High-grade glioma mimicking acute viral encephalitis – three case reports

J H Rees, R S Howard

Summary
The clinical features of viral encephalitis consist of headache, fever, seizures and encephalopathy. We report three patients with high-grade gliomas presenting with encephalitic illnesses. The diagnosis of brain tumour should always be borne in mind if definite evidence for a viral infection is not obtained.

Keywords: glioma; viral encephalitis; encephalopathy

Viral encephalitis is characterised by headache, confusion, altered consciousness with or without seizures, and fever. The most commonly identified cause of sporadic community-acquired encephalitis is herpes simplex virus (HSV) type I, which may be diagnosed on viral culture of cerebrospinal fluid (CSF), polymerase chain reaction (PCR) analysis looking for HSV DNA, serological testing for IgM antibodies, or brain biopsy. Characteristically the virus attacks the medial temporal lobes and insular cortex giving rise to a focal encephalitis which carries a significant mortality rate. Rarely, the patient presents with symptoms and signs of raised intracranial pressure, giving rise to the possibility of a temporal lobe abscess. The converse situation, whereby a high-grade tumour presents as an acute encephalitic illness, has only rarely been reported. The following three cases illustrate that this diagnosis should always be borne in mind if definite evidence for a viral infection causing the encephalopathy is not obtained, especially if the PCR test is negative for the presence of viral DNA.

Case reports

Case 1
A 41-year-old woman presented with tonic-clonic seizures, fever, drowsiness and confusion. Magnetic resonance imaging (MRI) showed extensive high signal abnormality in the left temporal lobe and posterior parietal region (figure 1). CSF was normal. A presumptive diagnosis of herpes simplex encephalitis (HSE) was made and she improved after intravenous acyclovir. However, her seizures persisted and 2 years later her MRI scan showed an increase in the size of the temporal lesion (figure 1). Approximately one month later, she was found unconscious at home. A CT scan now showed a large lesion with considerable midline shift. Emergency decompression was unsuccessful and she died without regaining consciousness. The histology of the lesion was glioblastoma multiforme.

Figure 1 Coronal T2-weighted MRI scans showing an area of high signal in the left medial temporal lobe which was identical to the images on presentation 2 years earlier. Two years later and one month before the patient’s death, the abnormality is more extensive and involves the inferior parietal lobe

Case 2
A 49-year-old man became acutely confused and then had serial tonic-clonic seizures. On admission he was pyrexial, tachycardic and unconscious. MRI showed extensive high T2 signal change affecting the right temporal and frontal lobes with some mass effect (figure 2). CSF examination including PCR for HSV DNA was negative. He was started on acyclovir and gradually improved. He had persistent simple partial seizures and a repeat MRI 3 weeks later showed no change. A stereotactic brain biopsy was carried out and revealed an anaplastic astrocytoma. He was treated with external beam radiotherapy followed by combination chemotherapy.

Case 3
A 72-year-old woman presented with a 2-week history of episodes of vagueness and confusion. MRI revealed a lesion involving the grey and white matter of the left temporal lobe but also a similar process involving the right temporal lobe and uncus (figure 3). There was some mass effect and mild contrast enhancement. It was felt that the appearances were more consistent with HSE than glioma. A CSF sample taken 5 days after starting acyclovir was normal, including PCR for HSV DNA. She remained persistently confused with frequent simple and complex partial seizures. One month later MR guided stereotactic biopsy of the left temporal lobe revealed a high-grade astrocytoma which was not treated at the request of her family.
Discussion

All three patients described above presented acutely with confusion and seizures. Fever was present in the first two cases and all patients had extensive areas of high signal on T2-weighted MR imaging within one or both temporal lobes. HSE was therefore diagnosed on both clinical and radiological grounds. Results of CSF examination were normal in each case, but as this was not incompatible with HSE, the patients were treated with a full course of intravenous acyclovir. No patient received steroids at presentation and the improvement in patients 1 and 2 presumably occurred as a consequence of achieving seizure control. It was only when the results of the PCR came back as negative that the initial diagnosis was reconsidered. In cases 2 and 3 a high-grade temporal lobe glioma was diagnosed using CT or MR-guided stereotactic biopsy within one to two months of the original presentation. The first patient, however, remained undiagnosed for over 2 years and biopsy was not carried out principally because the MRI scan appearances of the lesion had not obviously changed. Her final MRI scan showed that the tumour had increased in size but the decision to biopsy electively was never made as she presented soon after in extremis. Presumably the tumour had undergone transformation from low-grade to high-grade in a short space of time.

The distinction between acute viral encephalitis and a glioma is usually easily made on clinical and radiological grounds. PCR can be used to identify viral DNA in the CSF and allows early diagnosis of HSE. Occasionally the presentation is atypical and HSE may mimic a rapidly expanding temporal lobe mass, giving rise to considerable diagnostic difficulty. However, the converse situation whereby a malignant glioma presents as an acute encephalopathy is much rarer.

This ‘encephalitic’ presentation of temporal lobe tumours has been previously alluded to in a study of 432 patients who underwent brain biopsy for presumed HSE, in the pre-MRI era. Five out of 95 (6%) patients who were biopsy-negative for HSE had tumours, two metastatic colonic adenocarcinomas, one primary CNS lymphoma and two glioblastomas. In another study looking at the differential diagnosis and outcome of 65 patients presenting with an acute encephalopathy over a 17-year period, only one patient out of 34 in whom a definite or probable diagnosis could be made had a tumour. This was a 45-year-old man presenting with a short history of headache and confusion who had a right temporal lobe lesion. Three years later a biopsy revealed an oligodendroglioma. This very low frequency suggests that a primary CNS tumour rarely presents as an encephalitic illness. With the advent of PCR analysis brain biopsy is seldom undertaken to diagnose HSE, particularly as the most common differential diagnoses include other viral encephalitides for which no specific treatment is currently available. However, we believe that stereotactic biopsy should be considered in all patients with temporal lobe mass lesions if no definitive diagnosis is made after the results of PCR analysis for common viruses are available.

