INTRODUCTION

Fat embolism and Fat embolism syndrome (FES) are well-known complications of long bone fracture and surgery involving manipulation of skeletal elements. Many non-traumatic causes of FES have been suggested but they constitute only a small portion. FES presents with classical symptoms of petechiae, hypoxemia, central nervous system symptoms along with other features such as tachycardia and pyrexia. Diagnosis of FES relies on clinical judgment rather than objective findings such as emboli present in the retinal vessels on fundoscopy, fat globules present in urine and sputum, a sudden inexplicable drop in hematocrit or platelet values, increasing erythrocyte sedimentation rate.

KEY WORDS: Alveolar hemorrhage, fat embolism syndrome, bone fractures

CASE REPORT

A 25-year-old man was referred to us three days post road traffic accident. He was found to have open fracture shaft of left femur and fracture condyle of left humerus. There was no associated history of loss of consciousness, vomiting, convulsions or any bleed. Intra medullary nailing was done for fracture femur under general anesthesia in a peripheral hospital. Intra-operative and immediate post-operative period were uneventful. Within next 18 hours he developed altered sensorium and became restless. He was hypotensive and shortly became breathless and hypoxic. He was intubated and mechanically ventilated. Endotracheal bleed was detected post intubation. He was transferred to our center for further management. On evaluation in emergency department, he was found to be drowsy, restless, hypotensive, and there were signs of pulmonary hemorrhage in the form of blood stained endotracheal secretion with hypoxia. There were no obvious petechial rashes apart from small hemorrhagic spots in bilateral conjunctiva. Initial resuscitation was done and he was started on noradrenalin infusion in view of hypotension. Fiberoptic bronchoscopy showed blood in trachea and bilateral tracheobronchial tree. Bronchoalveolar lavage fluid was hemorrhagic suggestive of alveolar hemorrhage. Initial laboratory parameters showed: Hemoglobin 8.4 gm/dl, total leucocyte count 11300/cc, platelet count 80000/cc. Renal functions were deranged with serum urea 38 mg/dl and serum creatinine 1.2 mg/dl. His liver function tests showed serum bilirubin 2.1 mg/dl (direct 1.0 mg/dl), SGOT/SGPT 235 and 114 IU/L. His serum lactate level was 2.6 mmol/dl. Initial arterial blood gas analysis showed, pH 7.37, PaCO$_2$ 63.5 mm Hg, PaO$_2$ 70.8, SaO$_2$ 93% with FiO$_2$ of 1. X-ray chest was consistent with bilateral diffuse alveolar opacities consistent with ARDS. Other investigations include Serum d-Dimer 4.3 (Normal 0.063-0.245 mg/dl), Anti-CCP antibodies 0.43 (Normal < 5 RU/ml), ANA (Immunofluorescence) negative at 1:100 dilutions, PR-3ANCA (ELISA) 1.5 IU/ml (Normal < 6 u/ml). Bronchoalveolar lavage fluid showed numerous debris laden alveolar macrophages and...
Pearl’s stain was suggestive of intracellular positivity. Urine examination for fat globules was negative. His serum creatinine phosphokinase was 691 IU/L. EEG showed delta-theta coma pattern suggestive of modest encephalopathy.

On presentation there was evidence of subcutaneous emphysema with X-ray chest showing signs of pneumomediastenum. Initial high resolution computed tomography (HRCT) chest scan showed confluencing air space opacity with ground glass densities and interlobar septal thickening involving bilateral lung fields suggestive of acute respiratory distress syndrome (ARDS) [Figure 1]. He was electively ventilated for next three days, intercostal drain was placed in view of pneumomediastenum. Subcutaneous emphysema gradually subsided and elective tracheostomy was done on 7th day of admission. Repeat HRCT chest on day 6 revealed patchy ill defined air space opacity in bilateral lower lobes with ground glassing in bilateral upper and middle lobes. Right pneumothorax, pneumo mediastenum, excessive subcutaneous emphysema. ARDS changes in bilateral lung fields were significantly reduced [Figure 2]. His neurological status gradually improved over next 5 days and was put on T piece on 12th day, which he tolerated well. He underwent fixation of fracture humerus condyle, which was uneventful. He was decannulated on 15th day of admission and discharged on 18th day.

