Clinical aspects of the acquired immune deficiency syndrome in the United Kingdom

J N WEBER,* D J CARMICHAEL,† N SAWYER,† A J PINCHING,‡ AND J R W HARRIS*

From the *Praed Street Clinic and the Departments of †Medicine and ‡Immunology, St Mary’s Hospital, London W2

SUMMARY Between August and November 1983, seven new patients with AIDS were seen at this hospital; two with disseminated Kaposi’s sarcoma, and five with opportunist infections, of whom three have died. We present the case histories of the five patients with opportunistic infections, which show the wide clinical range of AIDS and suggest that the pattern of infection in the UK may differ from that reported in the USA.

Introduction

The acquired immune deficiency syndrome (AIDS) is an unprecedented epidemic of immunocompromise, resulting in opportunistic infections, Kaposi’s Sarcoma, and other unusual neoplasms.1-9 It was first documented in mid-1981, and over 3000 cases had been reported by December 1983, including 36 in the United Kingdom.10 11 Over 70% of cases of AIDS have occurred in homosexual men.12 The epidemiological features are compatible with a sexually transmitted, blood borne agent.13 The clinical features suggest a variable latent period, followed by prodromal non-specific symptoms of weight loss, fever, night sweats, and malaise.3 4 The most common opportunistic infection is Pneumocystis carinii pneumonia, which occurs in 50% of all cases of AIDS in the United States of America.12

Case reports

CASE 1

A 22 year old male homosexual prostitute was well until June 1982, when he presented to his general practitioner with fever, night sweats, and weight loss. A chest x ray showed an opacity in the right third interspace, and he was admitted to a local hospital for investigation. On admission, he was noted to have a swinging fever, oral candidiasis, ulcerating perianal herpes simplex, and pancytopenia. The opacity seen on chest x ray proved to be a bifid rib. Blood cultures were repeatedly sterile, and a bone marrow examination showed only normocellular marrow. A therapeutic trial of broad spectrum antibiotics failed to control the fever, and treatment with oxymethalone was started for the pancytopenia. The patient was discharged, and he returned to his home town. The fever, oral candidiasis, and pancytopenia persisted. Between January and June 1983 he was admitted to another hospital on seven occasions for investigation. Oesophageal candidiasis was confirmed by endoscopy, and two episodes of chest infection resolved on treatment with cotrimoxazole without an agent being identified. Over this period, seven grand mal epileptic fits were reported, although there was no history of epilepsy. A computed tomography scan of the head and an electroencephalogram showed no focal abnormality, and examination of the cerebrospinal fluid showed no cells, but a raised protein concentration (1·6 g/1). The patient became increasingly lethargic and apathetic and was transferred to this hospital in August 1983.

On admission, the patient was appreciably wasted (weight 38.5 kg) with oral candidiasis, ulcerating infection with herpes simplex perianally and in the anal canal, and with tender hepatomegaly. He was anaemic, with a blood haemoglobin concentration of 1·03 g/dl and a total lymphocyte count of 0·6 × 10⁹/l. There was anergy to three recall antigens, and an absolute depletion of T helper lymphocyte phenotype (0·23 × 10⁹/l). Gastroduodenoscopy showed candidal oesophagitis, with a 2 cm duodenal ulcer. Despite frequent small haemoptyses, the chest x ray remained clear. Lung function tests, however, showed a pronounced reduction in carbon monoxide...
transfer factor (TICO) (30% of that predicted), with preserved spirometry. A fibreoptic bronchoscopy with lavage and transbronchial biopsy showed no pathogens.

The herpetic ulceration and fever settled on treatment with oral acyclovir 1 g a day, and both returned on withdrawal of this drug, which has since been maintained at 200 mg three times a day. The candidiasis is controlled, but not eradicated, by amphotericin lozenges, nystatin suspension, and oral ketoconazole. Despite symptomatic healing of his duodenal ulcer the wasting has been resistant to treatment; however, there has recently been a 4 kg weight gain. His mental state improved spontaneously, and his fits have been controlled with anticonvulsant treatment.

The patient has never travelled outside the UK and had no known contact with AIDS, although clearly his previous occupation put him at risk of contact with a large number of casual partners. He is currently living independently as an outpatient.

