Superior dislocation of the patella is a rare diagnosis. A 72 year old woman attended the accident and emergency department of the hospital with a painful right knee after a knock to the knee. Clinical examination and radiographs confirmed a superior dislocation of the patella, which was reduced and closed with the aid of simple analgesia. The authors present the case report and discuss the relevant literature.

A 72 year old woman attended the hospital with inability to bend her right knee associated with pain. She had been standing and reaching out to grab something three hours earlier, when she felt a sudden pain in her knee. The woman was initially seen in the accident and emergency department where a diagnosis of ruptured patellar tendon was made. When seen by us, she was indeed unable to perform active straight leg raise. However, examination of the knee showed that there was no palpable gap in the patellar tendon. The superior pole of the patella was projecting anteriorly and there was a prominent dimple below the patella.

Plain radiographs showed that the patella was superiorly dislocated with interlocking osteophytes at the inferior pole of the patella and anterior surface of femur (fig 1). After the administration of intramuscular analgesia, the patella was gently moved from side to side and a click was felt. After this the patient was able to actively straight leg raise and flex the knee through its full range. Reduction of the dislocation was confirmed by plain radiograph (fig 2).

The woman was mobilised fully weight bearing without any restriction and at review four weeks later was found to be asymptomatic.

DISCUSSION
Superior dislocation of the patella is a rare diagnosis. Excluding this case there have been 14 previous such cases reported in the literature.1–12 The average age of these patients is 58 years (range 43 to 81 years) and the ratio of male to female cases is equal. Two of these cases have been reported within the past year.1,2 As previously predicted, it may well be that the frequency of this condition is increasing given the degenerative nature of the underlying cause and the increasing elderly population in our society.3

The woman in our case reports that she was leaning over a chair to pick something from the floor when the chair slipped and pushed her patella up. This is in keeping with previous cases where the underlying mechanism has been reported as a low energy posteriorly directed force on the inferior pole of the patella with or without eccentric contraction of the quadriceps.4 Atraumatic cases in which the mechanism has been active quadriceps contraction and hyperextension of the knee have also been reported.1,3–5

Figure 1 Lateral radiograph of the knee showing superior dislocation of the patella.

Figure 2 Post-reduction radiograph.
Superior dislocation of the patella needs to be distinguished from patellar tendon rupture. Both conditions cause an inability to perform straight leg raise. However, in the case of superior dislocation, the patellar tendon is intact. Also the patella is invariably tilted anteriorly because of the locking osteophyte in superior dislocation of the patella with a characteristic dimple below the patella.

In all previous cases reduction was achieved closed with one exception, which required open reduction after failure of closed reduction. In the case presented reduction of the dislocation was achieved without the need for general anaesthesia or sedation. Previously, reduction required general anaesthesia in four cases. The rest were reduced with simple analgesia or sedation. Details of the reduction method are given in eight cases. This invariably entails gentle upward pressure on the inferior pole of the patella or medial-lateral pressure or a combination of both.

In summary this case highlights a rare case of superior dislocation of the patella. It is probable that this condition is increasing in frequency. It is important to distinguish this condition from patellar tendon rupture, which can be done by careful clinical assessment. After diagnosis, superior dislocation of the patella can be reduced closed with simple analgesia or sedation in most cases.

Contributors
Rashpal Bassi initiated the writing of the report and the literature search. B A Kumar supervised the writing of the report.

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REFERENCES

Haemodynamic and electrocardiographic consequences of severe nicorandil toxicity

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A 35 year old woman was admitted to the emergency department two hours after ingesting 60×20 mg tablets of nicorandil, total 1.2 g. The dominant feature of nicorandil toxicity was profound peripheral vasodilatation associated with coronary hypoperfusion. Despite widespread electrocardiographic signs of myocardial ischaemia, there was no evidence of myocardial damage and no serious cardiac arrhythmia. Volume loading and pressor support proved to be an effective treatment strategy.

The following day the patient continued to vomit and was febrile with a temperature of 38°C. She had a transient increase in the serum creatine kinase (peak 176 µmol/l) and liver transaminases (aspartate transferase 164 IU/l, alanine transferase 71 IU/l), which normalised by 48 hours. Serum potassium, calcium and creatine kinase remained within the normal range; troponin assay was not available and hence values were not measured.
The case of an arrhythmic overdose of nicorandil is described. Studies so far have suggested that the drug may potentially shorten the action potential duration, and be potentially pro-cardiac pre-conditioning and cardioprotection. As far as the authors are aware, this is the first description of the effects of nicorandil in severe overdose.

