Traumatic Brain Injury and Intracranial Hemorrhage Induced Cerebral Vasospasm and Delayed Cerebral Ischemia in a Patient with a Fall - Significance of the High-Cost Tertiary Investigations in Medico-Legal Practice; a Case Report

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Abstract
Delayed morbidity and mortality following traumatic and spontaneous intracranial haemorrhages (ICH) are common in both neurological units and autopsy rooms. However, the causes of the delayed morbidity and mortality are complications of prolonged bed care such as aspiration or hypostatic pneumonia, urinary tract infection, dehydration, electrolyte imbalance, malnutrition and sepsis following bed sores. It can also be caused during the conservative management up to 14-21 days by delayed vasospasm and cerebral ischemia. A 67-year-old man with a history of hypertension was admitted to an ETU following a fall backwards. He lost his consciousness immediately after the fall for less than five minutes with retrograde amnesia, right sided paralysis and vomiting. On admission, blood pressure was 200/100mmHg and random blood sugar was 117mg/dl. Initial CT-brain revealed bilateral frontal lobe contusions with subarachnoid haemorrhages. In the second CT, an isolated left sided ICH that was not continuous with the brain surface was detected and the right sided hemiplegia was improved following evacuation. However, after 48 hours, he developed left sided hemiplegia. Repeat CT-brain, CT-angiogram and DSA-scan revealed no new hemorrhages, infarction, necrosis etc. There were no external injuries on the body. However, the MRI brain, showed a localized oedema of right supra-callosoal area. Contre-coup contusions identified by CT-scan were compatible with the given history of fall backwards. Isolated left ICH identified in the second CT was compatible with a spontaneous ICH resulting right sided hemiplegia and could have precipitated the fall backwards. This ICH may be due to rupture of vessels due to long standing hypertension. The delayed left sided hemiplegia could have been due to the oedema of right supra-callosoal area probably initiated by delayed cerebral ischemia following vasospasm. This case also highlights the importance of radiological investigations in clinical forensic medicine for reconstruction of the events.

Keywords: ICH, traumatic, CT-Scan, fall, blood

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Introduction
Sudden and delayed death and morbidity due to traumatic head injury and spontaneous intracranial haemorrhages are common in day to day practice in neurosurgical unit and autopsy rooms.¹ Traumatic head injury includes subdural haemorrhage, extradural haemorrhage, subarachnoid haemorrhage, intracerebral haemorrhage, diffuse axonal injury, and contusions and lacerations of brain.² Natural causes of intracranial haemorrhages are hypertension, infections, bleeding disorders (haemophilia, thrombocytopenia, DIC), ruptured aneurism, arterio venous malformations, congenital anomalies of blood vessels (cavernous malformation), Vascular tumors (angioma), intra cranial primary and secondary tumors (malignant) of the brain and drugs (thrombolytic agents, enoxaparin and warfarin and cocaine).³ Traumatic causes of intracranial haemorrhages should be differentiated from the natural causes in forensic practice.¹ Sometimes, it is a difficult task in clinical
cases when both conditions co-exist. It is extremely difficult when lack of external injuries and the unavailability of the tertiary investigation facilities.

Case report
A 67-year-old man was admitted to the ETU department of peripheral hospital with sudden fall on backwards while he was standing from the bed after sleeping. History revealed that he had immediate loss of consciousness for less than five minutes, retrograde amnesia of the event and severe vomiting. He had past history and family history of diabetes, hypertension and ischemic heart disease. He also had taken Aspirin 75 mg as a routine. He was transferred to a tertiary care hospital for further management due to the deterioration in the GCS scores.

BHT revealed that on admission, his GCS was 14, blood pressure was 200/100 mmHg, random blood sugar was 117mg/dl and ECG was normal. Neurological examination was unremarkable on admission except confused verbal response. Initial Non-contrast CT-brain (Figure 01) revealed that small bilateral frontal lobe contusions and thin subarachnoid haemorrhage but no cerebral oedema. His GCS level was reduced to 11-12 and developed right hemiplegia with aphasia at eight hours after the admission. Repeat CT-scan (Figure 02) revealed an isolated left intracerebral haemorrhage (ICH) near the supra-callosal region.

Patient underwent CT-angiogram (Figure 03) and digital subtraction angiography (DSA) of the brain (Figure 04) that revealed no aneurisms, rupture of main blood vessels or vascular malformations. He was medically managed and he was undergone evacuation of hematoma by craniotomy. Recovery was uneventful. Right side hemiplegia was improved.
Figure 03 (CT-angiogram) and Figure 04 (Digital subtraction angiography [DSA] of the brain) revealed absence of aneurisms, rupture of main blood vessels or vascular malformations.

After 48 hours he developed paralysis of left upper limb and lower limb. Repeat Non-contrast CT-brain, CT-angiogram (Figure 05) and DSA-scan (06) revealed no new hemorrhages, infarctions or necrosis. But MRI brain revealed oedema of the brain near to callosal region.

Figure 05 (Repeat CT-angiogram) and Figure 06 (Repeat DSA-scan) revealed no new hemorrhages, infarctions or necrosis.

Medico-legal examination revealed no external injuries on the body. Original clothing was not available for the examination. He was managed in ICU with prolonged bedridden care, insulin, a calcium channel blocker nimodipine, GTN infusion, noradrenaline and intravenous antibiotics. Aspirin and the antiplatelet medication clopidogrel were omitted. His blood pressure maintained at 190/110 mmHg in the ICU to maintain the perfusion pressure. His aphasia and left sided weakness was slightly improved on 12th day of admission.

