Chronic subdural haematoma is predominantly a disease of the elderly. It usually follows a minor trauma. A history of direct trauma to the head is absent in up to half the cases. The common manifestations are altered mental state and focal neurological deficit. Neurological state at the time of diagnosis is the most important prognostic factor. Morbidity and mortality is higher in the elderly but outcome is good in patients who undergo neurosurgical intervention.

Chronic subdural haematoma (CSDH) is an encapsulated collection of old blood, mostly or totally liquefied and located between the dura mater and arachnoid. It was first described by Virchow in 1837 as “pachymeningitis haemorrhagica interna”. Later Trotter put forward the theory of trauma to the bridging veins as a cause of what he named “subdural haemorrhagic cyst”. Since then trauma has been recognised as an important factor in the development of CSDH.

CSDH should be differentiated from acute subdural haematoma. Acute subdural haematomas generally occur in younger adults, after a major trauma, often associated with structural brain injury, and present within 72 hours. In contrast, CSDHs often occur in the elderly after a trivial injury without any damage to the underlying brain and usually there is a period of weeks to months before it becomes clinically evident. It has a peak incidence in the sixth and seventh decade of life. Fogelholm and Waltimo estimated an incidence of 1.72/100,000 per year, the incidence increasing steeply with advancing age up to 7.35/100,000 per year in the age group 70–79.

This incidence is expected to rise further due to the continuing growth of the older population.

**RISK FACTORS**

It has long been recognised that the elderly are more likely to develop subdural haematoma, particularly from minor trauma. Generalised cerebral atrophy and increased venous fragility associated with aging are the major predisposing factors. With aging, the mass of the brain decreases leading to an increase in the space between the brain and the skull from 6% to 11% of the total intracranial space. This causes stretching of the bridging veins and the greater movement of the brain within the cranium makes these veins vulnerable to trauma.

Trauma is an important factor in the development of CSDH. However, a history of head injury (direct trauma) is absent in about 30%–50% of the cases. Indirect trauma seems to be more important. About half the patients have a history of fall but without hitting their head on the ground. In many situations the trauma is so trivial that it is forgotten. Other predisposing factors include anticoagulation, alcoholism, epilepsy, bleeding diathesis, low intracranial pressure secondary to dehydration or after the removal of cerebrospinal fluid, and receiving renal dialysis, presumably due to platelet dysfunction. As many as 24% of patients with CSDH are on warfarin or an antiplatelet drug; 5%–10% have a history of alcoholism and epilepsy.

**PATHOPHYSIOLOGY**

The initial trauma to the bridging veins results in haemorrhage in to the subdural space. A day after the haemorrhage, the outer surface of the haematoma is covered by a thin layer of fibrin and fibroblasts. Migration and proliferation of the fibroblasts leads to formation of a membrane over the clot by the fourth day. The outer membrane progressively enlarges and the fibroblasts invade the haematoma and form a thin membrane during the next two weeks. Liquefaction of the haematoma occurs due to the presence of phagocytes. Then the haematoma may either resorb spontaneously or slowly increase in size resulting in a CSDH.

Two major theories have been proposed to explain the growth of a CSDH—namely, the osmotic theory and the theory of recurrent bleeding from the haematoma capsule. Osmotic theory was based on the hypothesis that the liquefaction of the haematoma increases the protein content and oncotic pressure in the encapsulated fluid. This attracts fluid from the neighbouring vessels into the cavity due to oncotic pressure gradient across the semipermeable membrane (haematoma capsule). However this theory was disproved by Weir, who demonstrated that the osmolality of the haematoma fluid was identical to that of blood and cerebrospinal fluid.

**Box 1: Risk factors**

- Advancing age.
- Fall.
- Head injury.
- Anticoagulants/antiplatelet drugs.
- Bleeding diatheses.
- Alcohol.
- Epilepsy.
- Low intracranial pressure.
- Haemodialysis.

**Abbreviations:** CSDH, chronic subdural haematoma; MRI, magnetic resonance imaging; TND, transient neurological deficits
Recurrent bleeding from the haematoma capsule is the proved and more widely accepted theory. The haematoma capsule has been shown to have abnormal and dilated blood vessels, the source of haemorrhage. This theory was supported by the study done by Ito et al. They administered Cr-labelled red cells intravenously six to 24 hours before the evacuation of haematoma and demonstrated that it contained 0.2%–28% of fresh blood. Also increased fibrinolytic activity and coagulation abnormalities have been demonstrated within the CSDH. This may also play a part in the expansion of CSDH.

