Unilateral proptosis as a clinical presentation in primary angiosarcoma of skull

SAMRUAY SHUANGSHOTI, PRACHA CHAYAPUM, NITAYA SUWANWELA, AND CHARUS SUWANWELA

From the Departments of Pathology, Surgery, and Radiology, Faculty of Medicine, Chulalongkorn University, Bangkok, Thailand

SUMMARY A case is reported of unilateral proptosis of the left eyeball as a unique clinical presentation of primary angiosarcoma of the skull involving the sphenoid and frontal bones and roof of the orbit on the left side of a 32-year-old man. The intraosseous, extradural non-encapsulated, and friable tumour contained cytoplasmic factor VIII-related antigen. After complete removal of the tumour the patient has been well for 10 months of the follow-up period. This case brings the total number of reported primary angiosarcomas of the cranium to nine. All patients were male and most of them were young. The average age was 24 years. The frontal bone seems to be the site of predilection for cranial angiosarcomas. Surgical extirpation, radiation, or combined surgical removal and radiation have been used for treating angiosarcoma of the skull with variable results. It is suggested that angiosarcoma of the skull has a worse prognosis than angiosarcoma of bones other than the cranium because the patient may die of secondary neoplastic involvement of the brain.

Primary new growths of the cranial bones constitute only 2-6% of 500 primary bone tumours in one series1 and 2.4% of 1610 primary bone neoplasms in another collection.2 Angiosarcomas are extremely rare among primary malignant tumours of the skull; only eight acceptable cases have appeared in the literature.3-8 In this communication we report an intraosseous angiosarcoma involving the sphenoid and frontal bones as well as the roof of the orbit and causing unilateral exophthalmos. The latter becomes a unique clinical presentation for angiosarcoma arising in the cranium.

Case report

A 32-year-old man was admitted to hospital because of progressive left proptosis without other associated symptoms or signs for one year.

EXAMINATION

With a slit-lamp the left eye was 15 mm compared with 12 mm for the right eye by measuring from the orbital rim to the most anterior aspect of the cornea. Ocular movements were normal bilaterally. Each pupil, 2 mm in diameter, was reactive to light. Visual acuity, visual field, eye ground, and the rest of the physical examination were unremarkable.

A plain x-ray of the skull showed destruction of the lesser wing of the sphenoid bone and posterior wall of the orbit on the left side. Computed tomographic (CT) scans of the cranium gave the following findings on the left side: proptosis, and destruction and expansion of the sphenoid and frontal bones in relation to an enhanced lesion which appeared to extend into the orbit (Figs. 1A, B).

OPERATION

A left frontotemporal craniotomy disclosed an intraosseous, extradural, non-encapsulated, highly vascular, and friable tumour involving the lesser and greater wings of the sphenoid bone, frontal bone, and roof of the orbit. It was firmly adherent to the adjacent dura mater. A small part of the lesion was also adherent to the superior aspect of the bulbar fascia of the left eyeball. The entire neoplasm was removed along with the affected parts of bones,
bulbar fascia, and dura mater. The dural defect was
grafted with temporal fascia. There was profuse
bleeding, which required 10 units of blood for
transfusion. The adjacent part of the brain was free
from tumour infiltration. The patient recovered
uneventfully.

PATHOLOGICAL EXAMINATION
Multiple pieces of haemorrhagic tissue with frag-
ments of bone, about 2.5 ml in combined volume,
were fixed in 10% formalin. The largest fragment of
tumour, 2-5 cm in greatest dimension, was attached
partly to dura mater; it had a spongy cut surface.
These tissue fragments were embedded in paraffin
and stained with haematoxylin and eosin (H-E).
Gomori’s silver impregnation for reticulin fibres was
also done. Moreover, the peroxidase-antiperoxidase
(PAP) indirect immunohistochemical method was
used on paraffin-embedded tissue sections for locali-
sation of factor VIII-related antigen, a protein
synthesised primarily by endothelium, in the
tumour cells. A fragment of the spleen was similarly
processed as a control.

