Cerebral fat embolism after bilateral total knee replacement arthroplasty
-A case report-

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Fat embolism syndrome is a rare and potentially lethal complication most commonly seen in long bone fractures and intramedullary manipulation. The clinical triad of fat embolism syndrome consists of mental confusion, respiratory distress, and petechiae. This study reports a case of cerebral fat embolism syndrome following elective bilateral total knee replacement. After an uneventful anesthesia and initial recovery, the patient developed neurologic symptoms nine hours postoperatively. (Korean J Anesthesiol 2010; 59: S207-S210)

Key Words: Arthroplasty, Fat embolism, Total knee replacement.

Fat embolism syndrome is a change in physiology resulting from mechanical causes, trauma, or sepsis. Fat globules generated within the systemic circulation induce pulmonary dysfunction, neurological changes, dermal symptoms, and dysfunction of several other organs. Although very rare, fat embolism syndrome is a fatal disease that develops in 12-72 hours. Acute dyspnea may appear, and if not treated appropriately, may progress to cardiac arrest in rare cases [1,2]. However, due to symptom diversity and unclear diagnostic methods, it is difficult to treat the fat embolism correctly. This paper reports on the case study of a patient who developed sudden dyspnea and deterioration of the nervous system 9 hours after bilateral total knee replacement arthroplasty. The subsequent diagnosis of brain fat embolism syndrome was determined, appropriate treatments were administered, and the patient recovered without sequelae. This study recounts this case, particularly for anesthesiologists, to emphasize the necessity of comprehensive patient management even after surgery for risk group patients.

Case Report

A 74-year-old female patient (height: 157 cm, weight: 56.8 kg) was diagnosed with degenerative osteoarthritis and admitted for bilateral total knee replacement arthroplasty. The patient was also previously diagnosed with angina and hypertension 2 years earlier and placed on drug treatment (isosorbide-5-mininitrate and atenolol). Blood and biochemistry tests performed prior to surgery were within normal ranges. Similarly, on simple chest radiograph, no special findings were observed except...
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Several bilateral small high signal intensity lesions in the cerebral and cerebellar cortices were detected on diffusion MRI. The apparent diffusion coefficient (ADC) in the area was reduced, suggesting fat embolism (Fig. 1). On 3D pulmonary arteriography, no special findings were detected.

Upon referral to the neurology department, the patient was transferred to the stroke unit the day after surgery. Immediately after transfer, her breathing became shallow and irregular and pulse oxygen saturation level decreased to 84%. She also exhibited eyeball deviation to the left side and nystagmus. The patient was unresponsive to even strong stimulation, and the conscious state was stuporous. The patient was intubated because, while being supplied with 7 L/min 100% oxygen via face mask, pulse oxygen saturation level decreased to 58% and blood pressure decreased to 70/40 mmHg. The patient was transferred to the intensive care unit, and placed on a mechanical ventilator. Blood pressure was measured at 95/60 and, at that time, a fever of 38°C was observed. After arriving on the intensive care unit, the neurology department considered administering low molecular weight heparin. Nonetheless, blood tests performed at that time revealed that hemoglobin was 8.3 g/dl and platelets had decreased to 34,000/μl. In a blood coagulation test, an increase in coagulation time was indicated; therefore the drug could not be administered. Packled RBCs and platelets were transfused, and inotropics as well as diuretics were administered. Three days after surgery, ischemic colitis was observed, and a petechial rash developed all over the entire body, including the abdomen.

Four days after surgery, blood coagulation test results were normal, so administration of low molecular weight heparin was initiated. However, the conscious state of patient was still stuporous, and neurological symptoms were unimproved. Five days after surgery, an electroencephalogram was performed from which moderate to severe diffuse cerebral dysfunction was found.

Seven days after surgery, due to improved conscious state
and arterial blood gas analysis, the patient was weaned off the mechanical ventilator. Thirteen days after surgery, the endotracheal tube was removed, and respiratory and neurological symptoms were improved. Seventeen days after surgery, the patient was transferred to a general ward. Twenty six days after surgery, brain MRI revealed that multiple brain infarctions had become normal, and the patient was discharged 44 days after surgery.

**Discussion**

It has been shown that fat embolism syndrome is caused by fracture of the ilium or other pelvic bones, and multiple trauma. However, other possible causes include severe burns, infection, kidney transplant, liposuction, cardiopulmonary bypass, and transfusions. Moreover, although rare, cases of fat embolism syndrome after gastrectomy have been reported [3-7].

Based on clinical symptoms, fat embolism syndrome is diagnosed by eliminating other possible causes. Major clinical symptoms are respiratory distress symptoms, central nervous system symptoms, and dermal symptoms. Such symptoms may be induced by various causes other than fat embolism syndrome, and thus differentiation is required. Pulmonary embolism or pneumonia may cause dyspnea and hypoxemia while hypoxemia or meningitis may cause deterioration of brain function [1]. Since our patient had a past history of ischemic heart disease, myocardial infarction or cardiac shock were considered as causes. However, since surgery was performed under CSE anesthesia, and blood pressure, pulse, electrocardiogram, oxygen saturation level, and arterial blood gas analysis were within normal ranges throughout the entire anesthesia period, these causes could be ruled out. In 75% of fat embolism syndrome patients, respiratory distress symptoms are detected first, with 10% progressing to respiratory failure [8]. Similarly, in our case study, dyspnea and hypoxemia were detected first. Regrettably, aggressive oxygenation treatments through immediate arterial blood gas analysis were not performed. Alteration of brain function is detected in 86% of patients, and the pattern varies from drowsy mental state to coma [7]. In 20–50% of patients, petechial rash is detected in the conjunctiva, around the neck, the axillary area, and the upper body. It appears within 24–36 hours and resolves spontaneously within one week [1].