Learning points

- A malignant brain tumour can present acutely as fever and encephalopathy with bilateral abnormalities on MRI scanning.
- Stereotactic biopsy should be considered in patients with temporal lobe mass lesions if no definitive diagnosis is made after the results of PCR analysis for common viruses are available.
Hypertensive encephalopathy in a patient with retroperitoneal fibrosis

D Das, J Brigg, C M Brown

Summary
A patient presented with retroperitoneal fibrosis but without any ureteric obstruction. The diagnosis was made by an abdominal CT scan and also at laparotomy. Post-operatively, she developed hypertensive encephalopathy. An isotope renogram with captopril was abnormal but not diagnostic of renal artery stenosis. The patient’s condition improved with steroid and antihypertensive treatment. A follow-up CT scan showed complete resolution of peri-aortic thickening. A causative link is postulated between retroperitoneal fibrosis, trauma during laparotomy, and onset of acute hypertension.

Keywords: retroperitoneal fibrosis; hypertensive encephalopathy

Retroperitoneal fibrosis, first described by Ormond in 1948, characteristically affects the peri-aortic tissues often involving the ureters, leading to ureteric obstruction. The pathogenesis of this disease, though not fully understood, is now recognised to be immune mediated. Although no therapeutic trials have been conducted because of low incidence of the disease, steroid treatment has been found to be effective in most cases. Patients often present with non-specific symptoms and occurrence of hypertension has been rarely reported in these patients. The present case report is related to a patient with retroperitoneal fibrosis complicated by hypertensive encephalopathy, the first case report of its kind. We discuss the possibility of a causal relationship between these two conditions and the role of steroid treatment in preventing such complications.

Case report
A 50-year-old woman presented with a 3-month history of anorexia and weight loss of 14 lbs. She had undergone hysterectomy and bilateral salpingo-oophorectomy several years earlier and was subsequently put on hormone replacement therapy. She specifically denied previous exposure to ergot, methysergide or practolol. Blood pressure, fundoscopy and a resting electrocardiogram (ECG) were all found to be normal at initial screening. On abdominal palpation, a para-umbilical mass was noted but without any evidence for hepatosplenomegaly or ascites. Full blood count and electrolytes were normal but erythrocyte sedimentation rate (ESR) was raised to 94 mm/h. Auto-immune screen for antibodies was negative and immunoglobulin levels were normal. Abdominal computed tomography (CT) scan showed a fair amount of soft tissue surrounding the aorta from the level of the renal vessels to the distal common iliac arteries on both sides of the pelvis with no abnormality of the other intra-abdominal organs (figure 1).

A laparotomy was performed to exclude lymphoma but extensive thickening of the aorta, superior mesenteric, renal and common iliac arteries was noted. Histology of tissue dissected from the peri-aortic region showed predominantly fibrous tissue infiltrated with plasma cells and lymphocytes, the latter often aggregated in places in the form of follicles.

Figure 1 Post-contrast CT scan of the abdomen showing thickening of the aortic wall due to extensive fibrosis
On the second postoperative day her blood pressure progressively rose to 230/120 mmHg, precipitating a generalised convulsion. CT scan of the brain demonstrated oedema over the vertex region with effacement of the cortical sulci but no haemorrhage. The total volume of intravenous fluid replacement in the postoperative period was only 2 litres. 24-Hour urinary catecholamine levels were normal. Systemic blood pressure was controlled with 50 mg intravenous labetalol followed by oral prednisolone (40 mg) and bendrofluazide (2.5 mg daily). An isotope renogram showed no initial abnormality but reduced bilateral renal function following a single dose of captopril. The classical delay in clearance, associated with renal artery stenosis, was not noted. More than a year after discharge from the hospital the patient remains well with normal blood pressure and an ESR of 5 mm/h. A repeat CT of the abdomen (figure 2) showed resolution of the inflammatory oedema and fibrosis around the aorta, seen in the previous CT. Her current treatment comprise atenolol 50 mg, nifedipine 20 mg and prednisolone 5 mg.

**Discussion**

The clinical condition of retroperitoneal fibrosis, presenting as ureteric obstruction and periaortiearcal thickening of the aortic wall has long been recognised, but wider use of CT scanning detects more asymptomatic cases. This condition is thought to be secondary to leakage of material from atheromatous plaque in the diseased aorta, generating an autoimmune inflammatory response and the presenting symptoms depend on the organs or vascular structures involved.

Mild hypertension has been previously reported with this condition, but no studies have been carried out to determine a causal relationship. However, hypertensive encephalopathy, manifest by cerebral oedema and a generalised convulsion has not been previously reported. The precise reason for severe hypertension in this woman is subject to speculation, but may involve inflammatory narrowing of the renal arteries, as evidenced by the abnormal post-captopril isotope renogram, with further local tissue oedema resulting from retroperitoneal biopsy at laparotomy. Steroid treatment may have controlled the inflammation improving the patency of the renal arteries. Our patient improved with steroids and antihypertensive therapy and could not be persuaded to undergo renal angiography.

This is the first case report documenting occurrence of hypertensive encephalopathy following operative biopsy in a patient with retroperitoneal fibrosis. The possibility of such a complication should prompt consideration of steroid treatment before invasive procedures or laparotomy is undertaken.

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**Figure 2** Post-contrast CT scan of the abdomen, 1 year after institution of steroid treatment, showing resolution of inflammatory oedema and fibrosis around the aorta.
Visceral leishmaniasis masquerading as tuberculosis in a patient with AIDS

Amitabh Yaduvanshi, Meenakshi Jain, S K Jain, Shyama Jain, Suneet Arora

Summary
We report a case of visceral leishmaniasis presenting as significant lymphadenopathy in a patient with acquired immune deficiency syndrome. The lymphadenopathy was initially suspected to be tubercular in nature on pathological examination. This report highlights the increasing incidence of acquired immune deficiency syndrome and Leishmania co-infection in India, and the importance of demonstrating tubercle bacilli on culture before suggesting a diagnosis of tuberculosis.