Microscopically the lesion was a highly vascular
neoplasm invading bone and dura mater (Fig. 2).
Neoplastic blood vessels were lined by plump cuboid-
to-ovoid endothelial cells having pleomorphic and
hyperchromatic nuclei. Scattered individual vessels,
anastomosing vascular channels, and papillary
formations were noted (Fig. 3). Nevertheless large
blood-filled spaces lined by endothelial cells that
were not strikingly atypical were also observed. The
stroma between recognisable blood vessels was com-
posed of loose myxomatous tissue that contained
stellate cells as well as elongate or spindle-shaped
cells (Fig. 4A) and neoplastic cells similar to those
lined the vascular channels. These extravascular
tumour cells tended to be in clusters or disseminated
individually (Fig. 4B). Gomori’s silver impregnation
disclosed that the tumour cells were within the
reticulin sheath of the vessels in the area where
vascular spaces were present (Fig. 4C). Reticulin
fibres also surrounded clusters of the neoplastic
cells in the zone where the tumour cells were grouped
within the myxomatous matrix (Fig. 4D). Encircling
of the argyrophilic fibres round the individual tumour
cells was not seen. Factor VIII-associated antigen
was often present in the cytoplasm of the tumour cells
that lined the vascular lumina and in the myxomatous
stroma (Fig. 5). Hence the endothelial nature of the
tumour cells was confirmed immunohistochemically.

The pathological diagnosis was an angiosarcoma
involving the sphenoid and frontal bones and roof of
the orbit on the left side and invading the adjacent
dura mater and superior aspect of the bulbar fascia of
the left eyeball.

POSTOPERATIVE COURSE
The patient was discharged 17 days after admission to
hospital. CT scans of the skull one month and nine
months after surgical intervention revealed no
evidence of tumour. There was a decrease in the size

Fig. 1  CT scan of skull. A: Plain CT scan at the base of the cranium shows destruction and expansion of the lesser wing of the
sphenoid and frontal bones on the left side with involvement of the diploic space. B: The enhanced mass extends into the
posterolateral part of the left orbit.
Unilateral proptosis as a clinical presentation in primary angiosarcoma of skull

Discussion

GENERAL CONSIDERATIONS

The term ‘angiosarcoma’ as used here is regarded as synonymous with the following terms found in the literature: angioendothelioma, malignant angioma, angiofibrosarcoma, haemangioblastoma, angioblastic sarcoma, malignant haemangioendothelioma, haemangioendothelioblastoma, malignant angioblastoma, haemangioendotheliosarcoma, intra-vascular endothelioma, malignant endothelioma, and malignant bone aneurysm.13 14

Anastomosing vascular channels lined by pleomorphic endothelial cells which often form papillae are diagnostic of angiosarcoma.14 These histopathological features were also seen in our patient’s tumour. Moreover, the endothelial nature of the tumour cells is further confirmed by localisation of cytoplasmic factor VIII-related antigen. Hence the pathological diagnosis of the angiosarcoma is considered as unequivocal in our case.

The main intraosseous position with minimal infiltration of the bulbar fascia, as noted surgically, makes us conclude that the current angiosarcoma is primarily intraosseous, with secondary invasion into the superior aspect of the bulbar fascia. It is hard to imagine reciprocally that a small primary angiosarcoma of the latter will extend widely to become an intraosseous lesion within the sphenoid and frontal bones as well as the orbital roof. A lesion of bones forming the orbit may cause proptosis.

of the shadow of soft tissues at the operative site. The left exophthalmos was improved. The patient could perform his daily work as usual.

