Pyoderma gangrenosum is a rare but serious ulcerating skin disease, the treatment of which is mostly empirical. Pyoderma can present to a variety of health professionals and several variants exist that may not be recognised immediately. This can delay the diagnosis and have serious clinical consequences. The mainstay of treatment is long term immunosuppression, often with high doses of corticosteroids or low doses of ciclosporin. Recently, good outcomes have been reported for treatments based on anti-tumour necrosis factor and infliximab proved effective in a randomised controlled trial. This article reviews the presentation of pyoderma gangrenosum and the therapeutic options available.

Methods

We used the keyword “pyoderma gangrenosum” to search Medline. We also searched the Cochrane database but found no Cochrane review on this disease.

How can pyoderma be recognised?

Several variants exist, but the most common one is classic pyoderma gangrenosum. This presents as a deep ulcer with a well defined border, which is usually violet or blue. The ulcer edge is often undermined (worn and damaged) and the surrounding skin is erythematous and indurated (fig 1). The ulcer often starts as a small papule or collection of papules, which break down to form small ulcers with a “cat’s paw” appearance. These coalesce and the central area then undergoes necrosis to form a single ulcer.

Classic pyoderma gangrenosum can occur on any skin surface, but is most commonly seen on the legs. Patients are often systemically unwell with symptoms such as fever, malaise, arthralgia, and myalgia. Lesions are usually painful and the pain can be severe. When the lesions heal the scars are often cribriform. Early diagnosis and prompt treatment reduce the risk of scars, and disfigurement may occur if the diagnosis is missed. Pathergy occurs in 25-50% of cases—lesions develop at the site of minor trauma, so surgery or debridement are contraindicated.

Peristomal pyoderma gangrenosum

Peristomal pyoderma, which occurs close to abdominal stomas, comprises about 15% of all cases of pyoderma. Most of these patients have inflammatory bowel disease, but peristomal pyoderma can occur in patients who have had an ileostomy or colostomy for malignancy or diverticular disease. A large questionnaire based study found a 0.6% incidence of peristomal pyoderma among patients with abdominal stomas. The ulcers in these patients have a similar morphology to classic pyoderma gangrenosum, but bridges of normal epithelium may traverse the ulcer base (fig 2). The lesions are painful and often interfere with the stoma bag adhering to the abdominal wall, which can cause the contents of the bag to irritate the skin more than usual.
Pustular pyoderma gangrenosum
Pustular pyoderma is a rare superficial variant of the disease. Pyoderma often begins as a pustule or group of pustules that later coalesce and ulcerate. This process stops at the pustular stage in pustular pyoderma, and the patient has a painful pustular lesion that may persist for months (fig 3). Pustular pyoderma seems to be confined to patients with inflammatory bowel disease and tends to occur on the trunk and extensor surfaces of the limbs.8–10

Bullous pyoderma gangrenosum
Bullous pyoderma is a superficial variant that affects the upper limbs and face more than the lower limbs. It is associated mostly with haematological conditions. This form of the disease presents as concentric bullous areas that spread rapidly in a concentric pattern. They may break down to form more superficial ulcers than those seen in classic pyoderma, although they still have the blue undermined edge. Prognosis is often poor because of the underlying haematological malignancy.11

Vegetative pyoderma gangrenosum
Vegetative pyoderma is a superficial form of disease that seems to be less aggressive than other varieties (fig 4). It usually occurs as a single lesion in patients who are otherwise well and may respond to local treatment more readily than other forms of the disease.9–10

What is the histopathology of pyoderma?
The histopathology of pyoderma gangrenosum depends on the timing and site of the biopsy.11 Biopsies taken early in the disease and from the advancing, erythematous border tend to show an infiltrate of chronic inflammatory cells confined to the dermis (fig 5). They often have features suggestive of vasculitis at the edge of the ulcer, with a perivascular lymphocytic infiltrate and fibrinoid necrosis of the dermal vessel wall. Occasionally, extravasation of red blood cells and areas of thrombosis are also seen. Ulceration of the epidermis tends to be secondary to the dermal inflammation. Biopsies taken later in the course of ulceration usually show a polymorphonuclear cell infiltrate with features of ulceration, infarction, and abscess formation.11

Who gets pyoderma gangrenosum?
About half of the cases are associated with underlying systemic conditions, such as inflammatory bowel disease, arthritis, and haematological malignancies. About 30% of cases occur in patients with inflammatory bowel disease. About 2% of patients with inflammatory bowel disease will develop pyoderma.12 Occasionally, the skin condition presents before the bowel disease. Pyoderma gangrenosum was thought to be associated only with ulcerative colitis, but both Crohn's disease and ulcerative colitis have a similar incidence.13 Pyoderma gangrenosum is not related to the activity of the inflammatory bowel disease, and it often occurs in patients whose bowel disease is in clinical remission.