The intracranial pressure is usually normal or only slightly increased. The atrophied brain and lack of tamponading effect contributes to the gradual expansion of CSDH. The nature of the subdural collection may vary between watery, altered blood and fresh blood clots, depending on the age of the CSDH and the frequency of recurrent haemorrhages. Onset of symptoms may be delayed by weeks or even months. Because of the many ways in which CSDH can present, it has been described as “the great neurological imitator”.

COMMON PRESENTATIONS

Altered mental state
The most common presentation in the elderly (50%–70%) is altered mental state. It may manifest as varying degrees of confusion, drowsiness, or coma. Acute delirium may be very difficult to differentiate from behavioural or psychotic symptoms. Some patients are even considered to be suffering from major psychiatric illness because of depressive and paranoid symptoms. Also the diagnosis may be very easy to miss in patients with psychiatric or neurological illnesses in whom any change in behaviour or functional state is usually attributed to their pre-existing illness. In the era before computed tomography, a postmortem study on 200 psychiatric patients revealed 14 subdural haematomas of which only one had been diagnosed in life.

Focal neurological deficit
Hemiparesis was found in 58% of cases in one series. Weakness of the limbs is usually mild but drowsiness is out of proportion to the degree of neurological deficit. Mostly the deficit is contralateral but there are reports of ipsilateral symptoms. Direct pressure on the cerebral hemisphere is thought to be the underlying mechanism. Fluctuating neurological symptoms are uncommon and usually the symptoms start insidiously and progress gradually.

Headache
The incidence of headache varies in different studies ranging from 14% to 80%. It is less common in the elderly when compared with a younger patient. It is partly due to the large available intracranial space for the haematoma to accommodate before creating pressure on the adjacent brain. Another reason is the earlier onset of confusion, which attracts medical attention before the development of headache in the elderly.

Falls
Interestingly falls have been reported to be a very common presenting symptom (74%) in a recent prospective study involving 43 elderly patients. It is a well known fact that recurrent fall is a significant risk factor for CSDH. Development of CSDH may lead to recurrent falls or increase the frequency of falls due altered mental state, neurological deficits, and postural disturbances.

Seizures
Epilepsy is traditionally thought to be a rare presentation, even though it has been reported in up to 6% of cases as an initial symptom. In patients with known epilepsy increasing frequency of seizures has been noted with the development of CSDH. Simple partial seizure has been reported as a sole manifestation of CSDH, and this could be easily mistaken for a transient ischaemic attack. Seizures usually occur in the presence of a large haematoma associated with focal neurological deficit.

Transient neurological deficits
Transient neurological deficits (TND) do not always imply cerebral ischaemia. The incidence of CSDH presenting with TND varies from 1% to 12%. The most common symptom is disturbance in language and the most frequent sign is hemiplegia or hemisensory deficit. An interesting case of intermittent paraparesis due to bilateral CSDH resolving completely after drainage has been reported. The mechanisms proposed to explain TND in CSDH are intermittent mechanical pressure on the neighbouring vessels, transient increase in parenchymal swelling causing vascular displacement and ischaemia, small repeated haemorrhages in the subdural space, seizure activity with postictal deficits, and spreading cortical depression.

Extrapyramidal syndromes
CSDH causing parkinsonian symptoms is a well recognised phenomenon. In a review of 20 cases the haematoma was found to be bilateral in nine and marked improvements were seen in most patients after surgical drainage. Reversible akinetic-rigid syndrome due to bilateral CSDH with complete resolution after surgery has also been reported. The mechanisms suggested are pressure on the basal ganglia, compresion of midbrain, and circulatory disturbances in the basal ganglia caused by displacement and compression of anterior choroidal artery.

Rare neurological syndromes
Gerstmann’s syndrome (right-left disorientation, finger agnosia, agraphia, and acalculia) and progressive quadriaparesis due to CSDH has been reported in the literature. These patients made a good recovery after the evacuation of haematoma.

Ease of falling
“Ease of falling” syndrome refers to acute onset contralateral postural deficit secondary to a lesion in the basal ganglia. It is usually associated with small ischaemic lesions. The falls are contralateral slow tilting motion either laterally or diagonally backwards. The patient shows a lack of awareness and does not make postural adjustments to avoid the fall. Wali has described a case of subdural haematoma presenting as “ease of falling” syndrome which resolved completely after treatment.