Fig. 2 Invasion of angiosarcoma into bone and dura mater. A: The angiosarcoma invades bone trabecula (upper arrowhead) and dura mater (lower arrowheads) (H-E, ×44). B: The area indicated by the upper arrowhead in A demonstrates focal invasion of the tumour cells into the edge of the bone trabecula (H-E, ×348). C: The region indicated by the left lower arrowhead in A exhibits infiltration of the neoplastic cells into the dura mater (H-E, ×348).
Differential Diagnosis
The present angiosarcoma of bones must be distinguished from such lesions as meningioma, haemangiopericytoma, and primary intravascular papillary endothelial hyperplasia. Attachment to the dura mater, bony invasion, and papillary formations suggest that the tumour is a meningioma, especially the papillary type. However, the absence of the tumour capsule, meningocytes, cellular whorls, and intranuclear vacuoles, and the presence of cytoplasmic factor VIII-related antigen in the tumour cells exclude a meningioma. We also do not consider our patient’s tumour to be a haemangiopericytoma of the meninges or of the orbit. The absence of reticulin fibres round individual tumour cells is not a feature of haemangiopericytoma. Moreover, the neoplastic cells were mainly intravascular rather than perivascular. Hence they were not pericytes. According to Nadji et al., tests for factor VIII-associated antigen are persistently negative in cells of haemangiopericytoma.

Primary intravascular papillary endothelial hyperplasia may mimic an angiosarcoma. However, the endothelium in this condition is benign, as the name ‘hyperplasia’ implies. Although the overgrown endothelium in this condition may form intravascular papillae, it lacks nuclear pleomorphism. By contrast, the current lesion had endothelium that showed pleomorphic nuclei and invaded the surrounding tissues. Hence the lesion should be named as angiosarcoma rather than as primary intravascular papillary endothelial hyperplasia.

Brief Review of Literature
To our knowledge 12 angiosarcomas of the skull have been reported. However, four of them are
Unilateral proptosis as a clinical presentation in primary angiosarcoma of skull

... excluded herewith because of inadequate pathological confirmation of the lesions. Thus nine angiosarcomas of the cranium including our case are considered in this discussion. The youngest patient was 3 months old and the oldest was 50 years. Most patients were young. Three were children less than 8 years old. The average age was 24 years. All persons were male. A local mass or swelling was the usual clinical presentation. Only our patient presented with unilateral proptosis.

The tumour involved the frontal bone in four cases, parietal bone in two cases, temporal bone in one case, sphenoid and frontal bones as well as the orbital roof in one case (current patient), and maxilla in one case. Thus the frontal bone seems to be the favoured site for angiosarcomas of the skull.

Resection, radiation, or combined resection and radiation were frequently used in treating the angiosarcomas of the cranium, with variable results. Surgical treatment seems to give encouraging results in some patients, in that they have been alive for 27 months and 3½ years, but one patient died post-

Fig. 4 Stroma and pattern of reticulin fibres in angiosarcoma. A: Stellate and spindle-shaped cells lie within the loose myxomatous matrix of the tumour (H-E, ×886). B: The tumour cells are clustered in the loose myxomatous stroma (H-E, ×354). C: Reticulin fibres surround blood vessels that are lined by neoplastic endothelium. Some blood vessels are partly filled by endothelial cells (Gomori's silver impregnation, ×354). D: Clusters of the tumour cells in the myxomatous matrix are surrounded by a few argyrophilic fibres (Gomori's silver impregnation, ×354).
operatively.\(^8\) A boy who received high doses of radiation for primary and recurrent angiosarcomas of the frontal bones has been well for 12 years,\(^4\) but another boy lived only 9-5 months with radiotherapy.\(^3\) Chow et al.\(^7\) believed that a 21-year-old man who had had an angiosarcoma of the frontal bone was cured by combined radiation and resection. The patient first had a biopsy and a course of radiotherapy totalling 6000 cGy over a six-week period. Seven months later he underwent resection, when the tumour site including the surrounding bone and dura mater was removed en bloc. Pathologically, only fibrosis and suture granuloma were seen.\(^7\) Nevertheless we believe that angiosarcoma of the skull may have a worse prognosis than angiosarcoma of bones other than the cranium, because the tumour may secondarily involve the brain and the patient dies with increased intracranial pressure, as did the boy reported on by Kinkade.\(^3\)

Professor Samruay Shuangshoti is in receipt of a grant from the Medical Section, Anandhamahidol Foundation, 1985-8, Bangkok, Thailand.

References

Unilateral proptosis as a clinical presentation in primary angiosarcoma of skull