Keywords: leishmaniasis; AIDS; tuberculosis

Visceral leishmaniasis (VL) is being increasingly recognized in human immunodeficiency virus / acquired immune deficiency syndrome (HIV/AIDS) patients. It occurs in epidemic form in Eastern India. Given the endemicity of VL in this area, physicians can expect to see VL in patients with HIV/AIDS. Resistance to sodium antimony gluconate, the paucity of alternative drugs such as pentamidine and amphotericin B, and woefully inadequate vector control measures in VL, make the situation positively explosive in developing countries. To the best of our knowledge there are only two case reports of HIV and VL co-infection from India, but this is the first report in which the patient presented with predominant lymphadenopathy, with no suspicion of HIV or leishmaniasis.

Case report
A 30-year-old man from Eastern Uttar Pradesh (in the VL belt in India) presented with moderate-grade intermittent fever associated with chills for the last 7 months. He also complained of swelling in the neck, axillary and inguinal regions for the past 6 months, and associated complaints of anorexia, weight loss and malaise. He had no other symptoms. He had undergone surgery for varicose vein removal 8 years earlier, during which he had received multiple units of blood. There was no history of any other high-risk behaviour.

He had been treated for tuberculosis for the past 6 months based on an inguinal lymph node fine needle aspiration cytology (FNAC) diagnosis of granulomatous lymphadenitis, possibly of tubercular nature. The patient was started on antitubercular drugs (rifampicin, isoniazid, ethambutol, pyrazinamide) reduced to two drugs after three months (RH). However there was no improvement in the patient's symptomatology.

Physical examination revealed generalised lymphadenopathy of 2–3 cm in size, firm, non-tender and non-matted. Pallor, cachexia, herpes simplex labialis and oral thrush were present along with a 3-cm hepatomegaly and a 5-cm splenomegaly. The rest of the systemic examination was normal.

Investigation revealed haemoglobin 8.7 g/dl, total leucocyte count 3200/µl with 70% polymorphs and 30% lymphocytes, platelets 99 × 10^10 g/l, erythrocyte sedimentation rate 75 mm/1st hour. Other investigations were within normal limits. Ultrasonographic examination revealed hepatosplenomegaly with abdominal lymphadenopathy.

A review of the previous FNAC revealed non-caseating granuloma (figure 1); acid-fast bacilli (AFB) staining was not done. At the same time lymph node biopsy from axillary lymph nodes showed non-specific dermatopathic changes. Mantoux test and ELISA for tuberculosis IgG and IgA were within normal limits. Bone marrow aspirations did not reveal any haemoparasites.

ELISA for HIV was positive and later confirmed by Western blot technique. Repeat FNAC from inguinal lymph node revealed numerous intracellular and extracellular amastigote forms of Leishmania donovani (LD bodies) (figure 2). The CD4 count (helper/inducer) was 2% and absolute CD4 count was 30/µl.

A final diagnosis of AIDS stage IVC with VL (kala azar) was made. The antitubercular drugs were withdrawn and the patient was started on intramuscular (im) sodium stibogluconate (20 mg/kg body weight/day), continued for 28
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from India. The possible reasons for the scarcity of cases from Southern Europe. However, this is only the third case of HIV/VL co-infection reported from India. The patient had lymphadenopathy as the predominant presentation, which is uncommon in Indian VL, and the lymphadenopathy had previously been misdiagnosed as tubercular in origin.

Tuberculosis is highly prevalent in India and is a common cause of lymphadenopathy. Due to lack of resources, the patient is usually started on antitubercular drugs on the basis of the cytohistological appearance, without resorting to AFB staining or culture. This strategy is generally helpful, as other granulomatous diseases are uncommon in India. However, lymphadenopathy and granuloma formation are rarely seen in VL. In response to the initial infection there is a migration of histiocytes into the lymph nodes, resulting in the formation of non-caseating granuloma. These granulomas may sometimes be thought to be tubercular in origin due to the paucity of LD bodies in the early stages, as happened in our patient. There is then reactivation of this latent infection when the patient is stressed or immunocompromised in any way. At this stage the granuloma and histiocytes are disrupted and numerous LD bodies are seen on cytology or histology, as was the case in our patient whose repeat FNAC showed numerous LD bodies.

In a recent WHO communication, 692 retrospective cases of HIV/VL co-infection were reported; 97.3% of these cases originated from Southern Europe. However, this is only the third case of HIV/VL co-infection reported from India. The possible reasons for the scarcity of cases from India are, firstly, that the seropositivity of HIV in India is restricted to the urban areas while VL has a predominantly rural and suburban distribution. A study from Bihar, the state with the highest incidence of VL, did not report a single HIV-positive sample from 4567 sera samples obtained from high-risk individuals, showing the low prevalence of HIV infection in rural and semi-urban societies. Secondly, in a developing country like India, other virulent infections are present in the environment (eg, tuberculosis, bacterial pneumonias, etc) and patients suffering from HIV infection usually die from such infections before they are immunocompromised enough to succumb to opportunistic infections like VL.

However, all this may change in the near future, with rapid liberalisation and economic growth, there has been a greater intermixing of populations from urban and rural areas, and due to better medical care and awareness, patients with HIV/AIDS are surviving longer. Thus India is on the verge of breaking through into the circle of developed countries and it will be facing unique health problems common both to underdeveloped and developed countries in the coming decades.

There is a sharp reduction in mean survival (13 months) in patients with Leishmania/HIV co-infection compared to other AIDS patients. There are also reports of frequent severe VL caused by Leishmania strains with lower or no virulence in HIV-negative individuals, and it has been suggested that LV should be included among the AIDS-defining diseases. The WHO has recommended treatment with
sodium stibogluconate (20 mg/kg/day for 28 days) as first-line therapy. Pentamidine at a dose of 4 mg/kg on alternate days for 3–4 weeks, sodium stibogluconate and allopurinol (10 mg/kg/day), and amphotericin B (0.5 mg/kg/day for 14–21 days) have been used as second-line drugs in HIV/VL co-infected patients, but have been associated with relapse, unresponsiveness and low efficacy. More recently, amphotericin B lipid complexes (AmBisome) have been found to be more effective and less toxic in immunocompromised VL patients. Pentavalent antimony given once a month was effective in prevention of relapse in HIV-infected leishmania patients after initial cure.