About 25% of patients have arthritis, most often seropositive rheumatoid arthritis, although the disease can occur in patients with seronegative arthritis or spondyloarthropathy. As with inflammatory bowel disease, the activity of the arthritis is not related to pyoderma.14 Haematological malignancies are the next most common disorders associated with pyoderma, and these tend to be myeloid rather than lymphoid in origin. Leukaemia is the most frequently reported...
malignancy, usually acute myeloid leukaemia and most commonly the myelocytic or monomyelocytic type.11

What is the differential diagnosis?
The diagnosis of pyoderma gangrenosum is based mainly on clinical findings because biopsies show no specific diagnostic features. In many cases, however, a biopsy can help exclude other conditions such as malignancy, infections, or cutaneous vasculitis. Swabs should be taken from the ulcer, as pyoderma is treated differently from infection. The differential diagnosis of pyoderma gangrenosum is wide (box). Special mention must be made of Sweet’s syndrome, which is characterised by sudden onset of fever and an erythematous, papular eruption. Patients have leucocytosis and skin biopsy shows a dense neutrophilic infiltrate. Sweet’s syndrome and pyoderma can coexist in the same patient as they are both neutrophilic dermatoses.12

How is it treated?
No single, specific treatment exists and few controlled trials of treatment have been done.16 Most clinicians use a stepwise approach and both topical and systemic treatments. Immunosuppression is the mainstay of treatment, and the most commonly used drugs are corticosteroids and ciclosporin. Several other immunosuppressive agents have been used with varied results, but treatment is largely empirical and the choice of treatment often depends on local experience.13 16

**Topical treatments**
Highly potent topical corticosteroids (occasionally underneath occlusive dressings) may be sufficient to induce remission.2 Triamcinolone 40 mg/ml may be injected into the ulcer edge, either alone or as an adjunct to systemic treatment.17 Recently, topical tacrolimus has been shown to be effective in patients with peristomal disease. This is now available as a 0.1% and 0.03% ointment.5

**Corticosteroids**
Most patients need systemic treatment to induce remission and doctors often start patients on oral corticosteroids at an early stage. Prednisolone is the drug of choice and is usually started at high doses (60-120 mg) (level B evidence).18 Patients exposed to these doses for a long time are at risk of steroid related side effects and may benefit from the addition of a bone protecting agent. Minocycline 100 mg twice daily may be of some benefit, usually as an adjunct to oral steroids (level C evidence).18 Rapid improvement has been reported in patients with severe disease given intravenous methylprednisolone as pulse therapy of 1 g daily for three to five days (level B evidence), and several series and reviews support this treatment.19

**Ciclosporin**
Other immunosuppressive agents may be used—firstly, to reduce the dependence on corticosteroids and, secondly, because pyoderma is often resistant to treatment. When corticosteroids fail, the most widely used alternative is ciclosporin. Several case reports and small case series have demonstrated a good clinical response to low dose ciclosporin (level B evidence). Most patients show clinical improvement within three weeks with a dose of 3-5 mg/kg/day. Ciclosporin has several serious side effects, including nephrotoxicity, hypertension, and increased risk of cancer. Such side effects are usually more severe in patients with renal impairment, which may make ciclosporin a less suitable option.13 16

**Ongoing research**
When is the optimal time to introduce treatments based on anti-tumour necrosis factor α? If remission is achieved with infliximab, should patients be placed on maintenance infusions? If so, for how long? How does infliximab compare with conventional treatments, such as corticosteroids or ciclosporin?

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**Differential diagnosis of pyoderma gangrenosum**

**Infections**
- Bacterial
- Mycobacterial
- Fungal
- Viral
- Parasitic

**Malignancy**
- Squamous cell carcinoma
- Cutaneous lymphoma

**Vascular ulceration**
- Venous or arterial disease
- Antiphospholipid syndrome

**Systemic conditions**
- Systemic lupus erythematosus
- Rheumatoid arthritis
- Behcet’s disease
- Wegner’s granulomatisis
- Sweet’s syndrome
Clinical review

GP tips

Pyoderma gangrenosum can occur on any skin surface and should be considered in any ulcer or surgical wound that does not heal. Pyoderma usually develops rapidly and can progress from a pimple to a crater in 24–48 hours. Pyoderma is usually painful and patients may have systemic features such as fever. If pyoderma is suspected, the patient should be referred urgently for a specialist opinion.

Other immunosuppressants

Azathioprine, used alone or combined with corticosteroids, has had variable results. Tacrolimus has also been used systemically, and two centres where patients received 0.1–0.3 mg/kg have reported sustained responses.

Anti-tumour necrosis factor α agents

Pyoderma has been reported to respond to infliximab, a monoclonal antibody against tumour necrosis factor-α. More recently, pyoderma gangrenosum was reported to resolve after treatment with etanercept, a recombinant protein that neutralises the soluble factor.

What is our approach to treatment?

Morbidity and potential for rapid progression are high in most patients, so once suspected we advise an urgent referral to the dermatology department. We recommend oral corticosteroids (with or without minocycline) as first line treatment. If patients do not respond promptly, we then use infliximab as this has fewer recognised side effects than ciclosporin and has been used widely in inflammatory bowel disease and rheumatoid arthritis. We recommend an induction dose regimen of 5 mg/kg at weeks 0, 2, and 6, followed by further treatments as necessary, depending on response. Another immunosuppressant, such as azathioprine, should be given at the same time as infliximab. This approach is not strictly evidence based but is now accepted practice in other inflammatory conditions, such as Crohn’s disease.

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A patient’s story

I first came across pyoderma about a month after having an ileostomy in December 2001. It started out as a small but sore spot rather like an insect bite, just under the stoma. Over the next 24–48 hours the spot took on a life of its own, growing and erupting into an ulcer with blue and red edges. I occasionally needed morphine for the pain. I was admitted to hospital where I was introduced to a doctor conducting a trial of infliximab, a new treatment for pyoderma. I decided to take up the offer of a place on the trial—I felt I had nothing to lose. The drug was given by infusion and the pyoderma was brought under control within 48 hours. Four years later I am on a maintenance regimen of infusions every eight weeks. I contracted pyoderma on my leg about two years ago, which flared up while I was not being treated with infliximab (during a six month break in treatment). Since being on the maintenance dose I have had no problems.

1 Harris AJ, Regan P, Burge S. Early diagnosis of pyoderma gangrenosum is important to prevent disfigurement. BMJ 1998;316:325.