Fronto-orbito-ethmoidal intradiploic meningiomas: A case study with systematic review

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ABSTRACT

Background: Primary intradiploic meningiomas, extra-axial tumors arising primarily in the skull, are rare. The authors reported a complex case of intradiploic intraosseous metaplastic meningioma of the left medial wall and orbital roof with the left frontal sinus invasion and left ethmoidal bone substitution. The authors also conducted a systematic review concerning diagnosis and management of patients affected by purely calvarial intradiploic meningiomas along with a focus on fronto-orbito-ethmoidal ones.

Methods: A literature search was conducted using PubMed and Scopus databases according to preferred reporting items for systematic reviews and meta-analysis statement and with the following Mesh terms: Intradiploic, intraosseous, calvarial, and meningioma. Eligibility criteria were limited by the nature of existing literature on intradiploic meningiomas, consisting of only case series, and case reports.

Results: A total of 128 published studies were identified through our search. 41 studies were included in this systematic review, 59 patients with a female/male ratio of 1.2/1. The mean age of the patients is of 47.69 years (range 3–84 years). Only seven out of 59 patients (11.9%) presented a complex intradiploic meningioma located in fronto-orbito-ethmoidal region like our case. In almost all patients, a gross-total resection was performed (96.6%) and only in two patients (3.4%) a subtotal resection was achieved.

Conclusion: The authors shared this successfully treated case to add to the overall clinical experience in the management of this rare subtype tumor, with the hope that more studies are conducted to further address the mechanism of intradiploic meningiomas development.

Keywords: Anterior skull base, Calvarial, Intradiploic, Intraosseous, Meningioma

INTRODUCTION

Meningiomas are one of the most common benign brain/intracranial tumors with an incidence rate of 15–20%.[12] Parasagittal and convexity location are the most frequent accounting all together for almost 45% of the cases. The most common extradural localizations are the paranasal sinuses, nasal cavity, skin, neck, glands, and intraosseous space.[3] Primary intraosseous lesions
are rare accounting for <1% of intracranial meningiomas.\[39\] Intradiploic meningioma is an extremely rare type of extradural meningioma and is generally localized in the frontoparietal and orbital regions. Due to their low incidence and lack of adequate preoperative diagnostic imaging, intradiploic meningiomas are generally mistaken for primary calvarial bone tumors, en plaque meningiomas, and fibrous dysplasia. It is postulated that they could arise from arachnoid cell remnants captured at inappropriate sites during embryonic development or implantation of these cells by some mechanical insult such as trauma or dural tear.\[23\] Their origin can also be attributed to cellular dedifferentiation within the diploic space or cranial nerve sheath cells during their course through diploe.\[37\] These tumors are observed mostly within the first two decades of life and histologically are usually of psammomatous type. From a surgical perspective, when located onto the convexity surgical excision is feasible. Often, the complex reconstruction of the anterior skull base can represent a limit in the case of gross total resection (GTR) of large meningiomas involving also ethmoid and sphenoid bones.

**MATERIALS AND METHODS**

The authors collected clinical, radiological, intraoperative, postoperative, and histological features of a patient harboring a rare case of left intradiploic fronto-orbito-ethmoidal meningioma. We also performed a systematic review in accordance with preferred reporting items for systematic reviews and meta-analysis guidelines [Figure 1]. A literature search was conducted using PubMed and Scopus databases from 1955 (first reported paper) to 2021. References of the included studies were searched for additional relevant studies. This search was limited to studies in English language. MeSH terms utilized were as follows: (intradiploic OR intrasossue OR calvarial) AND meningioma. Eligibility criteria were limited by the nature of existing literature on intradiploic meningiomas, consisting of only case series and case reports. Data items collected included the following: authors and year of the study, number of patients, sex, signs and/or symptoms, intradiploic meningioma location, treatment modality, histological examination, and clinical outcome. In brief, 128 articles were found after database searching and removal of duplicates. After full-text review, 87 articles were excluded, leaving 41 included studies.

**CASE DESCRIPTION**

**History**

A 47-year-old female patient was admitted to our department, following the progressive onset of left eye proptosis lasting for 4 years associated with occasional mild to moderate headache. She did not complain visual acuity decrease nor diplopia in any gaze direction. At the time of her first medical consultation (at another Institution), she was advised to perform an endoscopic trans-nasal biopsy; this procedure was then performed but the histology was not helpful (no more information about this surgical procedure is available). After this, she decided to have a second opinion with the aim of definitively cure her illness.