Nicorandil, a potassium channel activator, is the first agent in its therapeutic class to be licensed for the treatment of angina. Its oral bioavailability is 75%–80%, maximal plasma concentrations being achieved within 30–60 minutes after dosing, with a plasma half life of about one hour and therapeutic efficacy extending to 12 hours. Nicorandil is hepatically metabolised and renally excreted. Its mechanism of action is complex, and includes both a nitrate-like effect and in addition, activation of sarcolemmal and mitochondrial ATP sensitive potassium (KATP) channels. This leads to vasodilatation in coronary vessels and also in the venous and arterial systems, leading to a reduction in preload, afterload, and hence myocardial oxygen consumption and work. As a result of the above actions, nicorandil in double blind randomised studies, has been shown to be comparable as an anti-anginal agent to β blockers, calcium antagonists, and nitrates. In addition to its anti-ischaemic properties, nicorandil may also confer additional benefits related to myocardial pre-conditioning and cardioprotection.

The patient in this report had both symptomatic and ECG evidence of myocardial ischaemia, presumably as a result of coronary hypoperfusion. Volume loading and pressor support provided the most effective treatment strategy for reversing the effects of profound peripheral vasodilatation. It is interesting that in this case, despite ECG evidence of severe global myocardial ischaemia, there was no evidence of significant myocardial damage. Furthermore, it has been suggested that because of its mechanism of action, nicorandil could shorten the action potential duration, and be potentially pro-arrhythmic. However, studies so far have suggested this is not the case and, paradoxically, nicorandil may have anti-arrhythmic properties. Indeed, there is anecdotal evidence to suggest that nicorandil can shorten the QT interval and terminate torsades de pointes, and idiopathic VT. Certainly in this report of where almost 20 times the recommended daily dose was consumed, there was no evidence of sustained tachyarrhythmia.

Finally, the patient in this report experienced many of the reported side effects associated with nicorandil, most of which are attributable to its vasodilatory action. These include headache, flushing, dizziness, tachycardia, and hypotension. Other less common side effects have included oral aphthous ulceration, angioneurotic oedema, and photosensitivity. Although liver toxicity has been described, and elimination of nicorandil is largely dependent on hepatic metabolism, liver transaminases were only minimally increased in our patient. In summary, this is the first description of severe nicorandil overdose. The dominant feature of nicorandil toxicity was profound peripheral vasodilatation associated with coronary hypoperfusion. Despite widespread electrocardiographic signs of myocardial ischaemia, there was no evidence of myocardial damage and no serious cardiac arrhythmia. Volume loading and pressor support proved to be an effective treatment strategy.

Figure 1 ECG performed about six hours after overdose demonstrating widespread ischaemia with deep ST depression and T wave inversion.

was weaned off all inotropic support within 24 hours of admission and made a complete recovery. Post-discharge exercise stress testing using the standard Bruce protocol was uneventful.

DISCUSSION

The recently published IONA Study has shown a beneficial effect on major coronary events by the use of nicorandil in patients with chronic stable angina. As a result of this trial it is probable that the use of nicorandil in the treatment of angina will increase, which could also increase the potential for accidental or deliberate self harm with this agent. As far as we are aware, this is the first description of the effects of nicorandil in severe overdose. As a result of this trial it is probable that the use of nicorandil in the treatment of angina will increase, which could also increase the potential for accidental or deliberate self harm with this agent. As far as we are aware, this is the first description of the effects of nicorandil in severe overdose.

Nicorandil toxicity

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REFERENCES

Delayed diagnosis of foreign body aspiration in children

T Hilliard, R Sim, M Saunders, S Langton Hewer, J Henderson

Foreign body aspiration in children is common and usually presents with an initial episode of choking with subsequent respiratory symptoms. There may be cough, wheeze, or stridor, with decreased or abnormal breath sounds on examination. However, it can mimic other illnesses and cause difficulty in diagnosis. Radiological investigations may help to confirm aspiration but should not be used to exclude it. Three cases are presented of foreign body aspiration with a delay in diagnosis ranging from days to weeks. It is believed that delay could have been avoided with a more careful approach to the history and more appropriate use of investigations. These cases demonstrate that children with a history of choking and subsequent symptoms should be referred for bronchoscopy.

