Chronic Epidural Hematoma: Still a Rare Entity?

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Introduction

Extradural hematoma diagnosed more than 14 days after head injury is classified as a chronic extradural hematoma (CEDH). In the present study, we presented a series of 8 patients with CEDH in a span of 6 months.

Materials and Methods

In this article, we reported 8 cases of CEDH who presented to Nil Ratan Sircar Medical College, Kolkata, West Bengal, India, within a span of 6 months.

Discussion

Extratemporal epidural hematomas (EDHs) are often due to venous bleeding from the diploic veins or dural sinuses or to delayed rupture of a middle meningeal pseudoaneurysm. In these cases, cerebrospinal fluid may redistribute from the lateral ventricles and thus allowing room for the enlarging hematoma producing vague neurological symptoms and signs. The incidence rate of CEDH reported in the literature ranges from 3.9 to 30% of all EDHs. Computed tomography (CT) scan in CEDH often shows a low-density center surrounded by a high-density margin. Calcification of the displaced dura mater may also occur. Some are identified incidentally, whereas others are diagnosed when investigating for persistent and/or progressive neurological symptoms. Symptomatic CEDH should be surgically evacuated and has an excellent outcome the earlier it is done. In patients with no or mild symptoms, normal neurological status, and a small-sized CEDH spontaneous resolution may be expected.

Conclusion

In the post-CT era, it is always said that CEDH is a rare entity. However, in developing countries we still encounter a large number of such cases and the question arises whether CEDH is still a rare entity.

Keywords

► epidural hematoma
► uncal herniation
► neurodeficits
► head injury
► craniotomy

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Case Reports

Case 1
A 28-year-old female presented to our hospital with history of fall from moving vehicle 20 days back. She had two episodes of vomiting after the incidence with no LOC. The patient had persistent holocranial headache which was partially relieved with medications. Her general physical examination revealed healing abrasions over the left frontal region. The patient was neurologically intact. A CT scan was performed which showed left frontal EDH which was surgically evacuated (Fig. 1). Postoperative period was uneventful and patient was discharged with no neurological deficits.

Case 2
A 19-year-old boy was involved in an automobile accident 18 days back with history of transient LOC for approximately 30 seconds. He was taken to a local doctor who treated him conservatively. Due to persistent headache he presented to our hospital. He was neurologically intact with no signs of raised intracranial pressure. A CT scan was performed which showed left parietal EDH with hypodense center and hyperdense margins (Fig. 2). He was advised surgery but the patient was not willing for the same.

Case 3
A mentally retarded male patient aged 30 years presented to our hospital with history of progressive weakness of all four limbs since past 6 months. Patient had history of multiple falls due to his mental illness. At presentation, the patient was conscious, power all four limbs ⅗, and plantar extensor bilaterally. His CT scan showed EDH in the right frontoparietal, left frontal, and left parietal region with calcified walls with severe mass effect (Fig. 3). Patient underwent evacuation of EDH with removal of calcified membranes. Intraoperatively, the dura was densely calcified and could be partially removed. Patient was discharged with minimal improvement but with no added neurodeficits.

Case 4
An 11-year-old male child was brought to our hospital with history of fall from height 1 month back. He had two
episodes of vomiting following this episode. Patient was taken to a local doctor where he was treated conservatively. The parents brought the child to our hospital with complaints of depression in the left posterior part of the skull. On examination the child was conscious and oriented with no gross neurologic deficits. Local examination showed mild depression on left high parietal region. CT scan brain was done which showed left parietal EDH with depressed fracture (►Fig. 4). Patient underwent elevation of depressed fracture and evacuation of EDH. Patient was discharged in satisfactory condition.

Case 5
An 8-year-old male child with history of fall from swings while playing 20 days back presented to our outpatient department. He had a brief period of LOC and one episode of vomiting at the time of injury. Following the injury, patient showed to a local doctor and was prescribed some medications. However, the child complained of persistent headache despite taking medications. On examination child was conscious and oriented with no gross neurological deficit. CT brain was done in our hospital which showed right parietal EDH which was resolving (►Fig. 5). Patient was operated in view of large size and mass effect accounting to chronic headache. He was discharged after surgery in satisfactory condition with no added neurodeficits.

Case 6
A 32-year-old female patient with alleged history of assault 28 days back presented to our hospital with complaints of giddiness since the time of injury. Her husband reported that she has occasional irrelevant behavior with forgetfulness for 15 days. On examination the patient had a healed laceration over the right frontal region with no neurodeficits. Her mini–mental status examination was also under normal limits. CT brain was done which showed a right frontal EDH with local mass effect (►Fig. 6). Right frontal craniotomy with evacuation of CEDH was performed. Postoperative period was uneventful and patient was discharged with no added neurodeficits.
Case 7
A 14-year-old male child with history of fall from tree 19 days back presented to our hospital with history of two episodes of abnormal jerky movements of all four limbs followed by brief period of altered sensorium in the past 15 days. Patient had no history of LOC after fall. Patient also complained of mild holocranial headache since the time of injury. On examination the child was conscious and oriented with no gross neurological deficit. He had a right below elbow cast for a fracture after the fall. CT brain was done which showed left parietal EDH with mixed density of hematoma within it (►Fig. 7). Patient was advised surgery but the parents were not willing for the surgery.