**Physical examination**

On physical examination, there was axial proptosis on the left side, diplopia, with a restriction of the extraocular movements in the superior and medial side. The patient also presented hyposmia. She suffered from an amblyopia dating back to the adolescence in the contralateral eye (visual acuity: 7/10); in the left eye acuity was 8/10. No visual fields defect was detected, and the fundus oculi examination was normal. Rest of the neurological examination was normal. Routine hematological and biochemical tests were normal.

**Preoperative imaging**

A brain computerized tomography (CT) scan with thin cuts of the orbit and 3D-reconstruction showed a roughly ovoidal lesion centered at the left medial wall and roof of the orbit and invading the left frontal sinus and left ethmoidal body bone. The lesion’s maximal diameter was about 5 cm, and it was surrounded by a heterogeneous bone shell filled internally by hypodense tissue with multiple septations [Figure 2]. This lesion compressed and elevated the frontal lobe while, inferolaterally, the ocular bulb was dislocation with evident proptosis. Magnetic resonance imaging (MRI) showed that the mass was T1-weighted iso-intense and T2-weighted heterogeneously hyperintense with an intense postgadolinium enhancement. Medial and superior rectus and superior oblique muscles were compressed and dislocated [Figure 3]. A total body CT scan was negative for other neoplastic localization.

**Surgical procedures**

Neuronavigation was installed. A sterile surgical field was prepared at the level of the fascia lata to harvest a graft. A biconoral skin incision with subcutaneous two-layer dissection (to save a vascularized pericranial flap) was performed. A bilateral fronto-basal transfrontal sinus craniotomy was done to expose the tumor. Tumor was found to be completely extra-axial destroying the orbital roof and the orbital medial wall and eroding the ethmoid. Bilateral frontal sinus was invaded by the lesion (especially in the left side). The dura of the left anterior skull base was massively elevated together with the frontal lobe and totally substituted by a thin layer of fibrous tissue. The lesion presented a hard consistency due to an external hyperostotic bone shell. At the inside there was a cavity filled with numerous septations and blood clots. The periorbita was intact during the microsurgical
dissection with a good cleavage plan. The tumor was removed through a piecemeal technique. Peripheral hyperostotic bone was drilled away to obtain a good control of the bleeding and an efficient removal. Frontal skull base dura was removed and reconstructed with watertight technique with a synthetic patch. Left olfactory nerve was encased in the mass and was not possible to be spared. The tumor was completely removed obtaining a Simpson I resection. Frontal and ethmoidal sinus was closed by free temporal muscle fragments, gel foam, and fibrin glue. Then, orbital roof, medial orbital wall, and ethmoid were reconstructed using the split internal cortical bone of the frontal flaps, to avoid any postoperative pulsatile proptosis. This reconstruction was then covered with fascia lata and fibrin glue. Finally, the whole anterior skull-base until the planum sphenoidale was covered and sealed up with a vascularized pericranial layer. The bony defects of the frontal craniotomy were corrected using a methyl-methacrylate paste. A lumbar drain was placed to prevent any cerebrospinal fluid (CSF) leak (for a total of 8 days).

**Histological examination**

The histological analysis reported a metaplastic meningioma (the World Health Organization [WHO] Grade I) with...
extensive ossification with a bone angioectasia infiltration (immunostaining showed Epithelial Membrane Antigen +, Vimentine +, S100-, CKpool -, CD4-, Progesterone -) [Figure 4].

**Postoperative course and postoperative imaging**

At the recovery from the general anesthesia, the patient was drowsy and strongly apathetic. CT scan demonstrated a contusion and reactive edema at the basal frontal lobe. Intensive care stay was prolonged, and patient gradually recovered to a satisfying state of consciousness after eight days. Postoperative CT scan showed a total removal of the lesion, with an optimal reconstruction of the medial wall and the roof of the left orbit, in the absence of complications [Figure 5]. At the discharge, patient was neurologically intact and after 3 months got back to her job. Cosmetic outcome was particularly good, and the proptosis showed an evident reduction.

