Olecranon bursitis is a relatively common problem presenting to accident and emergency (A&E) departments. One third of cases are likely to be septic. The remainder are referred to in this paper as non-septic olecranon bursitis, which includes the cases sometimes termed traumatic or idiopathic olecranon bursitis. Occasional cases are related to rheumatoid arthritis or gout. As the clinical diagnosis can be difficult, septic cases are often not diagnosed; treatment with anti-inflammatory agents alone then leads to a delay in diagnosis and worsening suppuration. Incision and drainage can often be avoided by initial aspiration and antibiotics.

In this paper I discuss the aetiology of these conditions, the various tests which can identify the septic cases, the treatment options, and finally suggest an approach to management in A&E.

Epidemiology
The overall incidence of olecranon bursitis is not known. The vast majority of cases are seen in males, typically aged 30-60 years.

Aetiology
In non-septic olecranon bursitis, inflammation arises either because of bleeding into the bursa, or through the release of inflammatory mediators following trauma. This inflammation can lead to permanent damage to the epithelial lining of the bursa, predisposing to repeated attacks in the future.

In septic olecranon bursitis, trauma, breaks in the skin, or foci of infection provide a portal of entry from which bacteria appear to migrate across soft tissues to enter the bursa. However, there is not always a good history of trauma, so penetration through minute breaks in the skin must occasionally happen. Haematogenous spread does not appear to be the usual route of infection. Ninety per cent of the remainder are infected with β haemolytic streptococci. A range of more unusual organisms—Gram positive and Gram negative bacteria, mycobacteria, and fungi—have been identified in infected olecranon bursae, frequently in the context of systemic diseases. There is evidence that many of these are spread by the haematogenous route.

About one third of septic olecranon bursitis cases have a history of a previous episode of olecranon bursitis. It may be that in these cases the fibrosis and distortion of vascular architecture from previous inflammation prevents the bursa from healing efficiently with foreign material, such as bacteria. The olecranon bursa has a relatively poor blood supply, which is in contrast to the synovial membranes of joints. This may, in part, explain why septic olecranon bursitis is much commoner than septic arthritis. The bursal membrane is biologically different from that of joints. Impaired immunity is an important aetiological factor in up to half of all septic cases. The most common reason is alcohol abuse, but steroids, diabetes, renal impairment, and malignancy are also responsible.

Presenting features
The onset of symptoms can be over several hours or several days. The degree of inflamma-
tion can range from a painless inconvenient lump to a painful hot swelling with extensive cellulitis, accompanied by a fever.

**Diagnosis**

The clinical features may suggest gout or rheumatoid arthritis as the cause, and may help to differentiate septic from non-septic olecranon bursitis. Patients with septic olecranon bursitis usually seek help earlier, and about half are febrile. Bursae which are simply swollen with no pain are almost always non-septic. Pain, warmth, and tenderness, although worse in septic cases, are poor discriminators. Erythema, although seen in 63–100% of septic cases, is also seen in 25% of non-septic cases. On aspiration, purulent fluid indicates septic olecranon bursitis, but a serosanguinous or hazy appearance is often seen in both conditions.

Laboratory tests are essential to differentiate septic from non-septic bursitis when doubt remains after clinical examination, so the aspirate should be sent for microscopy and culture. Microscopy may identify urate crystals, as rarely the first presentation of gout can be with olecranon bursitis. Bacterial culture identifies a causative organism in practically all cases of septic bursitis that have not already received antibiotics.

Total and differential white cell counts in the bursal fluid are rapid, sensitive, and specific tests for identifying septic cases. Although these tests are not offered routinely, many laboratories are able to perform them. The total white cell count is usually greater than 100 × 10⁹/litre (range 1–300) in septic cases, and less than 5 × 10⁹/litre (range 0.05–11) in non-septic cases. Neutrophils average 85% (range 52–98%) in septic cases and 29% (range 0–90%) in non-septic cases. These tests can be performed with anticoagulated samples and the automated cell counters used in haematology. The commonly performed semiquantitative methods of reporting cells in wet preparations will usually report large numbers of cells in both septic and non-septic olecranon bursitis, and will not be able to differentiate between these two conditions.

Serum/bursal fluid glucose concentration differences are more marked in septic cases, but this has poor sensitivity as a diagnostic test.