4 Vishwas LA, Singh S, Wali JP. Visceral Leishmaniasis in AIDS. J Assoc Physicians India 1997;45:582.

Fatal phenytoin hypersensitivity syndrome

U Mahadeva, M Al-Mrayat, K Steer, E Leen

Summary

The phenytoin (hydantoin) hypersensitivity syndrome is rare but potentially fatal. Often, as in this case, it presents with nonspecific symptoms and signs, requiring a high degree of clinical suspicion for diagnosis.

Keywords: phenytoin; hydantoin; hypersensitivity.

Phenytoin (diphenylhydantoin) is an aromatic ring compound which is metabolised in the liver by hydroxylation of one of its phenyl groups and then excreted in the urine and bile as a glucuronide conjugate. It was first introduced as an anticonvulsant in 1938. Dose-dependent side-effects (neurological impairment, gingival hyperplasia and megaloblastic anaemia) soon became apparent but an idiopathic hypersensitivity syndrome to phenytoin (and other hydantoins) was first described in 1959.1

We report a case of unrecognised, fatal, phenytoin hypersensitivity syndrome with characteristic pathological changes in the heart, liver, skin and lymph nodes at autopsy.

Case report

The patient was an 85-year-old AfroCaribbean woman who was admitted following collapse at home, with drowsiness, fever, rigors and a pruritic rash. Her medical history included a hysterectomy, ischaemic heart disease, type II diabetes mellitus and paraphrenia. One month prior to admission she was diagnosed as having complex partial seizures and was commenced on phenytoin 300 mg at night. Her other medications included metformin 500 mg bid, tolbutamide 500 mg bid, nifedipine 10 mg bid and risperidone 1 mg od. She was a non-smoker and had stopped taking alcohol since the onset of the fits.

On examination she was unwell and confused with a temperature of 38°C, regular pulse rate of 90 beats/min and a blood pressure of 120/70 mmHg. She had a generalised maculopapular erythematous rash, marked neck stiffness but no photophobia or focal neurological signs. She had a systolic murmur at the apex. Her chest auscultation was normal. Abdominal examination was unremarkable.

The initial working differential diagnoses were septicemia, a drug reaction, or possible food allergy (the patient was allergic to salmon and she had consumed some one day prior to admission).

The results of initial investigations are presented in box 1, the most notable abnormal findings were a raised blood eosinophil count and C-reactive protein; a markedly elevated creatine kinase; and grossly deranged liver function tests. Skin biopsy was not performed.

She was treated empirically with intravenous antibiotics, corticosteroids and antihistamines. All her usual medication apart from the phenytoin was discontinued over a period of 3 days. Phenytoin was completely stopped on day seven and benzodiazepines were used to control any seizures. Unfortunately she developed renal, hepatic and cardiac failures and generalised skin desquamation. She deteriorated rapidly and died 12 days after admission.

At post-mortem examination there was florid desquamation of the entire skin surface. Internal examination revealed heavy lungs (right 520 g, left 420 g), a large, yellow liver (1640 g), a slightly enlarged spleen (140 g) and widespread soft, fleshy lymphadenopathy (up
to 3 cm in diameter). The heart (270 g) appeared normal. The brain (1000 g) was small and firm and showed yellowish discolouration of the cortex suggestive of amyloidosis. Histological examination of the heart showed an interstitial eosinophil-rich infiltrate with myocytolysis. There was no necrosis, vasculitis or fibrosis (figure). These features are characteristic of hypersensitivity myocarditis.2 The liver displayed a marked periportal acute and chronic inflammatory infiltrate including increased numbers of eosinophils, with lobular hepatocyte necrosis, consistent with a severe drug-induced hepatitis. The lymph nodes sampled showed loss of the normal follicular architecture with expansion of the paracortical areas. Immunoblasts were not increased in number and these features are consistent with phenytoin-induced pseudolymphoma.3 Biopsy of the skin showed a picture of toxic epidermal necrolysis (full thickness epidermal necrosis, separation at the dermo-epidermal junction and a sparse superficial dermal inflammatory infiltrate), also consistent with the phenytoin hypersensitivity syndrome. The sections of lung revealed pulmonary oedema only, with no evidence of infection. The brain displayed classical features of Alzheimer’s disease with amyloid angiopathy, extracellular amyloid plaques, neurofibrillary tangles and Hirano bodies.

Discussion
Phenytoin hypersensitivity syndrome is more common in blacks4 and typically has its onset 3 weeks to 3 months after initiation of therapy. Although variable in presentation, its hallmark clinical features are fever, a rash (erythoderma, progressing via a generalised maculopapular to a pustular rash and finally desquamation), lymphadenopathy and hepatosplenomegaly.5,6 Peripheral leucocytosis and eosinophilia are common. Potentially fatal complications associated with the syndrome include hypersensitivity myocarditis,2 drug-induced hepatitis with or without hepatocyte necrosis (which carries a mortality of up to 50%),2,7 and compromise of renal and pulmonary functions.4

As with any drug hypersensitivity reaction, prompt cessation of phenytoin is the first
essential step in management. A benzodiazepine should be substituted for seizure control, as was done with our patient. However, despite termination of phenytoin, the hypersensitivity reaction is often progressive and it is hence common practice to administer corticosteroids. The role of systemic corticosteroids in phenytoin hypersensitivity has not been studied in a randomised, placebo-controlled trial and its use is based only on anecdotal evidence.

The underlying mechanisms of the phenytoin hypersensitivity syndrome are unknown; immune complex, and delayed-type hypersensitivity allergic reactions, alteration of lymphocyte function and toxic metabolite production have been postulated. Genetic factors are also thought to contribute, since siblings are said to have a one in four risk of showing a similar reaction. Cross-reactivity with other aromatic ring anticonvulsants (eg, phenobarbital and carbamazepine) has been reported in patients with a history of phenytoin hypersensitivity.

We wish to acknowledge the assistance of Dr S Al-Sarraj, Consultant Neuropathologist, Institute of Psychiatry, London.