CASE REPORTS

Case 1

A 2 year old boy presented to the emergency department with wheeze and cough after a choking episode. A careful history and clinical examination can identify those children that need additional investigation including bronchoscopy. However foreign body aspiration can mimic other conditions and the link between choking and subsequent symptoms may not be made by parents and professionals alike. We present three cases with a delay in diagnosis, and discuss the appropriate management of suspected foreign body aspiration.

Case 2

A 2 year old boy developed noisy breathing after a choking episode while playing with a pistachio nut shell. The next day he had increasing difficulty in breathing and his general practitioner referred the child to the emergency department. He was thought to have acute asthma and was given nebulised bronchodilators, oral corticosteroid, and then an aminophylline infusion. After five hours of treatment there was little improvement and he continued to have noisy breathing, recession, and tachypnoea. A chest radiograph showed bilateral hyperinflation. On review he had biphasic stridor more likely to be attributable to upper airway obstruction. He underwent rigid bronchoscopy under general anaesthesia with removal of a pistachio nut shell from just under the vocal cords (fig 2).

Figure 1 Expiratory chest radiograph showing left sided hyperlucency.

Figure 2 Rigid laryngoscopy showing pistachio nut shell in trachea just below vocal cords.
Case 3
A 10 year old boy had a coughing fit while chewing on a pen top and subsequently realised that the inside of the pen top was missing. He was sent home from the local emergency department after a normal chest radiograph. Two days later he started to wheeze and cough. A chest radiograph was again normal, but he had wheeze that was louder on the right side of the chest and a fever. He was given intravenous antibiotics but he did not improve and the following day was transferred to the regional paediatric centre for assessment. With rigid bronchoscopy under general anaesthesia the pen top was removed from his right main bronchus.

DISCUSSION
Foreign body aspiration by children, especially those below the age of 3 years, is common.1 If it causes airway occlusion it may lead to asphyxia and it is unfortunately a leading cause of death in childhood.3 However, it more often presents with a history of an initial episode of choking and coughing with subsequent respiratory symptoms.1,2 These include cough, wheeze, stridor, or pneumonia. The most common physical sign is decreased or abnormal breath sounds.1,4 Most inhaled foreign bodies in children are food items, with peanuts being the most common.5

However, there is often significant delay until the diagnosis is made.1,4 In one series a delay of over three days between aspiration and removal of the foreign body was reported in almost 30% of children.1 This may be attributable to a high rate of initial alternative diagnoses and this occurred in 24% of cases in a separate series.6 Foreign body aspiration can be misdiagnosed as asthma, upper respiratory tract infection, pneumonia, or croup.1 Delay in diagnosis is associated with increased morbidity, especially respiratory infection.3

Most foreign bodies in children are radiolucent, but they may be associated with hyperinflation, atelectasis, or consolidation. In a series of 189 children with proven foreign body aspiration, 90 cases (47.6%) had normal chest radiographs.6 Inspiratory and expiratory films, and fluoroscopy can provide extra information, but even these may be normal in children who later are found to have an inhaled foreign body at bronchoscopy.1 Probably the most important feature of aspirated foreign bodies in children is the initial history of choking. In a series of 87 children who underwent bronchoscopy, a history of a choking episode was present in 67 of 70 with a foreign body and in only 4 of 17 without a foreign body.7 The choking history showed a sensitivity of 96% and a specificity of 76%. However, the episode may be un witnessed, or volunteered only after specific inquiry.1

When there is impaction of a foreign body in a major airway with acute respiratory distress and hypoxia the child should be resuscitated according to accepted guidelines.1 If urgent operative removal is required it should be carried out by the most experienced surgical and anaesthetic personnel available. However, when an emergency procedure is not indicated then transfer to a centre with regular experience of airway endoscopy in children should occur. In our centre we prefer to have available the option of both flexible and rigid bronchoscopy. If the diagnosis is in doubt flexible bronchoscopy can be used to examine more distal parts of the bronchial tree and more confidently exclude a foreign body than rigid bronchoscopy.5 However, flexible bronchoscopy plays little part in the extraction of foreign bodies.

