Introduction
Fat embolism occurs when fat globules from the bone marrow enter the circulation mostly after orthopaedic trauma causing fractures of long bones, pelvis or vertebra. It also occurs following surgical fixation such as intramedullary nailing, instrumented spine surgery or vertebroplasty. Fat embolism syndrome (FES) is a constellation of clinical symptoms characterized by a triad of respiratory insufficiency, altered sensorium and petechiae. Not all patient with fat embolism develops FES. FES generally occurs within 24 hours of onset of trauma or post-surgery. The incidence of FES after trauma varies from 0.25% to 35%. The diagnosis is based on clinical presentation and excluding other possible conditions. Beside histopathology, none of the investigations such as fat macroglobulinuria in urine or lipid laden macrophages in bronchoalveolar lavage (BAL) is 100% specific. At times FES can present with atypical presentation such as intra alveolar haemorrhage and only high index of suspicion can help in making diagnosis.

Case Report:
A 28 years healthy male sustained right closed femur shaft fracture in a road traffic accident. He had no history of loss of consciousness, vomiting, convulsions or any bleed. He underwent closed intramedullary nailing under spinal anaesthesia the following day which was uneventful. On 3rd post-operative day he developed sudden onset shortness of breath, low grade fever and streaky haemoptysis. He was shifted to our institute the next day and was seen in emergency. He had no history of bleeding from other orifices. His past history was not significant. He has no history of addiction. On examination he appeared pale. His mentation was normal. His pulse rate was 92/min and regular, BP was 130/90 mm Hg, RR was 18/min and temperature was 98.60F. His SpO2 was 93% in ambient air. His chest auscultation was clear with normal heart sounds. His abdomen was soft with no...
organomegaly. The right thigh was slightly swollen compared to left. There was no evidence of petechiae or deep vein thrombosis in lower limbs. Fundoscopy was normal too. On laboratory investigations his CBC showed: Haemoglobin 7.5g/dl, TLC 15400/cc with 75% neutrophils, platelets 231000/cc, Prothrombin time 13.8 (control 13.5), APTT 29.5 seconds (control 27.5). His RFT and LFT was within normal limits. His previous reports were not available during admission so the treating orthopedician was contacted over the phone and discussed the case. His haemoglobin during admission was 12g/dl and had no significant bleeding during perioperative and immediate post-operative period and post-operative haemoglobin was 11.5g/dl. An initial suspicion of fat embolism syndrome, pulmonary thromboembolism, pulmonary oedema and hospital acquired pneumonia were kept. His Chest X-ray taken in emergency showed bilateral infiltrate. An ECG showed normal sinus tachycardia and bedside 2D ECHO was normal. He underwent CT pulmonary angiogram that showed bilateral diffuse ground glass opacities with relative pleural sparing and no evidence of pulmonary embolism (figure 1). His urine routine microscopy was normal and no fat globules were identified. In view of sudden fall in haemoglobin, steaky haemoptysis and bilateral lung infiltrate a diffuse alveolar haemorrhage was suspected. He was admitted and received broad spectrum antibiotics (Meropenem and Teicoplanin) for MDR bacteria coverage, proton pump inhibitor. The next day he underwent flexible bronchoscopy with bronchoalveolar lavage (BAL) under conscious sedation that showed haemorrhagic effluent from both sides of lungs confirming diagnosis of alveolar haemorrhage (figure 2). BAL cytology revealed hemosiderin laden macrophages and few inflammatory cells predominantly neutrophils with full of RBCs. No lipid laden macrophages or fat globules identified. His autoimmune screening for ANA, ANCA and anti GBM was negative. BAL culture was sterile. His antibiotics were stopped and put on prophylactic oral antibiotics. He was transfused with one unit of packed RBC. His serial haemoglobin monitoring done daily showed no further fall and later discharged after 5 days of admission. A review literature provided us few case reports of fat embolism syndrome presenting as alveolar haemorrhage. Hence on dedication in an appropriate clinical scenario our patient too fits into the clinical criteria of fat embolism syndrome.