**Late rhinoliquorrhea management**

About 3 months from the previous CT scan control, the patient presented to the local emergency room for acute recurrence of rhinoliquorrhea. According to the COVID-19 re-organization model adopted by the Lombardy region, the patient was transferred to the neurosurgical reference hub in the area. When she arrived at the Neurosurgical department of the “Spedali Civili” Hospital in Brescia, the patient underwent a brain CT and MRI that showed the presence of pneumocephalus with a small defect in the posterior ethmoidal plane close to the spheno-ethmoidal suture with pneumocephalus [Figure 6]. After examining the radiological images, we decided to perform a combined transcranial and transsphenoidal approach. Such approach, in our experience, yields to a higher successful rate. Under balanced general anesthesia, the previous bicornal surgical incision was opened. After removing the bone flaps, no evident extradural CSF collection was found. The bone flaps and all the reconstruction materials (i.e., methyl methacrylate and orbital roof titanium mash) were removed and after meticulous exploration a defect at the margin of the duroplasty with the dura mater was found. The synthetic patch was completely removed, and the underlying subdural space was filled with a graft of abdominal fat tissue (paraumbilical incision). The dura was reconstructed with a free pericranium (taken from the calvarium posterior to the craniotomy) with a watertight technique and reinforced with fibrin glue. The reconstruction of the anterior skull base appeared intact except in its posterior ethmoidal portion. A 5 mm gap/defect was found at the level of the left ethmoido-sphenoidal suture. The adjacent pericranium was detached to adequately expose the defect and the surrounding bone. The posterior ethmoidal space, after thorough cleaning, was filled with free fatty tissue (till the intact periorbital plane).
laterally and medially to the mucosal graft and sealed with one layer of free pericranium. The margins of pericranial layer (positioned in the previous intervention) dehiscent have been incised and removed. The margins were slightly detached from the bone plane for about 3 mm; a first patch of free pericranium was placed, in contact with the free bone margin and to cover the fat layer. The pedicled pericranium was then repositioned to cover the whole floor of the left anterior cranial fossa and a second patch of free pericranium was applied to it by superimposing it on the vital pericranium (with multilayer wafer technique). The whole construct was then sealed with fibrin glue. The epidural space was filled with free fatty abdominal tissue. Through an endoscopic transnasal approach the free abdominal flap fat in the sphen-ethmoidal suture is easily identified. The nasal mucosa bordering the defect was removed and vital bone tissue exposed. A pedicled naso-septal flap, previously harvested, was then used to further seal and support the reconstruction.
Silicone coated pads were positioned to make the flap adhere better to hold the construct (the endonasal swabs will then be removed 4 days after surgery). The frontal bone operculum was positioned and fixed with plates and screw (Synthes Matrix). Postoperative course was uneventful. A control head CT scan showed good surgical results and resolution of the pneumocephalus. An endoscopic exploration showed good vitality and adhesion of the mucosal flap. After prophylactic therapy (vancomycin + cephalosporin for 10 days), the patient was discharged at home neurologically intact. The 3-month postoperative MRI showed no residual or recurrent disease [Figure 7].

RESULTS
A total of 128 published studies were identified through our detailed searches from 1955 to 2021. After a detailed examination of the titles, abstracts, and contents of these studies, 87 were rejected from our systematic review because they were not in English language, not purely intradiploic calvarial meningiomas, or did not include accurate data. Our case was not included in this systematic review. A summary of patients’ characteristics in selected papers are reported in [Table 1].

For about two-thirds of extradural meningioma. According to the above-mentioned classification, our case can be classified as Type 2. C. Cirak classified primary extradural meningioma (PEM) into three types: Type 1 (purely extracalvarial), Type 2 (purely calvarial), and Type 3 (calvarial with extracalvarial extension). Types 2 and 3 can be subclassified as B (convexity) and C (skull base). According to the above-mentioned classification, our case can be classified as Type 2 C. Cirak et al. reported that psammomatous variety is the most common histological subtype, but the results of our systematic review showed that the most frequent histology is the meningothelial subtype. At our knowledge, the current is the first case of intradiploic metaplastic meningioma. There are different theories as to the origin of intradiploic meningiomas. On the one hand, there is the concept that the meningiomas evolve from the terminal vessels of the diploic space; on the other hand, they are derived from the meninges enveloping the brain, especially at the level of the anterior cranial fossa or of the sphenoid lesser wing. Winkler in 1904 first described a meningioma originating from extradural location and extradural meningioma arising in the skull is usually referred as calvarial, intraosseous, or intradiploic location. Intraosseous meningioma denotes a subset of extradural meningioma arising in bone and accounts for about two-thirds of extradural meningioma. According to the literature, frontoparietal and orbital regions are the most common intraosseous meningioma locations. Lang et al. classified primary extradural meningioma (PEM) into three types: Type 1 (purely extracalvarial), Type 2 (purely calvarial), and Type 3 (calvarial with extracalvarial extension). Types 2 and 3 can be subclassified as B (convexity) and C (skull base). According to the above-mentioned classification, our case can be classified as Type 2 C. Cirak et al. reported that psammomatous variety is the most common histological subtype, but the results of our systematic review showed that the most frequent histology is the meningothelial subtype. At our knowledge, the current is the first case of intradiploic metaplastic meningioma. There are different theories as to the origin of intradiploic meningiomas. On the one