**CASE 2**

A 52 year old male homosexual executive was well until March 1983, when he developed anorexia, severe weight loss (>12 kg), and overwhelming malaise. A pancreatic neoplasm was suspected, but whole body computed tomography and endoscopic retrograde cholangiopancreatography gave negative results in August, although the latter procedure showed pronounced oral and oesophageal candidiasis. In view of the probable diagnosis and severe constitutional symptoms, treatment with prednisolone 40 mg a day was started and the patient went to Malta to convalesce. One week later he developed fever, night sweats, and confusion. On his return to the UK, blood cultures grew *Salmonella typhimurium*, which was also isolated from the stool, and the appropriate antibiotics were started. The septicaemia persisted, and the patient was transferred to this hospital.

On admission he was noted to have florid oral candidiasis and a fluctuating mental state. There was lymphopenia (lymphocyte count 0.8 x 10^9/l), T helpers 0.18 x 10^9/l) with otherwise unremarkable blood tests. Blood culture confirmed infection with *Salmonella typhimurium*, with extensive colonisation of the gut and skin. Computed tomography scanning of the head, electroencephalogram, and examination of cerebrospinal fluid failed to find a specific cause for the altered mental state. A polyomavirus, which was neither JC nor BK, was later isolated from the urine. The relevance of this virus is uncertain at present. The antibiotic regimen was augmented to include all three drugs to which the organism was sensitive (gentamicin, mecillinam, and colistin), but it was never possible to eradicate the resistant salmonella bacteraemia. He died of toxaemia after nine weeks of continuous septicemia.

The patient had visited the USA and Canada on numerous occasions over the preceding five years, with casual sexual contacts. There was no known contact with AIDS. Eighteen months before admission, he had experienced a febrile illness with pronounced lymphocytosis, atypical mononuclear cells, and negative Paul-Bunnell test reactions. This was thought at the time to have been glandular fever not caused by Epstein-Barr virus or cytomegalovirus.

**CASE 3**

A 28 year old male homosexual air steward was well until June 1983 when he developed sudden onset of diarrhea while in Australia. Stool culture grew *Salmonella ohio*; oral candidiasis was also noted. On his return to the UK, his diarrhea persisted, and his general practitioner treated him with ampicillin. A follow up stool examination by his employers showed persistence of the salmonella, but his symptoms settled; he had lost five kg in weight, but this was constant thereafter.

In August, his diarrhea returned, and he was noted to have oral candidiasis; in addition, he was lymphopenic (lymphocyte count 0·8 x 10^9/l). He was referred to the Praed St Clinic, where he was found to have anergy to three recall antigens, and to have T helper depletion (0·23 x 10^9/l); his chest x ray was clear. He was observed as an outpatient until September, when he complained of a sudden onset of dyspnoea over three days. He was admitted immediately, and his chest x ray showed bilateral mid-zone infiltrates, compatible with early atypical pneumonia. He was cyanosed and hypoxic, with an arterial oxygen partial pressure of 7·7 kPa (58 mm Hg). There had been no sputum production. Transtracheal aspiration was performed immediately, but showed no pathogens. Treatment with high dose co-trimoxazole was started (1·92 g 12 hourly) because of the risk of *Pneumocystis carinii* pneumonia. A fibreoptic bronchoscopy was performed 24 hours later with lavage and transbronchial biopsy. *Candida albicans* was seen free and within type II pneumocytes, and a diagnosis of probable candidal pneumonia was made. Treatment with amphotericin and miconazole was started, and erythromycin was added because of the possibility of infection with legionella.

Despite the appropriate antibiotic regimen, the patient's condition worsened clinically, and the hypoxia deepened. Serial chest x rays showed a rapid progression of the pulmonary infiltrates, causing generalised bilateral opacity by the fifth day. On day 6 he was moved to the intensive treatment unit for
ventilation, but had an asystolic cardiac arrest after a
short period of intubation and ventilation; prolonged
resuscitation was unsuccessful.

The patient had visited the USA regularly over the
preceding five years, and had had many casual sexual
contacts in cities throughout the world. There was no
known contact with AIDS.

**CASE 4**

A 34 year old male homosexual air steward was well
until August 1983, when he developed sudden onset
of diarrhoea. He attended his local STD clinic, where
Entamoeba histolytica was isolated from the stool.
He was treated with metronidazole 400 mg twice
daily for one week, and the diarrhoea abated. One
week after the completion of treatment, the
diarrhoea returned. The stool was watery, offensive,
and pale in colour and contained globules of fat. The
diarrhoea was episodic, lasting two or three days,
and the episodes were associated with fever and night
sweats. Weight loss of 5 kg was noticed over the next
month. A routine examination at work showed oral
candidiasis, and a full blood count showed
lymphopenia (lymphocyte count 0·8 × 10^9/l). He
was referred to the STD clinic at this hospital.

