Unusual presentation of infective endocarditis with cerebral infarction in a neuro-trauma patient

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One of the devastating neurologic complications of infective endocarditis is middle cerebral artery territory cerebral infarction due to embolic events causing ischaemia. We encountered an interesting case of infective endocarditis complicated by cerebral infarction in a patient with traumatic brain injury who had undergone craniotomy a month before the event. His head injury and its complications mislead the possibility of early suspicion of infective endocarditis.

Keywords: infective endocarditis; cerebral infarction; embolic brain infarction; traumatic brain injury; craniotomy

Introduction
Infective Endocarditis (IE) may present with devastating neurologic consequences, often manifested as the first sign of illness¹,²,³. The incidence of central nervous system (CNS) complications in infective endocarditis is 12% to 40%¹,²,³,⁴,⁵. Embolic event is the most frequent CNS manifestation and significantly associated with Staphylococcus aureus infection¹,⁷. Infective endocarditis–associated neurological complications have a mortality rate of about 20-25% in treated patients¹. The neurological manifestations of the IE include embolic brain infarction, Transient ischaemic attacks (TIA), cerebral haemorrhage, meningitis, brain abscess, toxic encephalopathy, seizures, and headache¹,³. Cerebral emboli commonly (> 90%) lodge in the distribution bed of the middle cerebral artery (MCA)⁶,⁷. The highest incidence of embolic complications is seen with aortic- and mitral-valve infections²,⁷. Here we present a case of embolic brain infarction predominantly in the middle cerebral artery territory associated with infective endocarditis, which manifested a month after a head injury. His previous history of head injury and craniotomy and the nature of it unusual presentation might have had significant impact on the diagnosis and the final outcome.

Case report
A sixty nine year old gentleman was admitted to the surgical casualty unit of our District General hospital following sudden onset of loss of consciousness. His GCS on admission was 4/15; he was intubated and admitted to our multi-disciplinary semi-open intensive care unit for further management. A month before, he had undergone left temporo-parietal craniotomy and evacuation of intracranial haematoma (ICH) causing a midline shift (MLS), following a road traffic accident. After surgery in a regional neurosurgical unit, he was transferred to our district General hospital for further care, and discharged home from the surgical ward two weeks later with a GCS of 11/15.

On the current admission to the ICU, he was febrile (101°F), tachycardic (130/min) and hypotensive (90/60 mmHg). Following investigation results were observed. White cell count (WCC) was 5550/mm³ with neutrophils (PNL) 86.1%. Platelet count and serum creatinine were 74000/mm³ and 1.4mg/dL, respectively. International normalised ratio (INR) was 2.08 and prothrombin time was 21.6 seconds. Non-contrast CT brain did not show MLS or any significant changes from his post-operative CT (Figure 1). Intracranial infection, meningitis or encephalitis were the differential diagnosis and intravenous ceftriaxone 2g twice a day and intravenous acyclovir 500mg thrice a day were started after taking blood and urine cultures. Intravenous noradrenaline infusion was started after giving fluid boluses and insertion of a central venous catheter. Patient was paralysed with atracurium, sedated with fentanyl and midazolam infusions, and ventilated for 48 hours. Despite the treatment, his WCC rose further to 15460/mm³, platelets dropped to 45000/mm³ and C Reactive Protein was 212mg/L. Platelet...
transfusion was given to keep the platelet count above 100,000/mm$^3$.

Three days later his blood culture isolated multi-resistant *Staphylococcus aureus* (MRSA) and tracheal secretions isolated coliforms. Intravenous vancomycin 1g daily was added and ceftriaxone dose was increased to 2g 8 hourly. He also had evidence of right lower-lobe consolidation in the chest radiograph.

His urine output reduced and serum creatinine was 1.61mg/dl. Ultrasound scan (USS) revealed evidence of renal parenchymal disease, right sided pleural effusion and the consolidation of inferior lobe of the right lung. He had no evidence of pericardial effusions or abdominal collections. Repeated non contrast CT of the head revealed extensive right side cerebral infarction. *(Figures 2 and 3)*

Reviewing the CT scan and the condition of the patient, consultant neurosurgeon advised on conservative management. Meanwhile the patient was increasingly becoming septic with WCC rising to 27,500/mm$^3$. He was also becoming more thrombocytaemic and coagulopathic which needed further platelets, fresh frozen plasma and cryoprecipitate transfusions. On day 6 in ICU, he developed supraventricular tachycardia (SVT) with a heart rate exceeding 160 beats/min. Intravenous amiodarone 300mg was given and 900mg infusion started.

The following day, 2D Echocardiogram was done as his cardiovascular status was increasingly unstable and he was becoming more septic (WCC 43000/mm$^3$). Furthermore, a cardiac murmur could clearly be heard. 2D echocardiogram revealed multiple vegetations on the mitral valve with grade III mitral regurgitation suggesting infective endocarditis.
On the same day despite intensive treatment he suffered a cardiac arrest and could not be revived.

Discussion
This patient presented with sudden loss of consciousness one month after a head injury and intracranial surgery. Thus, the focus of attention had mainly been drawn to secondary cerebral haemorrhage and cerebral infection, and other common sources of infection such as chest infection causing sepsis. Although infective endocarditis may present as neurological deficit as the 1st presentation, there was no evidence to suspect infective endocarditis until the 2nd CT scan of the head after 48 hours revealed extensive cerebral infarction. Renal impairment, pyrexia, and severe systemic inflammatory response he had, were non-specific and considered to be due to severe sepsis.

MCA embolic phenomenon, and cerebrovascular complications of staphylococcal infection (especially multiresistant *staphylococcus aureus*) increase the risk of mortality related to infective endocarditis. The MCA territory is the most frequently affected arterial territory in patients with IE. Most develop partial middle cerebral artery (MCA) stroke, and a few present with a complete MCA stroke. It has been shown that *staphylococcus aureus* infection has a significant relationship with the occurrence of a neurologic complications and death. This patient suffered almost complete MCA stroke and had MRSA infection in blood culture. As with the case of this patient, mitral valve vegetation has been found to influence the occurrence of neurologic events with a higher risk.

Conclusions
The presentation of this patient with infective endocarditis causing cerebral infarction was an unusual clinical scenario, as it followed traumatic brain injury and craniotomy. Many probable intracranial and extracranial complications of IE were not considered as those caused by infective endocarditis, as those complications were thought to be caused by his traumatic brain injury and secondary sepsis.

Infective endocarditis could be a cause of sudden neurological deterioration in some patients, especially, if they present with features of sepsis. Thus, high index of suspicion of IE is needed in such patients during the initial workup, to treat IE and to avoid devastating complications.

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References

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