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Management of pulmonary thrombo-embolism using catheter manipulation: a report of four cases and review of the literature

Peter S C Wong, Shyam P Singh, Robert D S Watson, Gregory Y H Lip

Summary
To date the management of pulmonary thrombo-embolism is still largely limited to anticoagulation. Heparin and oral anticoagulation have been shown to be effective in reducing recurrence and death in venous thrombo-embolism. During the acute stage, systemic thrombolytic therapy has also been advocated for the rapid dissolution of the thrombus in patients with haemodynamic instability. We describe four patients with acute pulmonary thrombo-embolism who were managed with catheter-based thrombus manipulation with intrapulmonary thrombolysis. This management strategy should be considered in patients with pulmonary thrombo-embolism who continue to deteriorate despite conventional management with anticoagulation or systemic thrombolysis.

Keywords: pulmonary thrombo-embolism; catheter manipulation; thrombus; thrombolytic therapy

Pulmonary thrombo-embolism (PE) is a common cause and consequence of hospital admission, with a significant mortality and morbidity. Male sex, old age, serious medical or surgical diseases, immobilisation and trauma are important predisposing factors in patients dying of such disease.1 Untreated PE is associated with a mortality rate of approximately 30%, but can be decreased to 1–10% with the institution of appropriate diagnosis and treatment.2–6

PE usually results from acute obstruction by thrombus, at any location of the pulmonary vascular tree. If severe, it leads to haemodynamic deterioration, systemic acidosis, right heart failure, cardiogenic shock, and ultimately death. To date, the management of PE is still largely limited to anticoagulation. Unfractionated heparin, nicoumalone and warfarin have been shown to be effective in reducing recurrence and death in venous thromboembolism,4,7 and more recently, low-molecular-weight heparins are increasingly used with their relative ease of administration and lack of requirement for anticoagulation monitoring.3 During the acute stage, systemic thrombolytic therapy has been advocated for the rapid dissolution of the thrombus, with some acute reduction of pulmonary artery pressure; nevertheless more evidence is needed for its long-term prognostic benefit, although it may have a role in patients with haemodynamic instability, as such patients have a mortality rate of approximately 20%, despite the use of anticoagulants and other supportive measures.7,8

Besides systemic thrombolytic therapy, there have been isolated reports on the use of pulmonary angiography and catheter-based manipulation with intrapulmonary thrombolysis to achieve clot fragmentation and dissolution (table 1).9–15 We describe four patients who presented to our centre with massive PE who were managed with catheter manipulation and intrapulmonary thrombolysis.

Case reports

Case 1
A 46-year-old Caucasian man was admitted after an assault resulting in a closed fracture of the neck of left humerus and left wrist. He had a medical history of mild asthma and was a non-smoker. Both arm fractures were reduced under general anaesthesia and a piece of autologous bone graft was used to seal the humoral fracture with a metal plate. He was mobilised and well for 3 days postoperatively, when he suddenly experienced acute onset dyspnoea, and became tachycardic (130 beats/min in sinus rhythm) and hypotensive (86/58 mmHg). His white cell count was 21.6 × 10⁹/l and arterial blood gases showed pH 7.34, pCO₂ 3.7 kPa, pO₂ 9.9 kPa, bicarbonate 17.5 mmol/l and base excess of −5.8. Central venous pressure was elevated at +29 cm water. A bedside transthoracic echocardiogram showed a dilated right atrium and ventricle with impaired right ventricular contraction. Left ventricular contraction was normal.

A clinical diagnosis of PE, originating from an axillary vein thrombosis, was made and 95% oxygen was administered through a rebreathing mask with subsequent arterial gases showing pH 7.36, pCO₂ 4.1 kPa, pO₂ 11.2 kPa, bicarbonate 19.7 mmol/l and base excess −5.8. A bolus of 5000 units intravenous (iv) unfractionated heparin was given, followed by an infusion rate of 25 000 units over the next 24 hours. Colloid (haemacel) and dopamine at 2–5 µg/kg/min were given iv to optimise systemic and renal output. However, the condition of the patient continued to deteriorate, with persistent hypotension and increasing hypoxia, and heparin was substituted for a systemic infusion of iv streptokinase at 100 000
He was referred to the on-call cardiologists for pulmonary angiography and clot manipulation. Via a right femoral venous puncture, a 6F Gensini catheter (Cordis) was manipulated into the heart. The mean right atrium (RA) pressure was 18 mmHg, right ventricle (RV) pressure 50/5 mmHg (mean 26) and pulmonary artery (PA) pressure 55/20 mmHg (mean 32). Pulmonary angiography confirmed a large thrombus obstructing the left main pulmonary artery. The Gensini catheter and a J guide wire (0.038”, USCI) were used under direct image intensifier control for attempted clot fragmentation, resulting in the clot successfully being dispersed to a more peripheral location. The catheter was then left in situ, in close proximity to the residual thrombus for further streptokinase infusion. A repeat right heart catheter the following day revealed a mean RA pressure of 15 mmHg and PA pressure of 42/16 mmHg (mean 26) with some persistence of the clot, although smaller in size. A further attempt at clot fragmentation and clot dispersion was made with an 8F pigtail catheter (Cordis) with the clot displaced to an even more peripheral location.

The post-procedure course was complicated by acute renal failure which was treated with haemofiltration for 7 days. The patient recovered fully and was discharged home on Day 30.

Case 2
A 18-year-old woman was transferred to our unit from a local orthopaedic hospital after developing a persistent sinus tachycardia, hypotension and poor arterial oxygen saturation in the immediate postoperative period. She had undergone an endoprosthetic replacement of the left tibia after excision of an osteosarcoma. She had been treated with subcutaneous heparin. On arrival in our hospital, 22 hours after her initial deterioration, her temperature was 37.8°C. She was in sinus tachycardia, with a heart rate of 150 beats per minute. Her blood pressure was 90/60 mmHg and her oxygen saturation was 92% on air. Her oxygen saturation on nasal cannula was 98% at 6 l/min. Her white cell count was 12.5 x 10^9/L with a neutrophil count of 8.5 x 10^9/L. Her creatinine was 0.9 mg/dL. Her haemoglobin was 10.5 g/dL.