On examination he was noted to be slightly wasted,
with mild oral candidiasis, and without lymphadenop-
athy; the abdomen was normal to palpation.
Repeated stool culture showed Entamoeba histo-
ytica (zymodeme 111) and a variety of non-
pathogenic amoebae. Sigmoidoscopy showed normal
mucosa, and a rectal biopsy showed areas of chronic
inflammation, with no evidence of amoebic in-
filtration. Barium enema, isotope liver scan, and abdo-
nominal ultrasound gave negative results. Studies of
cellular immunity showed cutaneous anergy to three
recall antigens and a reduction in T helper pheno-
types (0·22 × 10^9/l). Upper gastrointestinal intuba-
tion and biopsy were performed at the Central
Middlesex Hospital, and oocysts of Cryptosporidium
spp were seen on the jejunal mucosa and were later
isolated from the stool by a concentration technique.

The patient is currently being evaluated for treat-
ment of cryptosporidium, which can also cause
symptomatic diarrhoea in immunocompetent
homosexual patients.\textsuperscript{14,15} He has had numerous
visits to the USA and elsewhere, and has had many sexual
partners in the USA.

**CASE 5**

A 47 year old homosexual man was well until
October 1983, when he noted severe malaise and a
persistent sore throat. He presented to the Praed St
Clinic, where a routine examination showed slight
lymphopenia (lymphocyte count 1·3 × 10^9/l). He
declined to return for follow up, and one month later
began to develop a dry cough, night sweats, and
shortness of breath on exertion. His general practi-
tioner referred him to the chest clinic at this hospital,
where a chest x ray showed bilateral mid-zone
pulmonary infiltrates, consistent with an atypical
pneumonia.

On admission in November, cyanosis, florid oral
candidiasis, and tachypnoea were noted. The spleen
was palpable 3 cm below the costal margin. Blood
gases on air showed marked hypoxia. The
lymphocyte count was 0·8 × 10^9/l (T helpers 0·14 ×
10^9/l). A fiberoptic bronchoscopy with lavage and
transbronchial biopsy was performed immediately,
but no organisms were positively identified.
Treatment was started with co-trimoxazole 1·92 g 12
hourly for presumptive Pneumocystis carinii
pneumonia, and parenteral antifungal treatment with
amphotericin was started 48 hours later. The chest x
ray appearances continued to worsen, and the blood
gases deteriorated over the next four days; he was
transferred to the intensive treatment unit. A diffuse
maculopapular skin rash developed eight days after
starting co-trimoxazole. This drug reaction is
commonly seen in the context of AIDS.\textsuperscript{16} The
antibiotic regimen was augmented to cover legionella
with erythromycin, and the patient was ventilated
when unable to maintain his arterial partial oxygen
pressure. He died of pneumonia of unknown cause
on the 14th day.

The patient had visited Canada in 1978, but had
not been to the USA or to Haiti in the past five years.
He had had large numbers of casual sexual partners
in the UK, but had no known contacts with AIDS.

**Discussion**

Owing to the different environment, the pattern of
infectious diseases in the USA differs from that in the
UK. For this reason, the pattern of opportunistic
infections in AIDS in this country cannot be assumed
to be identical to that previously reported from the
USA. Analysis of the small number of cases at this
hospital indicates some notable differences from the
general American pattern.

Pneumocystis carinii was not isolated in any of
these cases, despite adequate transbronchial biopsy
and bronchial lavage with appropriate staining.
Open lung biopsy was not possible in the two fatal cases
of pneumonia, owing to the rapid development of
severe hypoxia; however, neither of these patients
responded to empirical treatment with high dose
cotrimoxazole. The cause of the fulminant pneumonia
was not identified in one case (case 5) despite
postmortem lung biopsy. In case 3, a probable
diagnosis of Candida albicans pneumonia was made
on cytological examination only. It would be
expected that Pneumocystis spp would be identified by fibroptic bronchoscopy in 85% of cases.22

The incidence of infection with Salmonella spp (cases 2 and 3) shows a noticeable variance from that in the USA, although it is similar to European reports of cases from Africa.17 Salmonella species are not currently listed as being associated with AIDS in the epidemiological protocol of the Centers for Disease Control.18 In case 2 Salmonella typhimurium was the principal pathogen, and AIDS was confirmed by the additional oesophageal candidiasis. In case 3 the salmonella was restricted to the gut, and blood cultures gave consistently negative results. It is reasonable to surmise that Salmonella spp may be of greater clinical importance in AIDS in the UK. As a facultative intracellular pathogen it is an appropriate secondary infection to this type of immune deficit.

