Delayed spontaneous extradural hematoma following bilateral decompressive craniectomy

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Abstract

Development of symptomatic extradural hematomas, in the absence of bone flap is quite uncommon. We herein report a case of spontaneous extradural haematoma following bilateral decompressive craniectomy for severe head injury with left frontal contusion and right frontal acute SDH. Traumatic brain injury (especially following parenchymal loss) is a dynamic process, with continual changes occurring till the normalisation of the internal milieu of the cranium, which should be planned earliest after normalisation of intracranial pressure.

Key words: Extradural hematoma, decompressive craniectomy, traumatic brain injury

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Introduction

The dynamic changes occurring in the brain are not just limited to the acute phase but do run into the postoperative period till the internal milieu of the intracranial compartment is normalized. Enlargement of small extradural hematomas, fracture hematomas and de novo development of hematomas on the contralateral side in the immediate post operative period have been well described in the literature with the reported incidence of 5.7%.[6, 9, 11, 12, 13,14] However, development of symptomatic extradural hematomas, in a delayed fashion without and fresh antecedent injury/insult has not been well reported. We herein present a rare occurrence of a symptomatic extradural hematoma occurring as a complication to bilateral decompressive craniotomy, after a month following first surgery.

Case presentation

A 55yr old male non-diabetic, non hypertensive suffered from head injury following road traffic accident and was admitted with a GCS-E3M5V2 (11/15). No injuries to other organ systems were noted. CT scan of head showed left frontal contusion and right frontal acute SDH. Traumatic brain injury (especially following parenchymal loss) is a dynamic process, with continual changes occurring till the normalisation of the internal milieu of the cranium, which should be planned earliest after normalisation of intracranial pressure. The patient improved in sensorium and was discharged on the 10th postoperative with a GCS of 15/15 and advice for bone flap replacement after 3 months. The CT scan brain plain showed the post craniectomy status, with bilateral subgaleal collections, gliotic changes in the left frontal lobe with mild dilatation of the ventricles and bilateral subdural hygromas (Figure 2). After a month of uneventful period, the patient presented to emergency department with GCS of E3M5V2 (11/15) and dilated pupil on left side. CT scan showed acute extradural haematoma of size
9.8X3.5X6.9cm in left frontal region with mass effect in the presence of bony defect (Figure 3). He was subjected to surgical evacuation of hematoma following analysis of bleeding parameters (platelet count 1.6 lakhs/mm³ PT-13.8 sec; control-13.0secs, aPTT- 30.40 sec; control-30.0secs, fibrinogen-288 mg/dl). Intraoperatively, after elevation of the scalp flap, 50cc of blood clot was noted over the intact dura. The source of active bleeding was noted to be extradurally from the medial margin of the craniotomy defect, towards the sagittal sinus. Following evacuation, durotomy and evertion of dura over the medial margin of craniotomy flap performed for haemostasis. There was no subdural clot. Lax duroplasty was performed using temporalis fascia. The patient improved in sensorium and was evaluated for bleeding tendencies, and no definitive causative factor was found. We reviewed the history for any minimal trauma to the unprotected brain and the same was negative.

Figure 1 (a, b, c)- Axial CT scan of brain showing left frontal contusion, diffuse cerebral edema with obliterated ventricles and basal cisterns and right fronto-temporo-parietal subdural hematoma.

Figure 2- Axial CT scan of brain with bilateral calvarial defects at the time of discharge from the hospital. Noted on the left frontal region is the loss of parenchyma and bilateral subdural hygromas and mild dilatation of ventricular system.

Figure 3 (a, b, c)- Axial CT scan of brain with left frontal extradural hematoma with mass effect, midline shift with bilateral calvarial defects.
Discussion

Intraoperative or immediate postoperative development of EDH contralateral to a decompressive craniotomy, with majority of them being enlargement of pre-existing small hematomas or having a fracture, is a well known entity with an incidence varying from 0-5.7%.[1, 5, 7, 10, 11]. Various pathophysiological mechanisms described for the occurrence of contralateral hematoma are-

1. Sudden relief of tamponade effect.[3]
2. Failure of cerebral blood flow regulation which is secondary to injury.[8]
3. Vaso motor mechanisms, including vasoparalysis and vasospasm interfering with the integrity of blood vessel walls resulting in intracranial haematomas.[2]
4. Coagulopathies.[4]
5. Dysregulation of the equilibrium due to raised ICP.[5]

In our case the exact cause of delayed EDH could be attributed to dysequilibrium in pressure across the scalp flap, only on one side (side of contusion and subsequent parenchymal loss) leading to gradual stripping of the dura from the bone and snapping of extradural bridging veins. However, the exact cause of the precipitating event could not be identified. The exact incidence of such occurrence of extradural hematomas following decompressive craniotomies is not well documented in the literature. It is also surprising to find a symptomatic extradural hematoma in the absence of a bone flap. However this can be explained by its large size.

Conclusion

Symptomatic EDH can also occur after decompressive craniotomy, which should be thought of a possibility in the event of new onset swelling in a sunken flap. Traumatic brain injury (especially following parenchymal loss) is a dynamic process, with continual changes occurring till the normalisation of the internal milieu of the cranium, which should be planned earliest after normalisation of intracranial pressure.

References