If there is any doubt about the diagnosis then treatment should be given as for a septic case until culture results are available.

**Treatment**

Antibiotic treatment alone will not prevent progression of many cases of septic olecranon bursitis, so initial aspiration to dryness is recommended. This should be repeated as fluid reaccumulates. If the fluid is viscous or particulate, then lavage with sterile normal saline through a large needle will improve drainage. Incision and drainage may be necessary if there is a pointing abscess, or aspiration and antibiotics fail to control sepsis. In some severe cases, placement of an intrabursal catheter followed by continuous irrigation with an antibiotic solution has been advocated as an alternative. However patients treated in this way have required lengthy admissions. In refractory cases, particularly those with a chronic discharging sinus, or in recurrent cases, bursectomy may be necessary.

Penicillins and erythromycin appear to penetrate the olecranon bursa well, reaching concentrations comparable to those in serum. Concentrations of oxacillin (related to flucloxacillin) greater than 10 times the minimum inhibitory concentration for Staphylococcus aureus can be achieved with standard oral doses. Much higher levels can be achieved with parenteral treatment. Standard doses are effective in sterilising the bursa in a mean of five days in immunocompetent patients, but tend to take much longer in those cases which begin treatment late. Similarly the time to sterilisation is longer, with an average of about 11 days in those with impaired immunity. Patients with systemic sepsis should have parenteral treatment. Olecranon bursitis caused by any of the more unusual organisms will have to be managed along with any underlying disease, and antibiotic treatment should be guided by bacterial sensitivities.

If non-septic olecranon bursitis is treated by aspiration alone, then half the cases will settle in two weeks, but a quarter will still have a bursal effusion at eight weeks, and 10% at six months. It is likely that the addition of non-steroidal anti-inflammatory drugs (NSAID) should lead to a more rapid resolution of symptoms, but the only controlled trial to compare these two options failed to prove this, perhaps because of the small size (n = 42) of the study, with patients randomised to one of four treatment options. This study did, however, show that intrabursal methylprednisolone led to a markedly more rapid reduction in bursal swelling than either aspiration alone or aspiration and NSAID, and there were also less recurrent cases over six months of follow-up. Concern has been raised that intrabursal steroids may cause long term problems, including skin atrophy, chronic local pain, and local infection. This trial did not find evidence of long term problems, but because of its small size and short period of follow up (maximum six months), this concern still remains.

**Prognosis**

With aspiration and adequate antibiotic treatment from early in the natural course, septic olecranon bursitis will usually resolve completely without the need for surgical drainage. However, symptoms may still be prolonged, with a mean of five weeks to complete resolution, and occasional cases may take five or more months. Persistent sinuses may follow incision or spontaneous rupture, and may necessitate bursectomy.

In non-septic olecranon bursitis effusion may persist for weeks, but pain usually settles more rapidly.

Recurrent episodes of non-septic bursitis may occur after either septic or non-septic bur-
Olecranon bursitis

Suggested approach to management in A&E

(1) Rheumatoid arthritis and gout need to be considered as causes.

(2) All cases of olecranon bursitis should be aspirated, whether inflamed or not.

(3) Microscopy for cells and crystals, Gram stain, and culture should be requested. In cases where the bursa is clinically inflamed antibiotics should be given until the culture results are available, when a firm diagnosis of septic or non-septic olecranon bursitis can be made. Cases which are clearly non-septic can be treated with NSAID and a course of antibiotics. Some cases will need admission, and a few will need surgical treatment. Non-septic olecranon bursitis can be managed with aspiration alone. Non-steroidal anti-inflammatory drugs probably hasten symptomatic improvement. Intrabursal corticosteroids produce a rapid resolution but concern remains over their long term local effects. Recovery from septic olecranon bursitis can take months.

I am grateful for the advice of Dr I O’Sullivan and Mr D P Watson, of the Accident and Emergency Department, and Prof R Grahame of the Rheumatology Dept, Goy’s Hospital.

Septic and non-septic olecranon bursitis in the accident and emergency department--an approach to management.

I M Stell

doi: 10.1136/emj.13.5.351

Updated information and services can be found at:
http://emj.bmj.com/content/13/5/351

*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/