Cytomegalovirus (CMV) was not excreted in the blood, stool, or urine in any of these patients, in contrast to the common finding of CMV viruria in the USA.19 In a study of asymptomatic homosexual men in San Francisco, CMV viruria was found in 32% of seropositive men, whereas a comparative study of 100 asymptomatic homosexual men at this hospital showed that only 1% of seropositive men excreted CMV in the urine.20 CMV was not cultured from any site in all the above cases.

Candida albicans was found in all the patients with AIDS at this hospital, both as oral candidiasis with large easily visible plaques on the buccal mucosa, and as oesophageal candidiasis at endoscopy or by radiology. The oesophageal candida was not associated with overt ulceration in these cases. The aetiology of the duodenal ulcer in case 1 was not established, but no opportunist agent was found in the ulcer crater. The presence of oral candidiasis was an early sign in all these cases, and was found before the isolation of other opportunistic infections.

In cases 3 and 5, a mild prodromal illness of less than eight weeks' duration without any significant weight loss was reported. This preceded fulminant pneumonia, which developed rapidly. There was a history of only three days of dyspnoea before admission to hospital in both these cases. This rapid development has been rare amongst the reported cases in the USA, and precluded open lung biopsy.3,5 Lymphopenia, with a total lymphocyte count in peripheral blood of less than 1·5 x 1091, preceded overt opportunistic infection in every case. In case 5 lymphopenia (lymphocyte count 1·3 x 1091) was the only laboratory abnormality six weeks before the onset of fatal pneumonia. By comparison, in case 1 lymphopenia has been present continuously for 18 months with oesophageal candidiasis and perianal herpes simplex but without life threatening infection.

In conclusion, we recommend that asymptomatic homosexual men should have a complete physical examination, with special attention to the buccal mucosa for oral candidiasis and lesions of Kaposi’s sarcoma; the entire skin area should be examined. A full blood count with differential and platelet count should be taken at every visit, and a serum sample stored at -80°C for future reference. A throat swab for candida should be taken. If facilities exist for the investigation of high risk patients, estimation of total T helper numbers and in vivo testing of cellular immune response with intradermal recall antigens (purified protein derivative, candida, and streptokinase-streptodornase) may be of value. Immunological abnormalities may occur in healthy, asymptomatic homosexual men.21 The finding of lymphopenia, oral candidiasis (in the absence of recent antibiotic treatment), or both, is important and warrants further investigation for occult opportunistic infection.

The investigation of homosexual men with symptoms of severe malaise, loss of weight, night sweats, fevers, or prolonged diarrhoea should be undertaken urgently. In addition to the tests above, strenuous efforts should be made to investigate the gastrointestional tract for pathogens, the chest should be x rayed, and lung function tests should be performed. The combination of symptoms with lymphopenia, oral candidiasis, or both is an indication for urgent admission to hospital and extensive investigation.

We thank the referring physicians, Drs Dawson, Green, and Chanarin; the physicians and staff of the respiratory laboratory; Dr I Trotman of the department of gastroenterology at Central Middlesex Hospital; the staff of the department of microbiology, particularly Drs D Jeffries and A Maddocks; Dr D Coleman and the department of cytology; and most importantly Sister M Anthony and the nursing staff of Almroth Wright Ward, St Mary's Hospital. The management of patients with AIDS in hospital requires a multidisciplinary approach, and it is not possible to thank by name all the physicians and technicians who have been concerned with the diagnosis and treatment of these patients.

References

Clinical aspects of the acquired immune deficiency syndrome in the United Kingdom


Clinical aspects of the acquired immune deficiency syndrome in the United Kingdom.
J N Weber, D J Carmichael, N Sawyer, A J Pinching and J R Harris

Br J Vener Dis 1984 60: 253-257
doi: 10.1136/sti.60.4.253