Involvement of Suboccipital Muscles in Plasmacytoma of Occipital Bone

Envolvimento suboccipital de músculos em plasmocitoma de osso occipital

Aravinth Kumar Ashok1  Mahendran Jolarpettai Venugopal1  Jothikumar Sethuraman1  Shankar Ranganathan1

1Institute of Neurosurgery, Madras Medical College, Chennai, Tamil Nadu, India

Address for correspondence Mahendran J. Venugopal, PhD, Institute of Neurosurgery, Madras Medical College, Chennai, Tamil Nadu, India (e-mail: mahendranvenugopal@gmail.com).

Abstract

Introduction  Suboccipital muscle involvement in a plasmacytoma of occipital bone is a rare clinical event. It leads to a dilemma of categorizing it into either Solitary plasmacytoma or extra medullary plasmacytoma. A working protocol is necessary to follow up these patients to detect their progression into multiple myeloma.

Clinical Presentation  A 38-year-old man presented with a progressively increasing hard swelling in the occipital region of five years duration. Brain CT revealed an osteolytic SOL over the torcula extending up to the foramen magnum. Brain MRI revealed an extraxial lesion. Brain MRV revealed no involvement of the torcula. Skeletal survey for metastasis was normal. After evaluation, we performed a surgical excision of the lesion. Intraoperatively, the tumor had invaded the suboccipital muscles and parietal bone and was adherent to the dura, without any intracranial extension. Histopathology reported Plasmacytoma. Immunohistochemical analysis was positive for CD-138. Bone marrow biopsy was negative for plasma cells. Bence Jones protein was absent and M protein absent in serum electrophoresis. USG abdomen normal. The patient underwent radiotherapy and is under continuous follow-up.

Conclusion  Plasmacytoma of occipital bone with suboccipital muscles involvement is categorized as Solitary Plasmacytoma of the bone. This provides convenience in diagnosis, management, and follow-up. We report on this case for its rarity in presentation. The protocol followed in our institute, based on our experience with such tumors, will aid in early identification of the progression of solitary plasmacytoma of bone into multiple myeloma and will augment prompt management.

Keywords  ► plasmacytoma  ► solitary  ► occipital bone  ► suboccipital muscle involvement  ► progression to multiple myeloma  ► follow-up protocol

Resumo

Justificativa e Importância  O envolvimento suboccipital de músculos em um plasmocitoma do osso occipital é um evento clínico raro. Isso leva a um dilema de categorizá-lo em qualquer plasmocitoma solitário ou plasmocitoma extramedular. Um protocolo de trabalho foi formulado para acompanhar esses pacientes para detectar a sua progressão para mieloma múltiplo.

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Introduction

Plasmacytoma of occipital bone with suboccipital muscles involvement is a rare clinical presentation. Such an entity could not be found in existing literature. Plasmacytoma of the head and neck is classified into two entities, namely, solitary plasmacytoma of bone and extramedullary plasmacytoma. The case in discussion does not fit the diagnostic criteria of either one. Suboccipital muscles involvement has rarely been encountered in cases reported to date. Hence, there is a dilemma in categorizing this case for the purpose of diagnosis and management. Moreover, such lesions have a high propensity to progress into multiple myeloma. We feel a longstanding need for a follow-up protocol to detect progression into multiple myeloma, which will help avoid any delays in further management. We report on this rare case for its unique presentation and to define the follow-up protocol designed and being followed at our institute.

Clinical Presentation

A 38-year-old gentleman came to our institute with complaints of progressive swelling in the occipital region of five years duration, headache of one year duration, and pain over the swelling of a year’s duration. The patient noticed a small swelling in the occipital region five years prior that was insidious in onset and grew progressively. The headache was mild, diffuse, on-and-off, more prevalent in the occipital region, and was not associated with symptoms of increased intra cranial pressure. Pain over the swelling was mild, on-and-off, and the swelling was not associated with trauma. There was no history of visual disturbance (Table 1) nor did the patient have any neurological deficit. Past medical history was not contributory to the current illness. There was no history of any radiation exposure. The patient was addicted to alcohol but had abstained from consuming alcohol for the past year. He was not a known smoker and was an agricultural laborer by occupation. History of any prolonged exposure to harmful chemicals was absent.

Upon examination, we saw a 10 × 10 × 6 cm swelling in the occipital region (Fig. 1). The consistency was variable with hard areas near the base and soft with firm areas near the apex. The skin over the swelling was not pulsatile. Serum calcium levels were normal.