Children who have a sudden onset of choking and coughing should be taken seriously. Most important is a thorough history of the initial episode and if there are persistent symptoms then the child should be referred for bronchosscopic evaluation.

Contributors
Tom Hilliard had the original idea, performed the literature search, and produced the draft manuscript. Richard Sim commented on the draft manuscript. Simon Langton Hewer and John Henderson were responsible for the management of the three cases and commented on the draft manuscript. John Henderson is guarantor for the article.

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REFERENCES
A 35 year old white man previously fit and well presented to the emergency department late one evening, with an episode of haemoptysis. He developed sudden left chest pain and breathlessness while having a drink in a local pub. The history obtained by the ambulance crew was that he coughed up half a litre of blood in the pub toilet.

On initial examination he was conscious but visibly distressed. He was noted to be tachypneic (respiratory rate 28 breath/min) and in severe pain over his chest and back. His initial pulse rate was 112 beat/min and his blood pressure was 141/93. The well healed scar of a thoracotomy was noted over the left side of his chest.

His partner who was present explained it was for the repair of a “hole in the heart” 23 years ago, when he was aged 12. He had been on follow up for a few years after the surgery by the tertiary cardiac centre and then discharged to his primary care practitioner. The last chest radiograph he had was more than 10 years ago. He had otherwise been fit and well and participated actively in sport.

The patient had 100% oxygen administered by non-rebreathing mask with a reservoir bag and intravenous access was obtained with wide bore cannulas. Blood was taken for grouping and cross matching.

Despite high flow oxygen the best saturation obtained was 89% by pulse oximetry. A blood gas sample taken showed marked hypoxia and mild acidosis (Table 1). Intravenous morphine was administered with temporary relief of pain requiring repeated titrated doses. An ECG showed sinus tachycardia. An urgent portable chest radiograph was done (Fig 1).

His condition rapidly deteriorated and he collapsed from torrential bouts of haemoptysis requiring endotracheal intubation, emergency blood, fluids, and cardiopulmonary resuscitation. He failed to respond to advanced life support resuscitation and was pronounced dead.

A postmortem examination showed an 8 cm thin walled aneurysm of the arch of aorta and proximal descending thoracic aorta that had ruptured into the upper lobe of the left lung and pleural cavity.

DISCUSSION

Haemoptysis as a presenting symptom of leaking aortic aneurysm after coarctation repair has been previously described in the literature. Aortic patch graft repair was first performed in the United Kingdom by Charles Drew in Westminster Hospital half a century ago.2 Between 1976 and 1982 patch aortoplasty was the routine procedure of choice.7 Aneurysm formation rate after the repair has been described in long term follow up studies (up to 15 years after surgery) to be between 3.8% and 27%.4-6 Our patient presented 23 years after his surgery with a ruptured aneurysm.

Some 32.8% of patients in one study who had patch graft aortoplasty for aortic coarctation underwent reoperation because of aneurysm formation at the site or opposite to the patch graft.7 The argument for lifelong radiological surveillance by plain radiology (chest radiograph), Doppler ultrasound, or computed axial tomography for asymptomatic patients with previous coarctation patch repair is very strong indeed from these reports. This can be along the lines of treatment for patients diagnosed to have early abdominal aneurysms.

General practitioners who have such patients on their lists are ideally suited to undertake this role and should review them on a regular basis with screening surveillance radiography.

In conclusion, vascular events like aneurysm ruptures and dissections are by their nature often catastrophic if they present late in the their clinicopathological course. They are literally like “ticking time bombs” pulsing to our heart rhythm. Early recognition of possible predisposing factors like age, atherosclerosis, hypertension, and previous risk related surgery (for example, aortic surgery) entails us to be vigilant in our follow up of the patient at risk of premature death and significant morbidity.

Contributors

Dilip Menon did the literature search, wrote the manuscript of the case report, and is guarantor for the paper. Tony Burdge did the postmortem examination and report and reviewed the manuscript of this case report.