DISCUSSION
A localization of meningiomatous tissue at the level of the orbita, it is most often due to secondary invasion of the orbit or of the orbital walls by a tumor starting from the meninges enveloping the brain, especially at the level of the anterior cranial fossa or of the sphenoid lesser wing. Winkler in 1904 first described a meningioma originating from extradural location and extradural meningioma arising in the skull is usually referred as calvarial, intraosseous, or intradiploic location. Intraosseous meningioma denotes a subset of extradural meningioma arising in bone and accounts for about two-thirds of extradural meningioma. According to the literature, frontoparietal and orbital regions are the most common intraosseous meningioma locations. Lang et al. classified primary extradural meningioma (PEM) into three types: Type 1 (purely extracalvarial), Type 2 (purely calvarial), and Type 3 (calvarial with extracalvarial extension). Types 2 and 3 can be subclassified as B (convexity) and C (skull base). According to the above-mentioned classification, our case can be classified as Type 2 C. Cirak et al. reported that psammomatous variety is the most common histological subtype, but the results of our systematic review showed that the most frequent histology is the meningothelial subtype. At our knowledge, the current is the first case of intradiploic metaplastic meningioma. There are different theories as to the origin of intradiploic meningiomas. On the one
Table 1: Schematic summary of the characteristics of 59 patients affected by intradiploic meningioma, in the context of our systematic review.

| Authors and year | Age (Years) | Sex | Signs and/or symptoms | Location | Treatment | Histopathological evaluation | Outcome |
|------------------|-------------|-----|------------------------|----------|-----------|-------------------------------|---------|
| Reale et al. 1978 | 19          | Male | Exophthalmos           | Medial and superior wall right orbit | GTR (right frontal flap) | Endotheliomatous and psammomatous meningioma | Good recovery |
| Pompili et al. 1983 | 40          | Male | Exophthalmos           | Upper part left orbita               | GTR (extradural subfrontal approach) | Psammomatous meningioma | Good recovery |
| Oka et al. 1989   | 79          | Female | Asymptomatic          | Vertex                               | GTR (skin flap over the vertex) | Transitional meningioma | Good recovery |
| Reale et al. 1978 | 19          | Male | Exophthalmos           | Medial and superior wall right orbit | GTR (right frontal flap) | Endotheliomatous and psammomatous meningioma | Good recovery |
| Pompili et al. 1983 | 40          | Male | Exophthalmos           | Upper part left orbita               | GTR (extradural subfrontal approach) | Psammomatous meningioma | Good recovery |
| Oka et al. 1989   | 79          | Female | Asymptomatic          | Vertex                               | GTR (bilateral parietal craniotomy) | Transitional meningioma | Good recovery |
| Van Tassel et al. 1991 | 41      | Female | Asymptomatic          | Right sphenoid bone                  | GTR | Meningothelial meningioma      | Not reported |
| Kulali et al. 1991 | 50          | Male | Asymptomatic           | Right occipital                      | GTR | Meningothelial meningioma      | Good recovery |
| Halpin et al. 1991 | 19          | Male | Exophthalmos           | Body and wings of the sphenoid bone  | GTR (bicoronal scalp flap) | Transitional meningioma | Good recovery |
| Ghobashy et al. 1994 | 65         | Female | Generalized dull headaches | Right frontal | GTR (bicoronal scalp flap) | Transitional meningioma | Good recovery |
| Prado et al. 1994  | not reported | Male | Asymptomatic           | Frontal                              | GTR | Meningothelial meningioma      | Good recovery |
| Monteiro et al. 1996 | 30          | Male | lump slightly painful  | Left parietal                        | GTR (left parietal skin flap) | Psammomatous meningioma | Good recovery |
| Muthukumar et al. 1997 | 55    | Male | Asymptomatic           | Right parietal                       | GTR | Meningothelial meningioma      | Good recovery |
| Anegawa et al. 1999 | 77          | Female | Transient episodes of left-sided weakness, vomiting, left-sided ptosis, lethargy | Bilateral parietal | GTR (external decompressive craniectomy) | Meningothelial meningioma | Good recovery |