**DISCUSSION**

Original paper describing fat embolism dates from 1873. Fat embolism is defined as a blockage of blood vessels by intravascular fat globules ranging from 10-40 μm in diameter.[1]

Fat embolism syndrome (FES) comprises a defined set of clinical pattern and is a serious consequence of fat embolism. Though the major cause of FES is skeletal fracture associated with trauma, small percentage (5%) of cases do have atraumatic etiology. These atraumatic causes include bone marrow transplantation, pancreatitis, sickle cell disease, burns, prolonged high-dose corticosteroid therapy, diabetes mellitus, hepatic trauma, liposuction, lipectomy, external cardiac compression, gas gangrene, decompression sickness, and lipid infusions etc.[2,3] Perioperative incidence of FES is between 3.5 to 5% in surgeries involving early fixation of fractures.[4] Age is considered to be a determining factor in the development of FES: Young men with fractures are at increased risk.[3,6] Our patient was a 25 year old male.

Single bone fracture has a rare chance of FES (1-3%). The incidence may increase with number of bones involved. Bilateral femoral fracture demonstrated an incidence of as high as 33 percent.[7] An overall mortality of 5-15% has been described.[8] Intramedullary fat embolises as a result of fracture or intramedullary procedure. This embolic phenomenon has been confirmed echocardiographically.[9]

The fat enters torn venules which are kept patent in the Haversian canals and makes way into circulation.[10] Fat globule ranging from 7-10 μm in diameter has been documented to traverse the pulmonary vasculature. Systemic embolisation has been documented due to a patent foramen ovale which may be present in 25% of individuals.[11]

Even larger size fat globule can traverse the foramen if severe pulmonary hypertension is precipitated by fat embolism. Pulmonary hypertension can cause a pressure difference between the right and left atria which leads to embolisation of larger fat globule.[10]

Systemic liberation of fat can cause FES and fulminant FES by different pathophysiological mechanisms. FES is

**Figure 1:** Computer tomography of chest showing bilateral alveolar hemorrhage

**Figure 2:** CT of chest showing recovering acute respiratory distress syndrome and pneumo thorax, pneumomediastenum with intercostal chest tube insitu
caused by perivascular hemorrhage and edema following the accumulation of fat in the pulmonary, cerebral or dermal microvasculature and local liberation of free fatty acids (FFAs). Obstructive shock caused by fat globules may result in fulminant FES.

The timing of symptomatic presentation of FES may be variable. Most typically it presents 24-72 h after the initial injury. In rare cases it may be delayed as much as 2 weeks after the insult. In our case, the patient developed respiratory distress and encephalopathy within 18 hours of surgical manipulation of the fracture.

Classically, it presents with asymptomatic interval followed by pulmonary, neurologic, and skin manifestations. It has a biphasic clinical course. The initial symptoms are caused by mechanical occlusion of blood vessels with fat globules that are too large to pass through the capillaries. Unlike other embolic events, the vascular occlusion in fat embolism is often temporary or incomplete since fat globules do not completely obstruct capillary blood flow. The late presentation is due to hydrolysis of the fat to more irritating FFAs which then migrate to other organs via the systemic circulation.

The most common system involved is respiratory (95%) followed by the central nervous system.

The most common presentation with fat embolism includes severe hypoxemia and respiratory distress. The incidence has been reported to be 50% to 96% of the affected patients and most of them require ventilation. Petechial rash in conjunctiva, oral mucosa, and skin folds in axilla and neck occurs in up to 60% of cases. The described mechanism for development of petechiae is embolization of small dermal capillaries leading to extravasation of erythrocytes. We noted oblivious petechiae in conjunctiva in both eyes of the patient [Figure 3].

A number of minor features of fat embolism syndrome may be present and these appear to result from the release of toxic mediators secondary either to the initial injury or to dysfunctional lipid metabolism. These include tachycardia, myocardial depression, ECG changes indicative of right heart strain, soft fluffy retinal exudates with macular edema, scotomata (Purtscher’s retinopathy), coagulation abnormalities (which mimics disseminated intravascular coagulation).

Fat embolism can lead to diffuse alveolar hemorrhage. The described pathogenesis is an inflammatory response caused by lipoprotein lipase which is activated by catecholamine surge caused by stress. It acts on the fat deposited in the pulmonary or systemic capillary network liberating high concentrations of toxic FFAs locally. It causes platelet aggregation, a mild disseminated intravascular coagulation, and disruption of the pulmonary and cerebral capillary walls.

Pulmonary histology usually reveals intra-alveolar hemorrhage, fat within pulmonary capillaries and oedema. Cerebral histology reveals diffuse cerebral edema with multiple hemorrhagic petechiae.