Table 1  Right heart catheterisation, pulmonary angiography and catheter manipulation for pulmonary embolus examples of studies in the literature

<table>
<thead>
<tr>
<th>Authors</th>
<th>Fava et al10</th>
<th>Stock et al11</th>
<th>Brady et al12</th>
<th>Essop et al13</th>
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<td>5</td>
<td>3</td>
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<td>49 (20–68)</td>
<td>50 (21–80)</td>
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<td>massive</td>
<td>2 shock</td>
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<td>Onset of symptoms to procedure</td>
<td>&lt;2 lobar arteries</td>
<td>&lt;5 days</td>
<td>≤10 hours</td>
<td>∞ &lt;5 days</td>
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<td>Catheters and guidewire</td>
<td>12 F stainless steel double lumen with suction cup</td>
<td>6 F pigtail; Grollman angioplasty balloon length (12–16 mm)</td>
<td>6–7 F pigtail; 0.035” Terumo; angiofistula balloon length (8–10 mm)</td>
<td>0.035” Goodale-Lubin pigtail; J guide wire</td>
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<td>1 died</td>
<td>all survivors</td>
<td>1 died 2 weeks after discharge</td>
</tr>
<tr>
<td>Complications</td>
<td>pulmonary infarct 11%; myocardial infarct 4%; rupture of pulmonary artery 2%; ventricular perforation 2%</td>
<td>haematoma 19%; haemorrhage 40%; minor bleeding 67%</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

Figure 1 Pulmonary angiograms demonstrating thrombus within the right pulmonary artery (case 2)

Figure 2 Pulmonary angiograms demonstrating thrombus within the left pulmonary artery (case 2)
**Table 2**  Summary of the four case histories

<table>
<thead>
<tr>
<th>Predisposing factors</th>
<th>Case 1</th>
<th>Case 2</th>
<th>Case 3</th>
<th>Case 4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sex</td>
<td>male</td>
<td>male</td>
<td>female</td>
<td>male</td>
</tr>
<tr>
<td>Age</td>
<td>46</td>
<td>18</td>
<td>30</td>
<td>36</td>
</tr>
<tr>
<td>Predisposing factors</td>
<td>fracture; axillary vein thrombosis</td>
<td>osteosarcoma</td>
<td>hospitalisation</td>
<td>pneumonia nphrotic syndrome</td>
</tr>
<tr>
<td>Severity of pulmonary artery obstruction</td>
<td>left main</td>
<td>all major intermediate branches</td>
<td>right upper and middle branches</td>
<td>right main</td>
</tr>
<tr>
<td>Catheters</td>
<td>6 F Gensini</td>
<td>6 F Gensini</td>
<td>8 F pigtail</td>
<td>6 F Gensini</td>
</tr>
<tr>
<td>Catheter manipulation</td>
<td>J guide wire</td>
<td>J guide wire</td>
<td>J guide wire</td>
<td>J guide wire</td>
</tr>
<tr>
<td>Index of improvement</td>
<td>mean PAP reduced from 32 to 26 mmHg</td>
<td>—</td>
<td>mean PAP reduced from 28 to 12 mmHg</td>
<td>—</td>
</tr>
<tr>
<td>Thrombolysis</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
<td>yes</td>
</tr>
<tr>
<td>Outcome</td>
<td>alive</td>
<td>alive</td>
<td>alive</td>
<td>alive</td>
</tr>
</tbody>
</table>

Case 1
A 46-year-old male smoker was admitted with a history of dyspnoea and collapse following a recent in-patient stay for a minor superficial leg injury. She was a smoker and was previously in good health. On examination she had a tachycardia of 138 beats/min, with a blood pressure 85/60 mmHg, and central venous pressure +16 cm water. Mechanical ventilation was in situ and arterial blood gases showed a pH 7.2, pCO₂ 6.8 kPa, pO₂ 20 kPa, bicarbonate 18.4 mmol/l and base excess of −7.6. An electrocardiogram (ECG) confirmed sinus tachycardia, with a frontal axis + 800, right bundle branch block and a S1Q3T3 pattern.

As a result of her critical condition, right heart catheterisation and pulmonary angiography was performed immediately upon arrival (figures 1 and 2). Mean RA pressure was 14 mmHg, RV 50/0 mmHg (mean 25), and PA 42/20 mmHg (mean 32). Bilateral thrombi were visualised to obstruct all intermediate pulmonary branches during pulmonary angiography. An attempt to disperse some of the clots to a peripheral location was made, and the cardiac catheter was left in situ for a continuous streptokinase infusion at 100 000 units/h. The patient was transferred to the Intensive Care Unit for continued ventilation, but remained in cardiogenic shock, with a blood pressure of 40/20 mmHg, despite ionotropic support (iv adrenaline boluses in addition to background dopamine 40 μg/kg/min) and died later that day with no response to therapeutic measures.

Case 2
A 18-year-old female presented with a right-sided pleuritic chest pain, dyspnoea and a productive cough. An initial diagnosis of left lower lobe pneumonia was made and treatment with antibiotics initiated. On day 2, he experienced a sudden onset of increasing dyspnoea. arterial blood gases revealed severe hypoxaemia with pH 7.4, pCO₂ 3.9 kPa, pO₂ 5.3 kPa, bicarbonate 22.6 and a base excess of −1.6. Chest X-ray confirmed pneumonic consolidation in the left lower zone whilst the rest of the lung fields was clear. An ECG showed sinus tachycardia without evidence of myocardial ischaemia.

In view of the mismatch between the degree of hypoxaemia and consolidation, and deteriorating clinical state (increasing dyspnoea, hypertension and hypoxia), a PE was suspected. Pulmonary angiography using a size 6 Gensini catheter (Cordis) revealed a major obstruction of the right main pulmonary artery with thrombus, and clot dispersion was attempted using the catheter and J guide wire (0.038”, USCI). Streptokinase was then infused via the catheter into the pulmonary artery. A repeat pulmonary angiogram 2 days later showed satisfactory perfusion to all areas of the right lung. The patient was subsequently discharged on Day 15.

Case 3
A 30-year-old woman was admitted with a history of dyspnoea and collapse following a recent in-patient stay for a minor superficial leg injury. She was a smoker and was previously in good health. On examination she had a tachycardia of 138 beats/min, with a blood pressure 85/60 mmHg, and central venous pressure +16 cm water. Mechanical ventilation was in situ and arterial blood gases showed a pH 7.2, pCO₂ 6.8 kPa, pO₂ 20 kPa, bicarbonate 18.4 mmol/l and base excess of −7.6. An electrocardiogram (ECG) confirmed sinus tachycardia, with a frontal axis + 800, right bundle branch block and a S1Q3T3 pattern.