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Competing interest: the first author was directly involved in the initial care of the patient.
First branchial cleft anomaly presenting as a recurrent post-auricular abscess

M A Siddiq

15 year old boy presented with a two year history of a recurrently infected cyst behind his right ear. He had initially presented to the accident and emergency (A&E) department with an infected swelling behind his right ear. This was treated in the department with incision and drainage under local anaesthetic. It recurred six months later and another presentation to the A&E department with an infected swelling behind his right ear. This lesion recurred again three months later and was treated in the A&E department with incision and drainage under local anaesthetic. This lesion recurred again three months later and was treated in the A&E department with incision and drainage under local anaesthetic. The lesion had become infected on two further occasions but had settled with a course of antibiotics from the general practitioner.

On presentation to the ear, nose, and throat clinic, there was a cystic swelling in the right post-auricular region inferiorly, with evidence of some scarring (fig 1). Examination of the external auditory meatus revealed the presence of a pit on the postero-inferior wall. The remainder of the ENT examination including examination of the neck was normal. A provisional diagnosis of a first branchial cleft anomaly was made. Computed tomography with contrast was performed, which showed the extent of the lesion with an associated tract running along the posterior wall of the external auditory meatus.

The patient underwent an exploration of the lesion, which confirmed the presence of a sinus communicating between the pit seen on examination and the cyst, thus confirming the diagnosis. Histological examination revealed the tract to be lined with stratified squamous epithelium thus establishing the ectodermal nature of the lesion.

DISCUSSION

First branchial cleft anomalies are uncommon and comprise 1%–8% of all branchial cleft anomalies. They often present in the first two decades of life and present a clinical challenge as they can easily be misdiagnosed and thus inappropriately treated. They are thought to arise as a result of developmental abnormalities of the branchial apparatus and may take the form of a cyst, sinus, or fistula.

Clinically they may present with repeated episodes of infection of the lesion. This may manifest itself with a cystic swelling or discharge from a fistulous opening either pre-auricularly or post-auricularly, in the cheek, or high in the neck. A thorough otological examination should be performed in all cases and may reveal a pit visible in the external canal at the site of entrance of a sinus or fistula. Such a lesion may result in otorrhoea or oitis externa with infective exacerbations. The meatus may be found to be partially obstructed by bulging of the canal wall because of a cystic swelling. Such patients may complain of hearing loss, as may those with oedema associated with an oitis externa. There may however be a complete absence of signs in the external auditory canal. The patient may also give a history of having to repeatedly undergo incision and drainage of an apparent...

REFERENCES


Scrotal pain in the absence of torsion; need for vigilance

S Venketraman, J P Gray, P A Evans

Epididymitis is a common presentation of acute testicular pain seen in the emergency department, the differential diagnosis being testicular torsion. The vast majority of young men with epididymitis have an infective aetiology and this settles with antibiotic treatment. The clinical course of a patient who presented with testicular pain is described. At ultrasonography, the patient was found to have the uncommon condition of testicular microolithiasis, a condition that has been linked to malignant disease. Emergency doctors should be aware of the potential consequences of returning scrotal pain consistent with epididymitis to the community on antibiotic treatment alone. All patients with probable epididymitis should have either a scrotal ultrasound or specialist follow up.

A 22 year old white man presented to the emergency department of the Leicester Royal Infirmary complaining of an increasingly painful left testicle, after mild blunt scrotal trauma sustained five days previously. The pain had not been alleviated by the counter non-steroidal analgesia. He described no urinary symptoms and had not had intercourse for some months. He gave a history of chronic bilateral testicular tenderness and had been treated for epididymitis in the past.

On examination, he was afebrile. Scrotal examination revealed a very tender left epididymis and a mildly tender right epididymis, clinically inconsistent with torsion. Urine analysis was negative. An ultrasound scan of the scrotum was arranged from the emergency department, which was reported as a bilateral epididymitis and bilateral testicular microlithiasis with no haematomata or infarct seen (fig 1). The patient was discharged with antibiotic treatment and urology outpatient follow up.