FES is a clinical diagnosis and none of the evaluated laboratory parameters found to be sensitive or specific enough to diagnose FES. Various definitive criteria (include Figure 3: Conjunctival hemorrhage

![Figure 3: Conjunctival hemorrhage](image)

**Chart 1: Diagnostic criteria for fat embolism syndrome**

<table>
<thead>
<tr>
<th>Major Criteria</th>
<th>Minor Criteria</th>
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<tbody>
<tr>
<td>Sustained PaO₂ &lt; 60 mm Hg</td>
<td>Sustained PCO₂ of &gt;55 mm Hg or a pH &lt; 7.3</td>
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<tr>
<td>Sustained respiratory rate &gt;35/min (despite sedation)</td>
<td>Increased work of breathing</td>
</tr>
<tr>
<td>Increased diuresis, accessory muscle use, tachycardia, and anxiety</td>
<td>Chest X-ray changes (diffuse alveolar infiltrates)</td>
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**GURD’S CRITERIA**

- Major
  - Asymptomatic retinal petechia (30%).
  - Hypoxemia (PaO₂ < 60 mm Hg).
  - CNS depression.
- Minor
  - Tachycardia (>110 beats/min).
  - Pyrexia (>38.5 degrees).
  - ECG changes indicative of right heart strain.
  - Baseline Sputum (fat globules).

**LINDEGUE’S CRITERIA**

- Sustained PaO₂ < 60 mm Hg
- Sustained PCO₂ of >55 mm Hg or a pH < 7.3
- Sustained respiratory rate >35/min (despite sedation)
- Increased work of breathing:
  - Dyspnoea, accessory muscle use, tachycardia, and anxiety
- Chest X-ray changes (Diffuse alveolar infiltrates)
- Hypoxemia (PaO₂ < 9.3 kPa)
- Fever (>38°C)
- Tachycardia (at least two of the following:)
  - Heart rate >120 beats/min
  - Tachypnea (>30 breaths/min)
- Cumulative score >5 required for diagnosis

**SDOFENFELD’S CRITERIA**

- Petechiae
- Chest X-ray changes (Diffuse alveolar infiltrates)
- Hypoxemia (PaO₂ < 9.3 kPa)
- Fever (>38°C)
- Tachycardia (at least two of the following:)
  - Heart rate >120 beats/min
  - Tachypnea (>30 breaths/min)
- Cumulative score >5 required for diagnosis
Clinical and laboratory parameters) have been described, which include Gurd’s (four major and three minor criteria), Lindeque’s criteria (four criteria) and Schonfeld’s criteria (14 out of 15 points) [Chart 1].

Investigations are usually performed to support the clinical diagnosis or to monitor therapy which include: Hematology and biochemistry: An unexplained anemia (70% of patients) and thrombocytopenia (platelet count < 1,50,000/cumm in up to 50% of patients). Hypocalcaemia (due to binding of free fatty acids to calcium) and elevated serum lipase have also been reported. Hypofibrinogenemia, raised erythrocyte sedimentation rate (ESR), and prolongation of prothrombin time may be present. Isolated case report suggests microemboli detected by Trans Cranial Doppler (TCD) and Magnetic Resonance Imaging (MRI) can add in diagnosing FES.

Corticosteroids act as anti-inflammatory agents and reduce the perivascular hemorrhage and edema. They have been used extensively and recommended by some for the management of FES. There is insufficient scientific evidence to support initiation of steroid therapy in established FES. An experimental study showed no beneficial effect.

Use of aspirin has been advocated. Heparin is thought to have potential for activating lipoprotein lipase thereby increases the clearance of lipemic serum. On the other hand, there is a potential risk of increase in free fatty acid level there-by possible enhancement of inflammation.

Bloody bronchoalveolar lavage specimens (with numerous erythrocytes and siderophages) establish the diagnosis of diffuse alveolar hemorrhage.

Treatment of alveolar hemorrhage include steroid. Majority of cases of pulmonary hemorrhage have an autoimmune cause and mainstay of the treatment include steroids and immunosuppression.

As in our case the cause is nonimmune, we used high dose steroids (methyl prednisolone started at 60 mg twice daily and tapered over 2 weeks with improvement of clinical state).

CONCLUSION

FES is a rare but devastating complication in patients presenting with traumatic fracture of long bones or procedure related and involving manipulation of the same. The diagnosis of FES is clinical, though use of diagnostic criteria may be helpful. Alveolar hemorrhage is a rare presentation of FES and the management is essentially supportive. Though none of the available studies have clearly supported the role of steroids, we found good result with early use of steroid in our case.

REFERENCES
