Avascular necrosis of the head of the femur in a case of retroperitoneal fibrosis

B. A. KAMDAR
M.B., B.S., F.R.C.S.

Orthopaedic Unit, Department of Surgery,
Hammersmith Hospital and Royal Postgraduate Medical School, London W.12

Summary
A case of avascular necrosis of the head of the femur is reported. The patient is known to suffer from retroperitoneal fibrosis. The possibility of an association between these two conditions is discussed.

In a review of the medical literature since 1948 when retroperitoneal fibrosis was first described by Ormond, no case of avascular necrosis of the head of femur has been reported in association with it. A patient is reported here with retroperitoneal fibrosis who 10 years later presented with avascular necrosis of the head of the femur.

Case report
The patient is a woman aged 67 years, known to suffer from retroperitoneal fibrosis, first diagnosed in 1960. She had had a left nephrectomy for hydronephrosis and pyelonephritis of the left kidney in September 1959. She was well until December 1959. She then developed low backache and on 18 December 1959 developed oliguria. She was treated by emergency measures and a spontaneous diuresis followed; the serum electrolytes and urea were normal by 21 December. An excretion urogram on 1 January 1960 showed a right hydronephrosis; the ureter was narrowed at the level of L5 vertebra and a similar appearance had been noted on the left side in September 1959. The ESR was constantly raised. A provisional diagnosis of retroperitoneal fibrosis was made and a right nephropexy, ureterolysis and a retroperitoneal biopsy were performed on 2 January 1960. She made satisfactory progress till November 1960 when oliguria developed. A right ureterostomy was performed as an emergency procedure which was later converted into an uretero-ileocystoplasty, at which time recurrence of retroperitoneal fibrosis was noted. Subsequently she developed oedema in both lower limbs which was controlled by regular use of diuretics.

Apart from occasional urinary tract infection the patient remained well until the end of July 1970, when she developed pain in her right groin and iliac fossa. She was admitted to the renal unit for assessment on 6 August 1970. The pain in her right groin settled with rest and as no urological cause for pain was found she was discharged home. However, the pain returned with increasing severity in October 1970 and the patient was referred to the orthopaedic clinic. At that time she could not bear any weight on her right hip; the movements of the right hip joint were extremely painful and grossly restricted. There was no history of previous trauma and no history of steroid therapy or the use of the methysergide group of drugs.

Radiographs of the right hip joint showed marked irregularity and depression of the supero-lateral part of the articular surface of the femoral head (Fig. 1). There was diminution of the joint space. Routine haematological investigations were normal. The ESR was 85 mm/hr. Though the appearances were consistent with avascular necrosis of the head of femur, infection could not be excluded. Aspiration of the hip was negative.

At operation the right femoral head and neck were excised and replaced by an Austin Moore's pros-

![Fig. 1. Radiograph of the pelvis showing the avascular necrosis of the head of right femur](http://pmj.bmj.com/)
Sections of the synovium and of the ligamentum teres (Fig. 2) showed a mild degree of chronic inflammatory cell infiltration and prominent cuffing by lymphocytes and plasma cells of many of the capillaries, venules and a few arterioles. There was endothelial swelling and infiltration by mononuclear cells and occasional polymorphonuclear neutrophils. There was no evidence of fibrinoid necrosis. The capsule showed no special features.

Histological examination of the retroperitoneal adipose and fibrous tissue (Fig. 3) showed a diffuse infiltration by lymphocytes, plasma cells, eosinophils and polymorphs. There was perivascular cuffing in some areas. In addition some new fibrous tissue had formed. The appearances were consistent with a diagnosis of retroperitoneal fibrosis.

Discussion
It cannot be concluded from one case that there is definite evidence of some association between retroperitoneal fibrosis and avascular necrosis of the head of femur, but the occurrence of the latter in a patient who has had retroperitoneal fibrosis for the past 11 years, and the presence of vasculitis in the tissues taken for biopsy from the retroperitoneum in 1960 and from the hip in 1970, does suggest a possible relationship between the two conditions. That the retroperitoneal disease in this case has been constantly active for the past 11 years is shown by the following facts: (1) continued involvement of the right ureter by the progressive retroperitoneal fibrosis, leading to several operations between December 1959 and December 1960; (2) continually raised erythrocyte sedimentation rate for the past eleven years; (3) the development of oedema in both lower limbs, more pronounced on the left leg than the right leg. Venography demonstrated narrowing of the inferior vena cava at the level of the L4 vertebra.

Acknowledgment
I would like to thank Mr W. H. Stephenson for his encouragement in the preparation of this paper and for allowing me to publish the case which is under his care.

Bibliography
An unexpected abdominal mass

MEIRION THOMAS*
M.B., B.S.

Masaka Hospital, Uganda

Summary

An unusual complication of ectopic pregnancy, presenting during post-menopausal life, is reported. As hospital services become available to previously medically isolated communities, early diagnosis and treatment will make such cases as this rare.

Case report

A Baganda woman was admitted in February, 1971, complaining of a painless abdominal mass present for over 20 years. She had not menstruated for over 2 years but, like most women in rural Africa, was uncertain of her age. She was probably 50 years old. Of her three children, two had died in the first week of life, and only the third, a 30-year-old male, was alive and well. One year after the birth of her third child, she had had amenorrhoea for 6 months and she thought she was pregnant again. Uneventfully normal menstruation was resumed and she was subsequently sterile.

On abdominal examination, a very large mass, the size of a 20 week pregnancy, was palpated, arising centrally from the pelvis. On vaginal examination, the site of the mass was confirmed and was thought to be in continuity with the uterus. A pre-operative diagnosis of calcified fibroids was made.

At laparotomy, the uterus was only slightly enlarged and studded with very small calcified fibroids. The right ovary was atrophic and there was a small right hydrosalpinx. Replacing the left ovary there was an oval, calcified mass, 20 x 10 cm, attached to the uterus by the left fallopian tube and left utero-ovarian ligament. The mass was densely adherent to small bowel and omentum. Adhesions were divided and a total hysterectomy and bilateral salpingo-oophorectomy was performed.

After operation, the mass was opened using a variety of orthopaedic instruments. The calcified capsule was 1–2 cm thick (Fig. 1). Some foetal parts were identified and removed (Fig. 2), but most of the skeleton had been incorporated into the capsule.

Comment

It was surprising to us that the patient denied any symptoms referable to a ruptured ectopic pregnancy.

The bone development suggested that the gestational age of the foetus was 5–6 months. Remarkably, this 'missed' ectopic had organized, and over a

* Present address: Department of Surgery, Royal Postgraduate Medical School and Hammersmith Hospital, Ducane Road, London, W12 OHS.

FIG. 1. Operative specimen described in the text.
Avascular necrosis of the head
of the femur in a case of
retroperitoneal fibrosis

B. A. Kamdar

doi: 10.1136/pgmj.49.567.59

Updated information and services can be found at:
http://pmj.bmj.com/content/49/567/59

These include:

Email alerting service
Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Notes

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/