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Case 4
A 36-year-old male smoker presented with left-sided pleuritic chest pain, dyspnoea and a productive cough. An initial diagnosis of left lower lobe pneumonia was made and treatment with antibiotics initiated. On day 2, he experienced a sudden onset of increasing dyspnoea. arterial blood gases revealed severe hypoxaemia with pH 7.4, pCO₂ 3.9 kPa, pO₂ 5.3 kPa, bicarbonate 22.6 and a base excess of −1.6. Chest X-ray confirmed pneumonic consolidation in the left lower zone whilst the rest of the lung fields was clear. An ECG showed sinus tachycardia without evidence of myocardial ischaemia.

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Subsequent clinic follow-up investigations revealed the presence of nephrotic syndrome secondary to membranous glomerulonephritis.

**Discussion**

The diagnosis of PE is often made too late, at the patient’s peril. When the thrombotic occlusion causes major haemodynamic compromise, as in our four cases, the diagnosis is seldom in doubt. The diagnosis is, however, less obvious in less severe cases, as symptoms are vague, non-specific or may even be absent initially. Simple investigations such as chest radiography and ECG are often inadequate. Bedside transthoracic echocardiography can provide some information to substantiate the diagnosis, as right ventricular dilatation with hypokinesis, abnormal septal position and paradoxical systolic motion, reduced left ventricular size, pulmonary artery dilatation, tricuspid and pulmonary regurgitation, and a lack of collapse of the inferior vena cava with inspiration are features suggestive of PE on the echocardiogram. The visualisation of
thrombus in the right side of the heart or pulmonary artery is less common, but confirms the diagnosis. More sophisticated investigations such as lung ventilation-perfusion scans, transoesophageal echocardiography and spiral computed tomography can add to the diagnostic accuracy in patients who are clinically stable enough to undergo these tests, which may add to the delay in making the diagnosis in more urgent situations.

Unfractionated heparin is no doubt the initial treatment of choice in the milder form of PE without haemodynamic instability or significant hypoxaemia. However, this form of treatment is associated with some uncertainty about the level of anticoagulation during the initial but crucial phase of PE. Systemic thrombolytic treatment has been advocated in some patients with PE associated with circulatory instability or hypoxaemia although larger studies with more outcome data are still needed. Although previous studies have demonstrated the short-term superiority of thrombolysis in terms of resolution of both radiographic and haemodynamic abnormalities when measurements were made within the first 24 hours, this does not appear to be sustained nor translated into a mortality benefit.

In selected patients where anticoagulation or thrombolytic therapy are contraindicated, or fulminant PE is evident, pulmonary angiography and catheter-based manipulation can be life saving, although more evidence from randomised controlled trials is needed. Pulmonary angiography provides the gold standard for initially visualising the presence of thromboemboli in the pulmonary vasculature. Our usual practice is to perform this under local anaesthetic from the femoral vein, using a Seldinger technique; visualisation is performed in the postero-anterior position with injection of 35 ml of dye (Nipam 340) at 14 ml/s and a pressure of 900 psi via a pigtail or Gensini catheter. The risks of undertaking pulmonary angiography relate to bleeding at the groin (especially with concomitant anticoagulants), possible reactions to the contrast (allergy, renal impairment) and rarely, perforation of the right ventricle.

As illustrated with our cases, intravascular catheters or balloon angioplasty can be used to dislodge or fragment thrombus in a proximal segment of the pulmonary artery, to be dispersed to a more peripheral location. Such manoeuvres can lower the pulmonary vascular resistance, reduce strain on the right side of the heart, and improve perfusion to the affected lung segments. Concomitant use of thrombolytic therapy either via pulmonary artery catheter or peripheral venous catheter can be synergistic with catheter-based manipulation for the treatment of PE, in the absence of contraindications. Our usual practice is to administer streptokinase, initially 250 000 units over 30 minutes, followed by 100 000 units/h for 24–72 hours. The introduction of newer agents such as the glycoprotein IIb/IIIa receptor antagonists may provide alternative antithrombotic strategies in the future, but further trials are needed to confirm the value of these agents.

A large number of different catheters have been used in clinical reports (table 1) since Greenfield originally described the technique of transvenous catheter pulmonary embolotomy. The pigtail catheter is familiar to all cardiologists and because of its round coil design, could be regarded as one of the least trauma-inducing catheter. Moreover, the pigtail coil design can be utilised to a greater advantage by maximising the surface area with the ‘loop upon axial rotation’ ability of the catheter, which facilitates the dislodgement and fragmentation of the thrombus. This approach with the pigtail catheter was used satisfactorily in two of our cases. In addition to the catheter shape, a larger size catheter with a bigger pigtail loop may be easier and more effective to manoeuvre than a smaller catheter of the same design; perhaps more purpose built catheters may be available in the future for this procedure.

At present, there are no data to support the routine use of pulmonary angiography and catheter-based manipulation in all cases of PE. This therapeutic measure should perhaps be reserved for cases whereby there are contraindications to anticoagulation or thrombolysis, or in patients with a rapidly deteriorating clinical course. Surgical embolectomy with or without inferior venae cava filters is another alternative to catheter-based manipulation in severe cases or in the presence of contraindications to thrombolysis, but its place in the management of PE remains controversial. Indeed, the operative mortality of surgical embolectomy in selected patients has been reported to be high, up to 40–70%, and the availability of an emergency cardiothoracic team would limit its widespread application.

In conclusion, we present four cases of acute PE where catheter-based thrombus manipulation with intrapulmonary thrombolysis were used, with a satisfactory outcome in three patients. We do accept that our patients had multiple interventions, including intravenous heparin, and intravenous and intrapulmonary thrombolysis, but we suggest that catheter-based thrombus manipulation with intrapulmonary thrombolysis could be considered in patients with PE who continue to deteriorate despite conventional management with anticoagulation or systemic thrombolysis, especially in centres with access to angiographic facilities.

