Anterior Clinoid Metastasis as First Presentation of a Signet Ring Cell Carcinoma: An Intriguing Diagnosis

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Introduction

The anterior clinoid process (ACP) may be affected by several pathological entities, being meningiomas the most common.¹⁻¹¹ Usually, patients harboring ACP lesions present with visual or oculomotor defects because of the involvement of optic nerve, superior orbital fissure, and/or cranial nerves in the cavernous sinus. Anterior clinoidectomy along with tumor removal is the treatment of choice, either performed intra- or extradurally. We herein report the case of a rare case of anterior clinoid metastasis from a silent signet ring cell carcinoma. The very unusual localization and the absence of any other sign of oncological disease made the diagnosis intriguing and difficult to achieve. The aim of this case report is to shed lights on the possibility of considering metastatic lesions among the differential diagnosis of pathologies involving the ACP.

Case Description

A 54-year-old female patient presented with right-sided visual disturbances due to optic nerve compression from a computed tomography (CT)-identified right anterior clinoid bone lesion. Contrast-enhanced magnetic resonance imaging showed an extra-axial, well-bordered enhancing mass extending from the right ACP toward the inner lumen of the optic canal. Pterional approach was adopted to remove the lesion and decompress the optic canal. Histological examination demonstrated a metastasis from a signet ring cell carcinoma. Postoperative CT showed near-total resection of the tumor and decompression of the optic canal. Visual defect remained unchanged.

Background

We report an extremely unusual case of anterior clinoid process (ACP) metastasis as the first presentation of a signet ring cell carcinoma.

Keywords

- anterior clinoid process
- metastasis
- signet ring cell carcinoma

Conclusion

Metastasis should be considered in the differential diagnosis of the ACP lesions. The early suspicion and identification of this extremely rare pathological entity can be helpful for the prompt management of patients, especially in the absence of any other signs of oncological diseases.
of 2/10 and visual field test revealed diffuse reduction of the retinal sensibility, both in the right eye. Contrast-enhanced magnetic resonance imaging showed an extra-axial, well-bordered enhancing mass, extending from the right ACP toward the inner lumen of the optic canal (Fig. 2).

Surgery was suggested to restore the visual function.

Fig. 1 Preoperative CT scans. Nonenhanced head CT scan, with axial (A) and coronal (B) reformats, demonstrates a lytic lesion of the right anterior clinoid process causing stenosis of the optic canal and marginal involvement of the superior orbital fissure. The lesion appears hyperdense with thin calcifications, representing “sequestra” of normal bone rather than dystrophic calcifications. CT, computed tomography.

Fig. 2 Preoperative MRI. Contrast-enhanced MRI: axial T2 weighted (A), coronal T2 weighted (B), axial T1 weighted (C), diffusion weighted (b 1,000 [D] and ADC map [E]), and contrast-enhanced VIBE (F) sequences. MR scan shows an extradural lesion with diffusion restriction (D, E), indicating high cellularity and enhancement. A mild linear dural thickening near the lesion can be also depicted (arrows on F). ADC, apparent diffusion coefficient; MRI, magnetic resonance imaging.
Surgical Technique
Right pterional craniotomy was performed and the lesser sphenoid wing was drilled away; the right ACP was reached extradurally and the lesion, which widely infiltrated the clinoid bone, was removed up to the achievement of a satisfactory decompression of the right optic canal—the right optic nerve and its vascularization were seen free of the pathology. Macroscopically, the tumor had reddish-gray soft appearance with calcifications in some regions and it presented a considerable vascularity. At the end of the removal, the dura mater was opened to confirm complete decompression of the optic nerve (►Fig. 3).

Pathology
Histological examination showed an infiltration of fibrous and bone tissue by a neoplasm arranged into nests and ribbons of medium-sized cells characterized by a broad cytoplasm containing large vacuoles full of mucin that displaced the nucleus to the cell’s periphery, the typical features observed in case of signet ring cell carcinoma (►Fig. 4). Immunohistochemical evaluation revealed positivity to cytokeratin 7, cytokeratin 20, cytokeratin AE1/3, epithelial membrane antigen (►Fig. 4), and vimentin. Focal reactivity was observed for CDX2. Progesterone receptor was negative. These findings were highly suggestive of a metastasis from an epithelial neoplasm that, in most cases, originates from the stomach. The major suspect was, therefore, for a gastric primitive, also because of the positive immunostaining to cytokeratin 7, cytokeratin 20, and focal to CDX2.

Follow-up
The postoperative CT showed near-total resection of the tumor and decompression of the optic canal (►Fig. 5). Visual defect remained unchanged. According to tumor board decision, the patient was immediately referred to oncological treatment for radiotherapy and chemotherapy.

Discussion
The present case exhibits three unusual features: the ACP as an extremely rare location for a metastasis, the origin of a suspected gastric cancer—one of the least frequent primary tumors causing intracranial metastases, and the occurrence of a skull base metastasis without any other sign or symptoms. To the best of our knowledge, the present case is the second report on anterior clinoid metastasis in the literature. In the previous case reported by Pojskić et al, the patient was under treatment with chemotherapy for a breast cancer. Upon the presence of an oncological disease, the occurrence of brain metastasis at the skull base and even at the ACP could be suspected. Albeit frank evidence of a primary disease, meningioma was suspected among the possible differential diagnosis.
In our case, visual defects were the only presenting signs of an undiagnosed, silent primary tumor; this has made the diagnosis of metastasis for the lesion of the ACP more troublesome. Neuroradiological findings allowed us to exclude common ACP lesions, such as meningiomas and mucoceles, favoring the hypothesis of an osseous neoplastic lesion. Indeed, extradural meningiomas are extremely rare, while mucoceles appear as expansile lesions with variable signal on T1- and T2-weighted images (depending on protein concentration) and are characterized by a peripheral linear enhancement. The differential diagnosis included both primary and metastatic bone lesions. Among primary ACP tumor, giant cell tumor has been seldom reported in this location, and therefore, it should be considered. Other pathologies that seldom may affect the ACP are pyocele, hemangiomas, hemangioblastoma, and dermoid cyst.

Pathology report disclosed a metastasis from a signet ring cell carcinoma, suggestive of a gastric primary tumor, although it has to be said that gastric cancer is one of the least frequent primary tumors causing intracranial metastases.

**Fig. 4 Histology.** The neoplasm was composed by nests of medium-sized cells, characterized by a cytoplasm with clear vacuoles that displace peripherally the nucleus (A, hematoxylin and eosin stain). Infiltration of bone tissue was evident (B, hematoxylin and eosin stain). Immunohistochemistry showed positivity to cytokeratin 7 (C), cytokeratin 20 (D), CDX2 (E) and EMA (F). All pictures were captured at ×40 magnification. EMA, epithelial membrane antigen.
Metastasis should be considered as possible differential diagnosis of ACP lesions, even in the absence of any sign of oncological disease. Safer and faster removal of ACP metastasis can be accomplished through extradural clinoidectomy. Albeit this pathological entity is extremely rare, the early suspicion supported by neuroradiological findings can be helpful for the prompt management of patients with such unusual clinical features as those described in the present case.

Conflict of Interest
None declared.

References


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