Case 8
A 28-year-old male patient was brought to our emergency department with history of injury to head due to unknown mechanism 15 days back. The patient was found unknown on road side and was taken to a local hospital where no neurosurgical facility was available. Once he was recognized his relative brought him to our hospital. On examination, patient had bilateral resolving black eye with multiple healing facial lacerations. He was in altered sensorium with strongly localizing to pain. CT brain was done which showed bilateral frontal EDH with mixed density hematoma within it with right frontal contusion with diffuse edema and midline shift to right (►Fig. 8). He was operated and evacuation of bilateral EDH was done. Intraoperatively, patient had no bleed from the midline sinus. Patient had a long stay in the hospital and was discharged from the hospital in satisfactory condition with persistent neurodeficits.

Discussion
EDHs are usually located in the temporal fossa (~57–83% of cases) followed by frontal fossa (11% of cases), in the occipital region (7% of cases), at the vertex (2–8% of cases), and in the posterior fossa (4–10% of cases). When located outside the temporal fossa, the usual rapidly progressive neurological deterioration caused by transtentorial uncal herniation does not occur and hence an EDH may not be suspected clinically.

Temporal epidural bleeding, usually originates from the middle meningeal artery causing rapid blood collection and early medial uncal displacement. This leads to progressive obtundation whether or not a lucid interval has been present and a high mortality rate is common. In contrast, extratemporal EDHs are often due to venous bleeding from the diploic veins or dural sinuses or to delayed rupture of a middle meningeal pseudoaneurysm. In these extratemporal cases cerebrospinal fluid may redistribute from the lateral ventricles and thus allowing room for the enlarging hematoma. Thus, the neurological symptoms and signs are usually vague and often late. Children particularly tend to demonstrate slow neurological deterioration, perhaps because their dura is loosely adhered to the inner table of skull and gets easily stripped off and distributes the pressure of the hematoma over a larger area.

Unlike subdural hematoma, which if identified more than 21 days after injury is termed as chronic subdural hematoma, there is no consensus on the precise time-based definition of CEDH. There are various definition of CEDH in
the literature. Sparacio et al.13 used the term CEDH for those extradural hematomas operated > 48 hours after injury and Clavel et al.14 defined CEDH as EDH which are diagnosed > 72 hours after injury. Bullock and van Dellen15 used the term CEDH as the EDH with injury to surgery interval of at least 7 days. Iwakuma and Brungraber,16 however, adopted anatomopathological criterion defining CEDH as those EDH operated more than 13 days after injury and was similar to that of Zuccarello et al.17 Recently, Bradley18 has defined CEDH based on hemoglobin breakdown products on magnetic resonance imaging as extradural hematomas identified more than 14 days after head injury. This definition seems to be more scientific, recent, and evidence-based.

The incidence rate of CEDH reported in the literature ranges from 3.917 to 30%16 of all EDHs. It occurs more commonly in the younger (< 40 years) age groups.11 In our series, all 8 patients were below the age of 40 years. The pathogenetic mechanisms that can be invoked to explain chronicity in extra-axial hematomas include the presence of associated skull fractures, frontally located hematomas, age-related diffuse cerebral atrophy, venous source of bleeding, and traumatic arteriovenous fistulae of meningeal vessels.18 CT scan in CEDH often shows a low-density center surrounded by a high-density margin.19,20 The mechanism of this rim enhancement is explained by granulation tissue forming a fibrovascular neomembrane on the outer surface of the dura mater in addition to the natural enhancement of the displaced dura itself.19,20 Calcification of the displaced dura mater may also occur.20 In our series, six patients had hyperdense rim with low-density center and one patient had calcified margins of the dura intraoperatively.

Some of the CEDHs are identified incidentally, whereas others are diagnosed when investigating for persistent and/or progressive symptoms like headache, dizziness, nausea, vomiting, memory impairment, weakness of limbs, and disturbance of consciousness. Symptomatic CEDH should be surgically evacuated and has an excellent outcome the earlier it is done. However, in patients with no or mild symptoms, normal neurological status, and a small-sized CEDH without any mass effect spontaneous resolution may be expected. A watchful wait may be appropriate in such cases, but implies a high cost of serial scans and long hospitalization.12 Surgical evacuation should be considered if CEDH is observed not to be naturally absorbed during serial scans even if the patient’s condition is good, because of the likelihood of calcification.21 All our patients had persistent and/or progressive symptoms/signs during presentation. We operated on seven out of the eight patients as one patient was not willing for surgery. With the advancement in surgical techniques, neuroanesthesia, and intensive care units, it does not seem to be judicious to wait for large acute hematomas to become chronic in case of hematomas due to sinuses bleed.

The mortality and morbidity rates in CEDH cases reported in other series are known to be very low except a single patient of Zhang et al.22 In our series, one patient with calcification of CEDH wall had minimal improvement after surgery probably due to inadequate brain expansion. The patient with diffuse brain injury with bifrontal CEDH also had delayed and partial improvement due to the diffuse nature of brain injury along with the CEDH.

Conclusion
Outcome of surgical management of symptomatic CEDHs is usually excellent, though smaller ones can be managed conservatively. A watchful wait may be appropriate in small CEDH cases, but implies a high cost of serial scans and long hospitalization. Any patient with head injury, even if alert, is having mild persistent symptoms and/or signs should undergo a CT scan to rule out CEDH. In the current era of advanced neuroimaging, it is always said that CEDH is a rare entity but the question arises whether in developing country where there is a shortage of specialists and facilities, especially in rural areas, is this really true.

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Conflict of Interest
None declared.

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