decision to proceed with surgical intervention must be influenced by the clinical presentation. The type of approach (e.g. biopsy versus total resection) should be discussed at a multidisciplinary level to mitigate the inherent risks of surgical intervention.

Conflicts of Interest

None declared
References

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