Brain Computed Tomography (CT)

We found an 8 × 8 × 6 cm lytic lesion in the right occipital region extending to the left side. The lesion was extra-axial and was not infiltrating into the brain parenchyma. There was involvement of surrounding soft tissue. Mass effect over the cerebellum was present. We observed no evidence of hydrocephalus (Figs. 2 and 3).

Magnetic Resonance Imaging (MRI)

We observed an 8.7 × 8 × 5.6 cm moderately defined lesion in the right occipital skull bone. The lesion was hyperintense in both T1 and T2 images. Extension into pericranial soft tissue and

Table 1 Vision

<table>
<thead>
<tr>
<th>Test</th>
<th>Right</th>
<th>Left</th>
</tr>
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<tbody>
<tr>
<td>Visual acuity</td>
<td>6/6</td>
<td>6/6</td>
</tr>
<tr>
<td>Visual field</td>
<td>Normal</td>
<td>Normal</td>
</tr>
<tr>
<td>Color vision</td>
<td>Normal</td>
<td>Normal</td>
</tr>
<tr>
<td>Fundus</td>
<td>Normal</td>
<td>Normal</td>
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extradural destruction of skull bone were present. We also observed mass effect with compression of right cerebellum. There was no intraparenchymal extension observed. Intense contrast enhancement was present. Both transverse sinus and sigmoid sinus were normal. We saw no evidence of cerebral venous thrombosis (Figs. 4–7).

**Surgical Procedure**

With the patient under general anesthesia and in the prone position, we made an inverted U-shaped incision based on both the Transverse Sinuses. We raised the skin flap and identified the tumor mass infiltrating into the occipital bone. We also identified firm areas in the periphery and soft areas in the apex (Fig. 8). Tumor mass was adherent to the dura, which we carefully separated and cauterized. We found extension of lesion to the parietal bone and suboccipital muscle. Parietal bone was nibbled and

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**Fig. 1** Clinical preoperative photo of the tumor mass in the occipital region.

**Fig. 2** Computed tomogram picture of the Bone Window.
suboccipital muscles infiltration was excised. We achieved total excision of the lesion. After confirming hemostasis, the skin was closed in layers and dressing was applied (Fig. 9). We sent the specimens (bone, suboccipital muscles, and tumor mass) for HPE (histo-pathological examination) (Figs. 10–12).

Post-Operative Status

Post-operative period was uneventful. The wound remained healthy. The patient underwent suture removal on the seventh post-operative day. Post-operative brain CT showed complete excision of the lesion (Fig. 13).

Fig. 3 Computed tomogram picture of the Brain Parenchyma.
Fig. 4 T1-weighted magnetic resonance Image of the brain.

Fig. 5 T2-weighted magnetic resonance Image of the brain.

Fig. 6 Contrast magnetic resonance Image of the brain.

Fig. 7 Magnetic resonance Venogram.
Fig. 8  Intraoperative photo of the tumor, prior to excision.

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Fig. 9  Intraoperative photo of the tumor, post excision.

Fig. 10  Photo of excised bone infiltrated with tumor.

Fig. 11  Photo of excised tumor mass.
Histopathology showed densely cellular infiltrate of tumor cells invading bony trabeculae. The tumor cells were plasmacytoid in appearance. The cells had large round eccentric hyperchromatic nuclei and were also seen with double nuclei in eosinophilic cytoplasm (Fig. 14). The investigation

Fig. 12 Picture of tumor infiltrated suboccipital muscles.

Fig. 13 Postoperative computed tomography scan of the brain.

Fig. 14 Microscopic view of the plasma cells in the excised specimen.

**Histopathology**

Histopathology showed densely cellular infiltrate of tumor cells invading bony trabeculae. The tumor cells were plasmacytoid in appearance. The cells had large round eccentric hyperchromatic nuclei and were also seen with double nuclei in eosinophilic cytoplasm (Fig. 14). The investigation
demonstrated the presence of suboccipital muscle infiltration with plasma cell (Fig. 15).

**Immunohistochemistry**

CD-138 was positive (specific for plasma cell neoplasm) (Fig. 16).

**Other Investigations**

Peripheral smear was negative for plasma cells. Bone marrow biopsy was negative for plasma cells. Urine Bence Jones protein was also negative. Serum M protein was reported to be negative. The abdominal sonogram was also normal. Total Skeletal survey of long bones and pelvis were normal (Fig. 17).
**Treatment and Follow-up**

The patient has undergone radiotherapy and goes to follow-up once every three months. Routine examination of the patient’s Bence Jones protein and serum M protein are negative to date.

**References**