Medico-legal examination form was issued by the police according to the request made by ward doctors, because of the suspicious circumstances due to missing of the bystander during the ward stay.
Discussion
He did not have external injuries on the body. Only
had bilateral small frontal contusion, subarachnoid
haemorrhage and isolated intracerebral hemorrhage.
Traumatic origin of intra cranial injuries should be
differentiated from not traumatic injuries for medico-
legal purpose. Even he had no external injuries to
compare, the CT evidence of bilateral frontal lobe
contusions (contra-coup injuries) and subarachnoid
haemorrhages were compatible given history of fall
backwards.[14] This is compatible with earlier studies
that one can develop head injury by intentional and
unintentional fall through his own height.[14]

There were no external injuries. It may be due to
interference by cloths, scalp hair and/or force
generated was not enough to cause tissue damage or
skull fracture.

Even he had a no aneurism, ruptured blood vessels or
vascular malformations on CT angiography and DSA
scan, ICH detected on the second CT was isolated in
location and it was not continuous with surface
indicates that it is unlikely to be originated from the
trauma.[14] It may be due rupture of weaken small
vessels related to long standing hypertension which is
very high on admission.[15] It may be aggravated by
blood thinning agents.[3]

His right side weakness was recovered after
evacuation of ICH by craniotomy, but he developed
left side hemiplegia after 48 hours while he was in the
ICU. Repeated radiological images excluded
atherosclerotic / thrombotic occlusion of blood
vessels,[6] new haemorrhages, repeated bleeding, brain
infarction/ necrosis, venous sinus thrombosis, tumor
and infection of brain and meninges[7] except oedema
on right cerebrum in localized area near to supra
callosal area. According to the neurosurgeon, the left
side weakness may be due to delayed cerebral
ischemia fallowing vasospasm.[8,10] Other possible
explanation was oedema near to corpus callosum
compressing the fibers in the white matter.

Theoretically ICH can cause vascular spasm within 7–
14 days, but can extended up to twenty-one (21)
days. [9] It is common in younger persons with ICH
than older. [9,10] If vascular spasm involves that artery
supplying to vital Centre of the brain can cause sudden
morbidity or death at any point even though amount of
ICH is not fatal. It can be demonstrated in clinical
cases but almost impossible in autopsy cases like
cardiac Reynaud phenomenon.[5]

This case revealed traumatic frontal lobe contusion
and SAH and non-traumatic ICH. Commonly SAH at
the base of the brain is causing the vascular spasm. But
delayed spasm can occur commonly with SAH at any
places.[10] It is less likely with intra cerebral hemorrhage.[10] If vascular spasm caused by
spontaneous ICH, criminal liability may be less when
traumatic and non-traumatic ICH co-exist and it may
be significant when considering the insurance and
compensation. In this case it is difficult to find the
cause of vascular spasm whether it was caused by non-
traumatic ICH or Traumatic ICH. It may be very
significant in the case of non-accidental falls
especially in assaults.

This patient had traumatic ICH without the evidence
of external injuries on body, scalp and skull bones.
This is compatible with earlier studies that one can
develop head injury without any external evidence of
fall.[4]

Interpretation of the findings is very important to
reconstruct the events medico legally. It will aid the
legal system to implement the justice in significant
cases when traumatic and non-traumatic ICH co-exist.
(a) Whether traumatic ICH was preceded the
spontaneous ICH or vice versa? (b) This patient
developed left sided hemiplegia after the craniotomy
at 48 hours of admission. Whether it was caused by
traumatic ICH or spontaneous ICH?

Even spontaneous ICH was visible on second non
contrast CT which was taken after 10 hours of first CT.
Spontaneous ICH might developed just before the fall
which was not visible on first CT scan and it may be
the reason for the fall. With time haemorrhage may be
expanded with the additional contribution of Aspirin
and it was visible on the second CT scan.

He already had ischemic heart disease and diabetes
mellitus which were under control with medication.
His random blood sugar was 117mg/dl and ECG was
normal on admission. These findings rule out the
hypoglycemia and cardiac causes as a cause for the
sudden fall most probably.

The possibility of stress induced hypertensive ICH
fallowing fall cannot be ruled out completely, but it is
a remote possibility.

He may have developed left sided hemiplegia after 48
hours most probably due to delayed cerebral ischemia
fallowing subarachnoid haemorrhage.[9,10] But it might
be due to fallowing intracerebral haemorrhage and/or
surgical/radiological intervention.[10] Vasospasm was
managed with nimodipine and nor-adrenaline[11] and
also by maintaining the blood pressure at higher level
to sustain the perfusion pressure. His aphasia and left
sided weakness was improved on 12th day which may support the theory of delayed cerebral ischemia following subarachnoid haemorrhage. This finding explain the delayed mortality and morbidity in a patient with traumatic and non-traumatic intracranial haemorrhages.

Most of the findings were detected and events are reconstructed medico-legally with the help of tertiary investigations such as CT scan - non contrast brain, angiography of brain, DSA scan and MRI brain. It highlights that importance of radiological investigations in Forensic practice in addition to the clinical management.

Analysis of the intra cranial hemorrhages with radiological investigations and medico legal examination confirmed that given history of non-intentional fall by close eye witness and excluded the foul play.

**Conclusions**

Contre-coup contusions identified by CT-scan were compatible with the given history of fall backwards. Isolated left sided ICH identified in the second CT-scan was compatible with a spontaneous ICH resulting right sided hemiplegia and could have precipitated the fall backwards. This ICH may be due to rupture of vessels due to long standing hypertension. The delayed left sided hemiplegia could have been due to the oedema of right supra-callosal area probably initiated by delayed cerebral ischemia following vasospasm. This case highlights the importance of radiological investigation in clinical forensic medicine for reconstruction of the events and also confirms the previous studies of delayed cerebral ischemia following traumatic brain injury and intracranial hemorrhage.

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