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19 Oakley CM. There is no place for acute pulmonary embolectomy. Br J Hosp Med 1989;41:469.
Mesenteric infarction due to combined protein C deficiency and prothrombin 20210A defects

C J Mainwaring, M Makris, W E G Thomas, K K Hampton, F E Preston

Summary
The prothrombin gene mutation, 20210A, a guanine to adenine substitution at nucleotide position 20210, has recently been described as an additional risk factor for venous thromboembolic disease. We describe the case of a patient with combined heterozygous prothrombin 20210A mutation and type 1 protein C deficiency who presented with massive mesenteric venous infarction of his small bowel and survived following the use of protein C concentrate and extensive small bowel resection.

Keywords: mesenteric venous infarction; protein C deficiency; prothrombin 20210A

A 49-year-old man presented with diffuse abdominal pain and no bowel action for 3 days. In 1975 surgery for a perforated duodenal ulcer had been complicated by a postoperative right leg deep venous thrombosis (DVT). Since then he had five further spontaneous DVTs related to erratic anticoagulation control. A thrombophelia screen performed in 1996 confirmed heterozygous type 1 protein C deficiency. Reinvestigation following this admission identified an additional thrombophilic defect, the prothrombin 20210 mutation. His baseline protein C antigen and activity levels, when not on warfarin, were reduced at 0.39 and 0.35 IU/ml, respectively (normal range 0.71–1.42). Family studies had shown that both his son and daughter were similarly affected though asymptomatic. Neither his parents nor his siblings had been tested for thrombophilic defects. Because of recurrent thrombotic events he had been maintained on long-term warfarin since 1993, but on the day of admission his anticoagulation was subtherapeutic with an international normalised ratio (INR) of 1.3. Clinical examination demonstrated diffuse abdominal tenderness only. Investigations revealed a raised white blood cell count of $11.8 \times 10^9/l$ with a neutrophilia, normal chest and abdominal radiology, biochemical screens and serum amylase. A gastroscopy demonstrated bile reflux and minimal gastric ulceration without bleeding. He continued to deteriorate despite full supportive care with increasing abdominal distention, absent bowel sounds and guarding. In view of these findings, in the context of a thrombophilic disorder, a small bowel infarction was suspected and a mesenteric angiogram confirmed extensive portal, splenic and proximal superior plus inferior mesenteric venous thromboses. Pre-operatively he was treated with 2850 units (40 units/kg) of intravenous protein C concentrate (Immuno, Vienna) by bolus injection which increased his levels from 0.1 to 0.57 IU/ml.

At operation, extensive venous infarction of most of the small bowel was noted, requiring resection, together with anastomosis of the remaining viable 4.5 cm of jejunum and 7.5 cm of ileum. Postoperatively he was treated with once daily protein C concentrate, by bolus injection aiming for a protein C level of greater than 0.50 IU/ml, from day +1 to + 4 (table) and intravenous heparin, until the activated partial thromboplastin time ratio was stably maintained at 1.5–2.5. There were no further thromboembolic events. The half-life of the infused protein C concentrate was calculated at 16.2 h on day +1 and 13.6 h on day + 4. The longer than anticipated half-life, normally 6 h, probably reflected recovery of endogenous protein C levels whilst not receiving warfarin. Conversion to subcutaneous low molecular weight heparin occurred on day +13 and he was discharged home on day +17. Warfarin was restarted some 6 weeks postoperatively with erratic anticoagulation control initially as a result of impaired absorption. Currently, he remains well on life-long anticoagulation, with no further thrombotic complications, aiming for a target INR of 2.5.

Discussion
Mesenteric venous thrombosis (MVT) causing small bowel infarction is an extremely rare cause of an acute surgical abdomen and often difficult to diagnose. Predisposing factors include acquired conditions such as congestive cardiac failure, atrial fibrillation, myeloproliferative disorders, oral contraceptive use and abdominal infections. Our patient had no
Protein C deficiency and prothrombin 20210A defects

Genetic factors predisposing to venous thrombosis

- antithrombin deficiency
- protein C deficiency
- protein S deficiency
- factor V Leiden mutation
- prothrombin 20210A mutation

Summary points

- inherited thrombophilia has an incidence of at least 5% in the general population
- combined defects are being increasingly recognised and associated with an increased relative risk of venous thromboembolic disease
- thrombophilic screening should be undertaken in patients presenting with mesenteric venous infarction
- protein C concentrate is available and should be considered for congenitally deficient patients presenting with major venous thromboembolic complications

acquired risk factors. MVT has also been described in inherited thrombophilias (box 1), including antithrombin deficiency and the factor V Leiden mutation. The prothrombin gene mutation 20210A is a recently described additional thrombophilic risk factor with an incidence of 2.3% in healthy control subjects and 18% in patients with a personal or family history of DVT. It is now recognised that some thrombophilic patients can be double heterozygotes for two inherited conditions with an increased relative risk for venous thromboembolic disease. Our patient had combined heterozygous type 1 protein C deficiency and the prothrombin 20210A mutation, which probably accounts for the severity of his thromboembolic complications. Additionally, on admission to hospital he had a subtherapeutic INR which would have provided inadequate protection combined with the fact that warfarin further lowers the levels of vitamin-K-dependent protein C, which was intrinsically reduced in this patient. The importance of strict anticoagulation control cannot be overstated and it is possible that this case may have been avoided if his INR had not been allowed to fall to 1.3. The severe protein C deficiency was corrected by the use of intravenous protein C concentrate. Prior to the commercial availability of protein C concentrate, fresh frozen plasma was traditionally used to raise protein C levels in cases of both inherited and acquired deficiencies with variable but generally poor therapeutic results.

Intravenous protein C concentrate has been successfully used in cases of purpura fulminans in congenital homozygous deficient neonates and during pregnancy in a patient with a history of previous venous thrombosis and foetal loss with successful outcomes. Our case is the first to use protein C concentrate successfully in the context of established massive MVT and it is noteworthy that there were no postoperative thromboembolic problems. In the three years since his bowel resection he has had no further thrombotic events whilst on long-term warfarin. In addition to screening patients with MVT for thrombophilia, we feel protein C concentrate should be considered in congenitally deficient patients with major venous thromboembolic complications.