(Contd...)
Table 1: (Continued).

| Authors and year | Age (Years) | Sex | Signs and/or symptoms | Location | Treatment | Histopathological evaluation | Outcome |
|------------------|-------------|-----|------------------------|----------|-----------|-----------------------------|---------|
| Cirak et al. 2000⁷ | 12          | Female | Exophthalmos, occasional diplopia | Fronto-orbitonasal | GTR (bicornal skin incision) | Psammomatous meningioma | Only exophthalmos |
| Lang et al. 2000²¹ | 41          | Female | Headache | Right greater sphenoid wing | GTR | Purely calvarial meningioma | Good recovery |
|                  | 49          | Female | Asymptomatic | Left frontal wing; superior & lateral orbital wall; left greater sphenoid wing | GTR | Purely calvarial meningioma | Good recovery |
| Muzumdar et al. 2001³⁶ | 63          | Female | Headache and exophthalmos, chemosis left eye | Both frontoparietal convexity and superolateral aspect of the left orbit | GTR (left frontoparietal craniotomy) | Meningothelial meningioma | Good recovery |
| Desai et al. 2004¹⁰ | 12          | Male | Exophthalmos | Left orbital roof | GTR (basal subfrontal craniotomy) | Psammomatous meningioma | Good recovery |
| Al-khawaja et al. 2007³⁷ | 50         | Male | Headache | Left parasagittal | GTR | Meningothelial meningioma | Good recovery |
| Bou-Assaly et al. 2007⁵ | not reported | Male | Not reported | Progressive vision loss and significant Exophthalmus | GTR (left fronto-temporal craniectomy) | Meningothelial meningioma | Good recovery |
| Borkar et al. 2008⁴ | 40          | Male | Asymptomatic | Exophthalmos | GTR (bifrontal craniotomy) | Transitional meningioma | Good recovery |
| Iannelli et al. 2008¹⁶ | 3           | Female | Asymptomatic | Left parietal | GTR | Transitional meningioma | Good recovery |
| Sujit Kumar et al. 2009²⁰ | 46          | Female | Not reported | Parasagittal | STR | Malignant meningioma | Wound infection with CSF leak |
| Yener et al. 2009⁴⁰ | 78          | Male | Asymptomatic | Right parietal | GTR | Meningothelial meningioma | Good recovery |
| Sambasivan et al. 2010³² | 13          | Male | Asymptomatic | Right temporal | GTR | Intradiploic meningioma | Good recovery |
| Mukherjee et al. 2010³⁴ | 55          | Male | Headache, exophthalmos | Large calvarial mass crossing the midline and with extension To the right orbit | GTR (bicornal skin incision) | Atypical meningioma | Reoperated Twice for small recurrences at bony edges. |
| Hong et al. 2010¹⁵ | 52          | Male | Asymptomatic | Parietal | GTR | Intraosseus meningioma | Good recovery |
|                  | 73          | Male | Asymptomatic | Occipital | GTR | Anaplastic meningioma | Good recovery |
| Yilmaz et al. 2010⁴¹ | 41          | Male | Headache | Frontal | GTR (right frontotemporal craniectomy) | Meningothelial meningioma | Not reported |
| Eras et al. 2011¹¹ | 16          | Female | Asymptomatic | Right parietal and posterior frontal | GTR (right frontoparietal craniectomy) | Transitional meningioma | Good recovery |
| Khalatbari et al. 2011³⁸ | 14          | Female | Exophthalmos, diplopia | Right orbital roof | GTR (bicornal skin incision) | Transitional meningioma | Good recovery |