DIAGNOSIS
The diagnosis of a CSDH is not usually suspected at the time of initial presentation in majority of cases. In a series of 194 cases (in 1979), CSDH was suspected only in 28% of patients. Other suspected diagnoses at the time of presentation include tumour (27%), subarachnoid haemorrhage (10%), and cerebrovascular accident (6%). However in our recent study involving 40 patients (unpublished data), cerebrovascular accident was the most common initial diagnosis (48%) followed by CSDH (20%) and others including tumour (32%).
The most important step in the diagnosis of CSDH is a high index of suspicion. It should be considered in any patient with or without a history of trauma presenting with (1) a change in mental status or worsening of pre-existent neurological or psychological illness, (2) focal neurological deficit, and (3) headache with or without focal neurological deficit. Computed tomography of the brain should be strongly considered in these patients to exclude a CSDH.

In the era before computed tomography, the diagnosis was usually made by angiography or diagnostic burr holes. The advent of computed tomography has made a major impact on the radiological diagnosis of CSDH and nowadays most of the cases are diagnosed on cranial computed tomography. A CSDH is a dynamic lesion and its appearance on computed tomography is dependent on its age (figs 1 and 2). Soon after a haemorrhage (acute phase), the haematoma looks hyperdense when compared with the normal brain, due to the presence of fresh blood. During the next few weeks (subacute phase) resolution occurs due to fibrinolysis so the haematoma appears isodense. After about four weeks (chronic phase) it appears hypodense due to the resorption of fluid. However repeated microhaemorrhages into a CSDH can increase the density, giving rise to a heterogeneous or a hyperdense picture. So classification of CSDH based on the appearance on computed tomography is far from reliable.

Hyperdense haematoma can be readily recognised but an isodense haematoma may be difficult to visualise on the computed tomogram. A specific finding is the displacement of the brain parenchyma away from the skull and the usual convex border appears flattened or even concave. Also several other indirect features due to the displacement of the brain—for example, effacement of the sulci, compression of the ipsilateral ventricle leading to midline shift, deformity of the normal ventricular anatomy, and obliteration of the basal cisterns—could aid in the diagnosis. Bilateral haematomas may lead to medial compression of both ventricles resulting in a narrow, slit-like elongated ventricle (so called “squeezed ventricle” or “rabbit’s ears”). If in doubt, a contrast computed tomogram may show displacement of cortical vessels and a delayed scan may reveal accumulation of contrast material in the subdural collection.

However magnetic resonance imaging (MRI) scan may be required in patients with isodense haematoma without midline shift and in identifying small collections at the vertex, base of the skull and in the posterior fossa. It has been clearly shown that MRI is better than computed tomography in identifying small and transversely oriented collections where the computed tomogram has failed to identify a collection in as much as 80% of cases. Even though some of these lesions may not need surgical intervention, they have significant therapeutic implications such as prevention of anticoagulation in these patients.

Even though MRI has advantages, computed tomography remains the procedure of choice in the acute setting because of shorter examination time, which is important in acutely ill patients, reliability in identifying other parenchymal lesions, no magnetic interference (especially in patients on life support machines) and the ready availability.

MANAGEMENT

Treatment of CSDH is by surgical evacuation, although small haematomas may resolve spontaneously. A recent study has shown that 23% of the patients did not warrant surgery...
because the haematoma was small. Patients treated conservatively should be carefully monitored and the scan should be repeated if there is a clinical deterioration. Some studies have shown that concurrent use of high dose steroids accelerate the resolution of subdural collection. But these studies were done in 1970s involving small number of patients and there is no strong evidence to advocate the routine use of steroids in CSDH.

The commonly followed surgical procedures include drainage by twist drill/burr hole craniostomy or craniotomy. Twist drill trephination was associated with lower mortality rate, reoperation rate, and duration of inpatient stay compared with burr hole craniostomy. Craniotomy is usually reserved for those patients in whom there is reaccumulation with recurrence of symptoms or where there is a solid haematoma. Recently Reinges and colleagues have described a less invasive bedside technique for treating CSDH. They performed twist drill craniostomy under local anaesthesia and drained the fluid through a cannula by gravity. The procedure was repeated if reaccumulation occurred and it was done up to five times in unilateral haematoma and up to 10 times in bilateral haematoma. If there was no improvement, insufficient evacuation or development of subdural empyema, they proceeded to burr hole evacuation or craniotomy (9% of patients). With very low complication rates and good results, they recommend this minimally invasive procedure especially for severely ill patients. However wider acceptance of this treatment remains to be seen.

