Cerebellar Infarct Accompanied by Acute Hydrocephalus: A Case Report of 1-Year Follow-up in Rural Neurosurgical Practice

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Introduction

Cerebellar infarctions account for about 2–3% of all ischemic strokes [1], [2], [3]. Massive cerebellar edema with acute hydrocephalus due to brainstem compression or compression of the cerebrospinal fluid (CSF) flows is a rare manifestation of a stroke of the posterior circulation. The condition is considered one of the most life-threatening complications in cerebellar infarct due to the possibility of transtentorial and upward transtentorial herniation. The management of patients with cerebellar infarct is challenging, because the patient usually presents with non-specific signs and symptoms until the patient loses consciousness. Standard management should be provided by a stroke unit team or neurointensive care unit. The precision timing of treatment and evaluation with close observation is crucial, even when there is no lifethreatening condition at initial presentation, but sometimes, it is difficult to fulfill in rural areas due to the substandard facilities and lack of resources.

Case Report:

Here, we report a case of cerebellar infarct with massive edema in association with acute hydrocephalus with the progressive deterioration that happened in a rural area. A 59-year-old male patient complained about an episode of sudden headache which was followed by dizziness, vomiting, and loss of balance. A head non-contrast computerized tomography (CT) scan in the emergency room is performed 4 h after ictus, showed a slightly hypodense lesion in the left cerebellum, without accompanying edema and hydrocephalus. The patient was then managed conservatively in the ward. In the next 36 h, his consciousness level was reduced and a head CT scan evaluation showed the development of massive edema of cerebellar infarct with acute hydrocephalus. The patient underwent an emergency surgical procedure with suboccipital decompressive craniectomy with stereotactic, expanded duraplasty, and ventricular drainage (ventriculoperitoneal shunt).

CONCLUSION: Satisfactory results with rapid resolution of Glasgow Coma Scale were seen at daily follow-up after surgery. A 1-year follow-up also showed remarkable outcomes.
followed by dizziness, vomiting, and loss of balance. On arrival, he was still conscious (Glasgow Coma Scale [GCS] 15/15). His vital sign was unremarkable, except for the blood pressure (i.e., 150/90 mmHg). Both of his pupils were normal in size and reactive. However, he developed ataxia of the trunk with a tendency to fall toward the left. Muscle strength was normal in all extremities. He was a smoker and had a history of hypertension with irregular use of antihypertensive drugs, but there was no history of STEMI (ST elevation myocardial infarction) or atrial fibrillation.

The patient was then treated in the ward with close observation. About 36 h after initial treatment, the patient's level of consciousness began to decrease (GCS 7/15), accompanied by an increase in blood pressure (170/98 mmHg). Computerized tomography (CT) scan was performed to evaluate the progress of the patient's disease.

**Radiological features**

A CT scan of the brain in ER was performed 4 h after ictus, showing a slightly hypodense lesion in the left cerebellum, without accompanying edema, and signs of hydrocephalus (Figure 1).

A CT scan of the brain evaluation was then performed after 36 h following the initial management on the ward when the patient was showing some signs of clinical deterioration. It showed a hypodense lesion in the left cerebellum which corresponds to the territory of the superior cerebellar artery that is posterior circulation. This was also accompanied by signs of compression to the brainstem and the fourth ventricle, as well as the development of signs of hydrocephalus that was characterized by a widening temporal horn, the third ventricle, and ballooning of the lateral ventricles (Figure 2).

**Surgical finding and post-operative evaluation**

Suboccipital decompressive craniectomy (SDC) with expanded duraplasty and ventriculoperitoneal (VP) shunt insertion was performed. A midline incision was made approximately 4 cm above the inion to the spinous process of VC2. Harvesting of the fascia was carried out to prepare for the duraplasty procedure. Two burr holes on the occipital bone were made, then followed by thinning of the bone with high-speed drill and craniectomy using Leksell and Kerrison rongeurs. After the bone was removed, the dura mater appeared tense and did not show any presence of pulsations. Before the dura incision was performed, we performed VP shunt insertion using Frazier’s point as the entry point. After CSF was diverted, a Y-shaped incision was performed on the dura, and the necrotic cerebellum parenchyma came out spontaneously. Strokectomy was done, then continued with duraplasty using fascia as a replacement.

The procedure was completed in 150 min without any complications during the operation. Postoperatively, the patient was treated in the intensive care unit (ICU) with improvement in GCS 13/15. Patients were treated for a day in ICU and subsequently transferred to the ward on the next day with clinical improvement. In the ward, the patient complained of dizziness and pain in the surgical wound. Both were conservatively managed. On day 3, the patient complained of double vision when seeing with both eyes, with improvement in headache and dizziness. We suspected the patient with bilateral sixth nerve palsy, which we managed conservatively. The patient was discharged on the 5th day. During the treatment, the patient did not experience fever, CSF leakage, and other complications.