(Contd...)
| Authors and year | Age (Years) | Sex | Signs and/or symptoms | Location | Treatment | Histopathological evaluation | Outcome |
|------------------|-------------|-----|------------------------|----------|-----------|-----------------------------|---------|
| Cruz et al. 2013<sup>[9]</sup> | 43          | Female | Ptosis | Right orbital roof and later portion of the frontal | Gtr (coronal approach) | Meningothelial meningioma | Not reported |
| Kariyattil et al. 2014<sup>[17]</sup> | 40          | Female | Headache and frontal swelling | Left frontal | GTR (left frontal craniotomy) | Meningothelial meningioma | Good recovery |
| Yun et al. 2014<sup>[4]</sup> | 65          | Female | Asymptomatic | Right frontal | GTR | Atypical meningioma | Good recovery |
| Velazquez Vega et al. 2015<sup>[36]</sup> | 60          | Male | Asymptomatic | Right frontal | GTR | Microcystic meningioma | Good recovery |
| 62 Male | Asymptomatic | Left frontal | GTR | Microcystic meningioma | Good recovery |
| 56 Female | Previous right optic Nerve meningioma and Radiation Asymptomatic | Right frontal | GTR | Microcystic meningioma | Good recovery |
| 68 Female | Asymptomatic | Right parietal | GTR | Microcystic meningioma | Good recovery |
| 71 Female | Asymptomatic | Right frontal | GTR | Microcystic meningioma | Good recovery |
| 71 Female | Diplopia | Left parietal | GTR | Microcystic meningioma | Good recovery |
| 79 Female | Asymptomatic | Left parietal | GTR | Microcystic meningioma | Good recovery |
| 79 Female | Diplopia, exophthalmos, intermittent Ptosis and visual field Defects | Right sphenoid bone | GTR | Microcystic meningioma | Good recovery |
| 84 Female | Asymptomatic | Left frontal | GTR | Microcystic meningioma | Good recovery |
| Vital et al. 2015<sup>[38]</sup> | 62          | Female | Headache | Right parietal | GTR (right parietal incision and craniotomy) | Meningothelial meningioma | Good recovery |
| 64 Male | Pain And swelling on the right side of head | Right frontotemporal | GTR | Intradiploic meningioma | Good recovery |
| 16 Female | Exophthalmos, headache | Right orbital roof | GTR (right fronto-temporal craniotomy with orbitotomy) | Intradiploic meningioma | Good recovery |
| 63 Female | Pain in the right eye | Right frontal | STR | Fibroblastic meningioma | Good recovery |
| 62 Male | Exophthalmos | Lateral wall of the orbit and the greater wing of the sphenoid | GTR (left frontopterional approach) | Meningothelial meningioma | Good recovery |

(Contd...)
hand, it is thought that they arise from arachnoid cell rests inappropriately laid down during embryonal life or deposited in unusual sites by trauma and dural tears. On the other hand, they may arise from the sheath cell of cranial nerves as they course through the diploic space or their origin may be due to cellular dedifferentiation within the diploe; a similar mechanism is proposed for the equally uncommon cutaneous meningioma.\textsuperscript{7} The common age of presentation was in the early second decade of life and although intraosseous meningiomas commonly involve the calvarial bone, they rarely occur in the orbital roof and sphenoid bone.\textsuperscript{1,10,11} Intradiploic meningiomas of the orbital roof are more frequent in males and in young people.\textsuperscript{19} These tumors may be considered as a subgroup of intraosseous meningiomas and it does not fit any of the intraorbital meningiomas described by Craig and Gogela.\textsuperscript{8} PEM grows slowly over time and produces signs and symptoms due to compression of adjacent neural structures, especially exophthalmos is the predominant symptom.\textsuperscript{122} In addition, intraosseous PEMs appear on tomography scans as expansile masses or provoking osteolytic lesions that mimic metastatic disease.\textsuperscript{9} X-ray, head CT, and brain MRI are methods of radiological screening in diagnosing intradiploic meningioma. Radiologically intradiploic meningiomas are typically either osteoplastic or osteolytic, and although the majority are osteoblastic, extremely rarely are osteolytic or very extremely have a mixed pattern. These can present with opacity of the orbital roof on X-ray skull study. The osteolytic subtype of intradiploic meningiomas is more likely to be malignant than the osteoblastic subtype. Intradiploic meningiomas should be considered in the differential diagnosis of patients presenting with osteoblastic or osteolytic skull lesions.\textsuperscript{107} The differential diagnosis of the osteoclastic lesion should include chondroma, chondrosarcoma, hemangioma, epidermal cyst, eosinophilic granuloma, metastasis, and fibrous dysplasia, whereas the radiological differential diagnosis of osteoblastic lesions should include metabolic diseases including osteoma, hyperparathyroidism, metastasis and A and D hypervitaminosis. Head CT scan can be used