Several investigators have reported criteria mainly based on clinical symptoms for the diagnosis of fat embolism syndrome. Criteria developed by Gurd and Wilson in 1974 [5] are widely used (Table 1). At least one of the major criteria and four of the minor criteria are necessary for diagnosing fat embolism syndrome [5]. Our patient exhibited all of the major symptoms of Gurd’s diagnostic criteria, as well as the minor criteria of tachycardia, oliguria, fever of 38°C, thrombocytopenia, and the elevation of erythrocyte sedimentation rate.

Radiological findings are also useful for diagnosis. Brain CT shows normal findings in most cases. As brain MRI is more sensitive, small high signal intensity lesions scattered in the cerebral white matter, cerebellum, and brain stem are detected on T2 weighted imagings or diffusion weighted imagings [4,6,7]. In our case, brain CT revealed no specific findings. Nevertheless, on T2 weighted imagings and diffusion weighted imaging of MRI, many bilateral small lesions in the cerebral white matter and the cerebellum were observed.

In fat embolism syndrome, prevention, early detection, and appropriate treatment are important. First, anesthesiologists and surgeons should recognize patients at-risk for developing fat embolism syndrome. Patients considered high risk for the development of fat embolism syndrome are both young and old, with closed or multiple fractures, as well as patients treated with only conservative therapies for iliac fracture. Particularly, it has been reported that in cases of bilateral total knee replacement arthroplasty with long tourniquet time, the risk of developing fat embolism syndrome increases [9]. In contrast, types of anesthesia, past medication history of patient prior to surgery, and type or duration of patient controlled analgesia are not associated with complications or development of cognitive dysfunction after knee arthroplasty [10]. When performing surgery on high risk patients, continuous hemodynamic monitoring by arterial cannulation and regular arterial blood gas analysis should be performed in addition to basic monitoring. In bilateral total knee replacement arthroplasty, measurement of pulmonary arterial blood pressure is also useful as a means of diagnosis and prognosis. In a recent study, during surgery, if cardiovascular system instability caused by fat embolism is detected, it is recommended to maintain perfusion pressure and use vasoconstrictors to maintain the function of the right sided heart [11]. After surgery, similarly, in conjunction

| **Table 1. Gurd’s Criteria for Diagnosis of Fat Embolism Syndrome** |
|---------------------------------|
| **Major criteria (one necessary for diagnosis)** |
| Petechial rash |
| Respiratory insufficiency |
| Cerebral involvement |
| **Minor criteria (four necessary for diagnosis)** |
| Tachycardia > 120 beats/min |
| Fever |
| Retinal changes: fat or petechiae |
| Jaundice |
| Renal signs: anuria or oliguria |
| Thrombocytopenia |
| Anemia |
| High erythrocyte sedimentation rate |
| Fat macroglobulinemia |

From Gurd and Wilson’s study [5].
with basic pulse oxygen saturation level monitoring, comprehensive monitoring of patients through regular arterial blood gas analysis should detect the early deterioration of the respiratory system of patient, and sufficient oxygen should be supplied. In our case, bilateral total knee replacement arthroplasty was performed, and a tourniquet was used for the right and left sides for 55 minutes each, which is thought to raise the risk of fat embolism syndrome. In addition, considering that consciousness altered suddenly nine hours after surgery, patients with high risk for the development of fat embolism syndrome, even without the presentation of special problems during surgery and anesthesia, should be sent to intensive care unit for continuous observation and early detection of symptoms as well as rapid and appropriate treatment.

It is important to perform conservative treatments for hypoxemia, hypotension, and reduction of the blood flow to peripheral organs simultaneously for the treatment of fat embolism syndrome. Among these treatments, securing the airway and oxygenation in tissues through oxygen supply are important [5]. In arterial blood gas analysis, PaO₂ higher than 90 mmHg should be maintained, and if required, endotracheal intubation and mechanical ventilation can be considered. Moreover, appropriate hydration is also vital. If effective blood volume within blood vessels is not maintained, pulmonary injury caused by fat embolism may be exacerbated. Albumin administration is helpful for maintaining the blood volume. In addition, it has been reported that early administration of high-dose steroid reduces the development of hypoxemia and fat embolism. Nonetheless, therapeutic and prophylactic effects of high dose steroids are still controversial [12,13]. It was reported that heparin [14] and dextran [15] exerted fat degradation effects that help treatments, but it was found ineffective in subsequent studies. Hence, further studies are required [11].

In our patient, the department of neurology recommended the administration of low molecular weight heparin during the early onset period, but because of the risk of postsurgical hemorrhage and impediment of wound healing, treatments for brain fat embolism syndrome were limited. In addition, during surgery, regional anesthesia was performed and the epidural patient controlled analgesia method (PCA) was applied. Hence, potential hemorrhage to the epidural space should have been considered. We consider that, for brain fat embolism cases, anesthesiologists, neurologists, and surgeons should communicate closely, and perform the most appropriate treatments for patients.

Our case study was a patient who developed brain fat embolism after total knee replacement arthroplasty. If such patients show a sudden deterioration of consciousness, a fat embolism with a fatal prognosis should be suspected. As shown in our case, since symptoms may manifest even nine hours after surgery, anesthesiologists should recognize high risk patients by presurgical evaluation and aggressive efforts be made to obtain relevant information on patients in advance. To prevent fat embolism syndrome, risk factors should be corrected if possible, and careful anesthetic management should be undertaken. Furthermore, early diagnosis and appropriate management of fat embolism are important and prior to and following surgery, patients should be monitored comprehensively in the intensive care unit.

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