Discussion:

FES following trauma presenting as diffuse alveolar haemorrhage (DAH) is an unusual manifestation. A search in PubMed, Google and ResearchGate using search item “Fat embolism OR Fat embolism syndrome AND alveolar haemorrhage” lead to 11 cases published till date with 3 in French and remaining 5 in English.3-11 One case report was published as abstract presentation2 and another 1 as image of the week.5 The clinical characteristics of these cases is tabulated in table 1. The onset of symptoms is as early as 2 hours to 96 hours after the onset of trauma. Six cases had FES after performing surgery. The onset of FES after surgery was as early as during perioperative period to as late as 96 hours of surgery. Except for 1 patient whose data is not available rest all had recovered including our patient. The pathogenesis of FES is a two-phase process and takes into consideration both the mechanical and biochemical hypothesis. Immediately after trauma or surgical intervention, the fat globules get embolized into circulation. From here these droplets gets mechanically trapped in pulmonary capillary bed leading to acute rise in pulmonary arterial pressure. If massive emboli occur, it may lead to circulatory collapse and even death rapidly. Smaller size fat droplets can pass through pulmonary capillaries to systemic circulation or via patent foramen ovale in the heart. The biochemical hypothesis explains FES due to degradation of fat globules by tissue lipase into fatty acids and other intermediaries that are proinflammatory. It leads to release of various cytokines damaging endothelial lining in lung, brain, skin and other organs. In lungs, it leads to damage of alveolocapillary endothelium producing interstitial oedema and haemorrhage with subsequent development of acute lung injury or ARDS.12 The biochemical theory explains the latency period by hours to days in development of FES post trauma or surgery as well as in nontraumatic FES. However, both mechanical and biochemical theories seem mutually inclusive rather than exclusive.

A suspicion of alveolar haemorrhage is made when an unexplained anaemia or sudden fall in haemoglobin with lung infiltrate is seen.13 Sequential haemorrhagic effluent during bronchoscopic BAL confirms the diagnosis. Depending upon the clinical background, laboratory findings and/or lung histology; the aetiology of DAH is made. Alveolar haemorrhage occurs due to disruption of alveolo-capillary basement membrane that can occur either due to injury.
or inflammation of pulmonary capillaries, venules or arterioles. Three distinct histopathological pattern is seen: pulmonary capillaritis, bland pulmonary haemorrhage and diffuse alveolar damage.13 Surprisingly, none of the articles pertaining to DAH cite FES as a cause of DAH.13-15

But is alveolar haemorrhage an unusual presentation in FES? A PubMed search was made to find out research articles mentioning alveolar haemorrhage incidence in FES, but none of these literatures have focused on alveolar haemorrhage.16-18 FES is a clinical diagnosis based on an appropriate clinical scenario and the use of clinical criteria as proposed by either Gurd and Wilson,19 or Lindeque 20 or Schoenfeld. 21 However, none of them are prospectively validated or standardized. 22 The most widely used Gurd and Wilson criteria is less sensitive but more specific than other criteria.2 In diagnosis of FES by Gurd and Wilson, unexplained anaemia or fall in haemoglobin by 20% is one of its minor criteria. The fall in haemoglobin due to lack of bronchoscopy in many of cases the diagnosis of alveolar haemorrhage remains elusive. The current author feels that it is underreporting rather than an unusual event. As the focus on bronchoalveolar lavage was to look for lipid laden macrophages rather than hemosiderin laden macrophages. Moreover, the presence of lipid laden macrophages was nonspecific in diagnosis of FES hence the routine use of bronchoscopy and BAL has gone against favour in those who meet the clinical criteria in the appropriate setting.

In a Serbian journal published in 2000 by Nikolic S et al, alveolar haemorrhage was found in about two third of autopsy cases of 56 patients who died of post traumatic FES.23

Conclusion:

Rapid fall in haemoglobin with new lung infiltrate in a patient with trauma can be a sign of fat embolism syndrome.

Conflict of interest:
None

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References

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<tr>
<th>Case</th>
<th>Year</th>
<th>Age (years)/ Sex</th>
<th>Type of fracture*</th>
<th>Onset of symptoms (hours) after trauma(T)/ surgery(S)</th>
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A Case report and review of literature.