DISCUSSION

Epididymitis itself is not an uncommon emergency department presentation. However, because of a lack of epidemiological data the actual incidence is unknown. Symptoms are usually unilateral and patients generally present with testicular pain accompanied by a tender swollen epididymis. They may present with dysuria, fever, scrotal erythema, and orchitis. Epididymitis is mostly infective in aetiology, with Chlamydia trachomatis and Neisseria gonorrhoeae being the two most common sexually transmitted organisms. From an emergency department perspective: the important diagnosis to exclude is testicular torsion. Simultaneous bilateral epididymitis as in this case is rare. In a review of 610 cases, bilateral epididymitis was noted in only 9% and in a proportion of these
the initial presentation was unilateral and thereafter developed bilaterally.¹

First documented in the 1960s as microcalcification within the lumina of seminiferous tubules, testicular microlithiasis is a rare diagnostic entity.² As an essentially asymptomatic pathology, the prevalence in the male population remains unknown. Radiological studies assessing the incidence on ultrasonography have quoted figures ranging from 0.16% to 4%.³ Specific postmortem investigation to assess the incidence suggests a figure of 4% for adult men.³ Electron microscopy in the 1980s confirmed these deposits as consisting of a central calcified core surrounded by cellular debris, glycoprotein, and collagen.⁴ On ultrasound the deposits are less than 2 mm in diameter, hyperechoic, and tend not to cast an acoustic shadow. Reported associations include cryptorchidism, Kleinfeilter’s syndrome, infertility, testicular pain, and testicular neoplasm.⁵ It has therefore been suggested that testicular microlithiasis should be considered a premalignant condition, and patients should have clinical and ultrasonographic follow up.⁶

As far as we are aware, there is no previous published case of bilateral epididymitis and bilateral microlithiasis occurring simultaneously in the same patient. This case reports standard emergency department management of acute testicular pain, following a pathway geared to excluding a diagnosis of testicular torsion. Once excluded, many patients are discharged from the emergency department with antibiotic treatment and often no specialist follow up.

This case highlights the importance of vigilance among emergency clinicians with regard to the painful scrotum in the absence of testicular torsion. Indeed, the take home message for us to learn here is that a clinical diagnosis of epididymitis without a scrotal ultrasound and or urology follow up is potentially hazardous.

Contributors
Shakthi Venketraman and Jim Gray realised this case merited reporting and conducted a literature search. Philip Evans edited the paper, stands guarantor, and oversaw the project. All authors contributed to writing the case report.

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REFERENCES
Unusual presentation of atrial fibrillation

P Muthu, G Oduro, M Sakr, D A Esberger

A case is reported of atrial fibrillation in a young healthy man after head injury and the possible causes are discussed. The atrial fibrillation reverted spontaneously to normal rhythm in two days. The authors are not aware of a similar report in the literature.

A 33-year-old male police officer was brought to the accident and emergency department after being found unconscious in the street. The exact mechanism of injury was not known, though the paramedics had obtained the history from bystanders that the patient had been trying to apprehend a suspect. He was confused and disoriented and could not remember how he had sustained his injuries. He was complaining of headache and was vomiting repeatedly. He was not complaining of neck pain or chest pain.

On examination his airway, breathing and circulation were normal. His Glasgow Coma Score was 13/15. His pupils were equal and reacting to light. His pulse was 96/minute and blood pressure was 136/60 mm Hg. His cardiovascular system, respiratory system, and abdomen were normal. Bleeding from the left ear and swelling and tenderness over the nasal bones was noted. No chest injury or any other injuries were seen. A clinical diagnosis of basal skull fracture and fracture of nasal bones was made.

There was no significant past medical history particularly in relation to the cardiovascular system and he was not taking any medication. He smoked 10 cigarettes a day and consumed 10 units of alcohol per week. Radiographs of the cervical spine, chest, and pelvis were normal. An ECG showed atrial fibrillation (fig 1). A CT scan of the head revealed no abnormality. Full blood count, urea, and electrolytes were normal.

He was admitted for neurological observations and during the review on the following day he was able to recollect the events and remembered chasing and apprehending a suspect and escorting him to the police vehicle. He was not able to recall details of the incident subsequently and remembered being in the accident and emergency department. He was also reviewed by the medical team in view of his atrial fibrillation.

Figure 1  ECG showing atrial fibrillation.