| Authors and year | Age (Years) | Sex | Signs and/or symptoms | Location | Treatment | Histopathological evaluation | Outcome |
|------------------|-------------|-----|-----------------------|----------|-----------|-------------------------------|---------|
| Zhang et al. 2017\textsuperscript{43} | 48 | Male | Headache | Frontal | GTR (bilateral frontal craniotomy) | Intraosseus meningioma | Good recovery |
| Sundblom et al. 2018\textsuperscript{34} | 39 | Female | Headache, bilateral anosmia, bilateral papilledema, diminished visual fields bilaterally, headache, exophthalmos, visual blurring in her left eye | Left temporal | GTR (left frontotemporal orbitozygomatic craniotomy) | Intradiploic meningioma | Good recovery |
| Mankotia et al. 2018\textsuperscript{22} | 27 | Female | Headache, exophthalmos, visual blurring in her left eye | Occipital | GTR (midline suboccipital craniectomy) | Meningothelial meningioma | Persisting minimal gait disturbance |
| Satyarthee 2018\textsuperscript{32} | 40 | Male | Headache, gait imbalance, bilateral papilledema exophthalmos, diplopia, headache, hyposmia | Left fronto-orbital and left ethmoidal | GTR (bicornoral skin incision and bilateral fronto-basal craniotomy with orbital roof and medial wall orbital reconstruction | Metaplastic | Good recovery |
| Present Case | 47 | Female | Exophthalmos, diplopia, headache, hyposmia | Occipital | GTR (bicornoral skin incision and bilateral fronto-basal craniotomy with orbital roof and medial wall orbital reconstruction) | Metaplastic | Good recovery |

GTR: Gross Total Resection, STR: Sub-total resection
to best define the margins of meningiomas in the bone algorithm and brain MRI can be used to identify adherent tissue involvement. Particularly, MRI is helpful to evaluate the extraosseous extension and the soft tissue component. It is like the primary intradural meningiomas except for the dural tail sign, which could not be seen in intradiploic meningiomas. Furthermore, scintigraphy and positron emission tomography CT (PET-CT) can be used for the differential diagnosis, with bone scintigraphy that shows an increased metabolic activity of the specific bone region when positive. Tumor excision with wide surgical resection and meticulous bony reconstruction is the primary choice for treatment in symptomatic patients and in this kind of tumor surgery, if the shell of the tumor facing the dura mater is not intact, the most important rule is to prevent postoperative CSF leakage. This holds especially true for our case, given that the large dimension and the infiltration of multiple compartments (namely the roof and medial wall of the orbit, the ethmoid, and the frontal sinus) set a particularly high risk of CSF fistula. More, the absence of a dura sheet between the tumor and the basal frontal brain provoked a superficial damage to fronto-basal cortex. Nevertheless, these factors did not prevent us to accomplish a Simpson I resection and avoid CSF leak. Although most intradiploic meningiomas are benign, they could present malignant growth, which might lead to erosion of the skull. Therefore, in most cases, total resection of lesions seems to be impossible, and the residual tumor should be treated with radiation therapy and chemotherapy, especially for the patients who had malignant or atypical features on histological examination. The present chemotherapeutic agents mainly include molecule-targeted drugs (imatinib, sunitinib, and vatalanib), cytotoxic chemotherapeutics (hydroxyurea, irinotecan, and temozolomide), hormonal inhibitors (targeting estrogen and progesterone), and other drugs. Hence, postoperative imaging will become vital for monitoring tumor recurrence or progress and prolonged follow-up of patients will be necessary.

CONCLUSION

Primary intradiploic meningiomas are rare tumors, which are used to describe the subset of extradural meningiomas that arise in the skull. To the best of our knowledge, only seven cases of complex intradiploic fronto-orbito-ethmoidal meningioma were reported in the literature. Regular monitoring is essential in the management of intradiploic meningioma and surgical treatment with wide incision of the lesion is the goal of the primary therapy. We share this successfully treated case to add to the overall clinical experience in the management of this rare subtype tumor, with the hope that more studies are conducted to further address the mechanism of intradiploic meningiomas development.

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Ethical approval

All procedures in studies involving human participants were in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards.

Declaration of patient consent

The authors certify that they have obtained all appropriate patient consent.

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Conflicts of interest

There are no conflicts of interest.

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