**Patient follow-up**

During the follow-up, 1 week after hospital discharge, the patient visited the outpatient clinic without any neurological deficits and improvement in the sixth nerve paresis. We evaluated the wound and the stitches were removed. Four months after surgery, the patient visited the outpatient clinic with dizziness but without any neurological deficits, we performed a head CT evaluation with a favorable result (Figure 3). One year after surgery, the patient visited the outpatient clinic without any neurological deficits, we performed a head CT evaluation to ascertain the progress of his hydrocephalus and the result is remarkable (Figure 4).

**Discussion**

The management of patients with cerebellar infarct remains a challenge for the physician because the patients usually present with non-specific complaints which
continue until the patients lost consciousness [8]. Even if in the conscious condition, the patient required treatment in the stroke unit or neuro-ICU [4] and is medically treated with anti-thrombolytic intravenous therapy to improve the condition [8]. Brain CT scan sometimes does not show a sign of the infarction if it is done several hours after ictus. The magnetic resonance imaging examination with DWI sequences tends to be superior in this case and also as an assessment of the infarction degree when endovascular procedures are to be performed [13], [14], however, to perform such measures in a rural area with minimal facilities is complicated.

**Surgical consideration**

SDC with expanded duraplasty and VP shunt insertion was performed. Postoperative evaluation shows that satisfactory results with rapid resolution of GCS were seen at follow-up after surgery. Four months and 1-year follow-up evaluations also showed remarkable outcomes.

Infarct in the posterior fossa is unique and the surgeon is faced with the possibility of worsening conditions from several mechanisms. First, the compression of the brainstem by the lesion could lead to transforaminal and upward transtentorial herniation. Second, compression of CSF flows might produce secondary obstructive acute hydrocephalus, which subsequently suppresses the brainstem and also might trigger transforaminal and upward transtentorial herniation or a mixture of both. In this case, emergency management must be taken [15].

Conservative management is not recommended in cases with severe mass effects, which tend to give unsuccessful in clinical practice. On the other hand, surgical treatment is widely accepted. One study mentioned a high survival rate in patients undergoing surgery, specifically around 81.6% in patients who underwent external ventricular drainage (EVD), 76.8% in patients who underwent SDC, and 77.5% who underwent EVD and SDC [4]. The other case report also reported a good outcome in the patient with cerebellar infarction complicated with acute hydrocephalus that underwent VP only [16] and SDC with VP shunt [5].

Figure 2: Non-contrast head computerized tomography scan, 36 h post-ictus, infarct area of the left cerebellar with massive edema accompanied by hydrocephalus
The main concept of the procedures in patients with cerebellar mass effect is decompression of the posterior fossa to prevent the brainstem compression and the occurrence of transforaminal herniation. Furthermore, ventricular drainage with VP shunt is essential to release the high intracranial pressure in the supratentorial compartment. However, such a procedure should be performed after the pressure on the posterior fossa is reduced to prevent upward transtentorial herniation because of immediately significant pressure changes between the infratentorial and supratentorial compartments [15].

**Regional referral rural hospital**

This case occurred in Dr. Yuliddin Away General Hospital, Tapak Tuan, a district hospital in South Aceh which was located about 440 km from Banda Aceh, the capital city of Aceh Province, Indonesia. This hospital
is a regional referral rural hospital from three districts around it (Subulussalam General District Hospital, Aceh Singkil General District Hospital, and Simeulue General District Hospital) with a total population of four districts which were 536,827 people [17], [18]. We started neurosurgical services in April 2019 to reduce disparity in neurosurgery health services in Indonesia, particularly in Aceh, with collaboration among Neurosurgery Division of Universitas Gadjah Mada, local government, health insurance, and other institutions. Neurosurgical services provide by a senior neurosurgical resident under supervision by consultant neurosurgeons on the whole process of services [18]. Nevertheless, this is certainly full of great challenges to start neurosurgery services in rural setting, these challenges include minimal facilities, infrastructure, and lack of resources, particularly related to neurosurgical services, such as having only one neurologist, without neuro nurses, neuro-physiotherapists, neurophysiology, and also without improper neurointensive care and neuromonitoring.

The option to transfer the patient to another hospital with a better facility could make him lose the

Figure 4: Non-contrast head computerized tomography scan 1 year after surgery
golden period of brain and life-saving, and also was denied by the family members due to the long transport distance (estimated 9 h away by ground ambulance).

**Conclusion**

The patients with cerebellar infarct require close observation for general condition and neurological signs in the stroke unit or neuro-ICU, even when there is no life-threatening condition at initial presentation in the ER. Inevitably, this is a challenge for rural hospital areas, to improve the practice of neurosurgical service and improve the standard of care in rural settings. Collaboration among hospitals, government, and related institutions is needed to improve facilities, infrastructure, and training programs among healthcare staffs, to improve the quality of neurosurgery services to reduce morbidity and mortality. In patients in whom surgery is indicated, the choice of surgery with SDC with expanded duraplasty and ventricular drainage (EVD or VP shunt) might provide satisfactory results.

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**References**