Figure 2  Normal ECG after 48 hours.
Atrial fibrillation is one of the most common arrhythmias seen in accident and emergency departments. It is most commonly attributable to medical disorders and usually occurs in the elderly population. It is associated with detectable organic heart diseases in about 70% of patients and may also arise secondary to severe chest trauma. There are various reports in the literature regarding uncommon aetiologies of atrial fibrillation such as electrical injury, high protein diet, high alcohol intake, excessive exercise, electroconvulsive therapy, hypoglycaemia, sigmoidoscopy, and dental extraction. A patient with newly diagnosed atrial fibrillation warrants a full investigation of the aetiology of this common arrhythmia because it may be associated with unusual pathology. It has been reported in the literature that cardiac rhythm disorders can occur after head injury as well as after cerebral stimulation.

There are various pathophysiological mechanisms that may cause atrial fibrillation. It may be attributable to morphological changes in the heart such as acute or chronic stretch or possible changes in the cellular electrophysiology. It may also arise from disorders of autonomic tone with several studies emphasising the importance of autonomic nervous system in the initiation and perpetuation of atrial fibrillation.

We considered the possibility of three causes for the atrial fibrillation in our patient such as paroxysmal atrial fibrillation attributable to an autonomic disorder, blunt chest trauma, and head injury with basal skull fracture.

Vagally mediated atrial fibrillation occurs more frequently in young healthy men. The age of onset is usually between 30–50 years and it occurs in subjects with normal heart where vagal influence predominates. Atrial fibrillation usually occurs at night, and reverts to normal sinus rhythm in the morning. It is not triggered by physical exertion and emotional stress. However, the relaxation that follows physical efforts or emotional stress is frequently associated with the onset of atrial fibrillation. This seems to be consistent with the occurrence of atrial fibrillation in our patient, but we ruled out this aetiology as the atrial fibrillation lasted for two days.

We considered the possibility that our patient might have fallen on his radio and sustained chest trauma, which would have caused the atrial fibrillation. However, there was no evidence of injury to the chest. Furthermore, the most common arrhythmias after myocardial contusion are sinus tachycardia, supraventricular tachycardia, atrial or ventricular premature contractions, conduction disorders, and non-specific ST segment and T wave changes. Atrial fibrillation after chest trauma is very rare and reported only in elderly patients. Large studies of patients with chest trauma or sternal fractures found no cases of isolated atrial fibrillation particularly in young patients. It is theoretically possible that a heavy blow to the anterior chest, timed appropriately in the cardiac cycle, could propagate an ectopic impulse, precipitating an atrial or ventricular dysrhythmia. However, its applicability to humans is unclear, as ectopic rhythms are distinctly less common in clinical practice. Therefore, it is not likely that atrial fibrillation in our patient was attributable to the chest trauma.

Recent research on the pathology of head injury has focused on the changes occurring at cellular level in the first few hours after head injury. Trauma is hypothesised to produce widespread depolarisation of neurons and excessive release of excitatory neurotransmitters, which cause excitotoxic effects on postsynaptic neurons. In experimental animal studies, Mauck et al noted that stimulation of the distal cut end of the right vagosympathetic trunk evoked bradycardia with hypotension, and in one instance, a brief run of atrial fibrillation. There were no other arrhythmias. It has also been reported that sub-arachnoid haemorrhage causes a stress response with increased concentrations of plasma catecholamines and serious cardiac arrhythmias.

Atrial fibrillation in young, healthy patients without pre-existing heart disease may account for up to one third of all cases. It may be attributable to physiological stress of trauma such as hypovolaemia, acidosis, electrolyte imbalances, and excessive catecholamine release. Alcohol ingestion may promote atrial fibrillation by increasing catecholamine release or by increasing the vagal outflow because of the associated nausea and vomiting. It may also be attributable to thyrotoxicosis and pneumonia. Idiopathic or lone atrial fibrillation may also be considered in younger patients with atrial fibrillation.

This case illustrates that atrial fibrillation may be detected in otherwise fit young patients with an isolated head injury and no other obvious precipitating factors. It is very important that full investigations are carried out to exclude other causes before atrial fibrillation is attributed to a head injury. However,
it is not always possible to determine the exact cause of atrial fibrillation.

Contributors
PM initiated the idea, did the literature search, and wrote the paper. GO helped in the literature search and writing the paper. MS and DAE involved in the management of the patient and helped in writing the paper. PM acts as the guarantor of the paper.

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