**COMPPLICATIONS**

The following complications are encountered in addition to the usual postoperative problems such as infection and inappropriate secretion of antidiuretic hormone.

**Recurrence**

Reaccumulation of the haematoma is the most common postoperative problem. Residual fluid can be detected on computed tomography in as many as 80% of the patients, a majority of them asymptomatic and clinically insignificant. Symptomatic recurrence has been noted in 8%–37% of postoperative patients. It usually occurs between four days to four weeks with an average interval of 12 days. Clinical deterioration with radiological evidence brings attention to this condition. It is more common in the elderly and inadequate expansion of the brain following the evacuation of the haematoma is thought to play a part.

**Seizures**

Around 11% of patients develop seizures after surgery. Patients with a previous history of epilepsy are at particular risk to develop postoperative seizures. It has been recommended that prophylactic anticonvulsants should be started preoperatively and continued for six months.

**Tension pneumocephalus**

Development of tension pneumocephalus after burr hole evacuation of CSDH is a rare postoperative complication. The chronically compressed brain is thought to contribute to the ingress of intracranial air. The slow re-expansion of the brain and trapped subdural air leads to increase in intracranial pressure leading to neurological deterioration. This complication has been reported in as many as 8% of patients after surgical intervention. Cranioectomy and aspiration is the usual treatment.

**PROGNOSIS**

The morbidity and mortality in CSDH varies widely in the literature. The overall in-hospital mortality during index admission was found to be 15.6% for patients with CSDH in a large series involving 157 patients. However the outcome is good in patients who undergo neurosurgical intervention where the morbidity and mortality after surgery is around 16% and 6.5% respectively. The significant difference is due to the fact that critically ill patients are not considered fit for surgery resulting in a higher overall mortality.

In a recent prospective study of 43 patients, only 16 (37%) were able to undergo surgical intervention, four were too ill, and one died soon after the scan. The rest of the patients were treated conservatively for either the haematoma was small (10 patients) or for other reasons not clearly identified in the study. Six month mortality was 31% (13 patients) of which only one death occurred in the operated group. In patients who died, CSDH was the direct cause of death in half and the rest were due to underlying disease.

Neurological status at the time of diagnosis is the most significant prognostic factor. The influence of age on the morbidity and mortality is controversial and several studies have shown no relationship with age. However in a multivariate model, increasing age was significantly associated with mortality, but its contribution was small compared with the level of consciousness. In general, morbidity and mortality increase with advancing age and a major contributing factor to a poorer prognosis is frailty and the presence of multiple concomitant medical problems.

**QUESTIONS (TRUE/FALSE; ANSWERS ON NEXT PAGE)**

1. The following are risk factors for CSDH:
   - Advancing age.
   - Antidepressants.
   - Aspirin.
   - Falls.

2. The following statements are true in CSDH:
   - Occurs after a trivial injury.
   - Associated with brain contusion and damage.
   - Evident soon after injury.
   - Mechanism involves trauma to the bridging veins.

3. The common presenting features of CSDH in the elderly are:
   - Fluctuating neurological weakness.
   - Drowsiness.
   - Headache.
   - Falls.

4. The uncommon manifestations of CSDH are:
   - Diplopia.
   - Ataxia.
   - Parkinson’s disease.

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**Box 4: Key references**

- Cameron MM. Chronic subdural haematoma: a review of 114 cases. *J Neurol Neurosurg Psychiatry* 1978; 41: 834–9.
5. The diagnosis of CSDH should be suspected in patients with:
(A) Increasing confusion.
(B) Recurrent falls.
(C) Neurological deficit.
(D) Rapidly worsening extrapyramidal symptoms.

6. The important prognostic factors are:
(A) Age.
(B) Clinical picture at the time of diagnosis.
(C) Coexistent medical problems.
(D) Bilateral haematoma.

REFERENCES


ANSWERS (T = TRUE/F = FALSE)
1. (A) T, (B) F, (C) T, (D) T
2. (A) T, (B) F, (C) F, (D) T
3. (A) E, (B) T, (C) T, (D) F
4. (A) T, (B) T, (C) T, (D) F
5. (A) T, (B) T, (C) T, (D) T
6. (A) T, (B) T, (C) T, (D) T

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