Ulcerative conditions

Mouth ulcers are common and are usually due to trauma such as from ill fitting dentures, fractured teeth, or fillings. However, patients with an ulcer of over three weeks' duration should be referred for biopsy or other investigations to exclude malignancy (see previous article) or other serious conditions such as chronic infections.

Ulcers related to trauma usually resolve in about a week after removal of the cause and use of benzylamine hydrochloride 0.15% mouthwash or spray (Difflam) to provide symptomatic relief and chlorhexidine 0.2% aqueous mouthwash to maintain good oral hygiene.

Recurrent aphthous stomatitis (aphthae, canker sores)

Recurrent aphthous stomatitis typically starts in childhood or adolescence with recurrent small, round, or ovoid ulcers with circumscribed margins, erythematous haloes, and yellow or grey floors. It affects at least 20% of the population, and its natural course is one of eventual remission. There are three main clinical types:

- Minor aphthous ulcers (80% of all aphthae) are less than 5 mm in diameter and heal in 7-14 days.
- Major aphthous ulcers are large ulcers that heal slowly over weeks or months with scarring.
- Herpetiform ulcers are multiple pinpoint ulcers that heal within about a month.

Some cases have a familial and genetic basis, but most patients seem to be otherwise well. However, a minority have aetiological factors that can be identified, including stress, trauma, stopping smoking, menstruation, and food allergy.

Aphthae are also seen in haematinic deficiency (iron, folate, or vitamin B-12); coeliac disease; Crohn's disease; HIV infection, neutropenia, and other immunodeficiencies; Neumann's bipolar aphthosis, where genital ulcers may also be present; and Behçet's syndrome, where there may be genital, cutaneous, ocular, and other lesions. The mouth ulcers in Behçet's syndrome are often major aphthae with frequent episodes and long duration to healing.

In children aphthae also occur in periodic fever, aphthous stomatitis, pharyngitis, and cervical adenitis syndrome. This syndrome resolves spontaneously, and long term sequelae are rare. Corticosteroids are highly effective symptomatically; tonsillectomy and cimetidine treatment have been effective in some patients.

Diagnosis of aphthae is based on the patient's history and clinical features since specific tests are unavailable. A full blood picture (haemoglobin concentration, white cell count and differential, and red cell indices), iron studies, and possibly red cell folate and serum vitamin B-12 measurements and other investigations may help exclude systemic disorders, which should be suspected if there are features suggestive of a systemic background. Biopsy is rarely indicated.

Main systemic and iatrogenic causes of oral ulcers

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<td>Other dermatoses</td>
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Minors aphthous ulceration (top) and major aphthous ulceration (bottom)
Management—Predisposing factors should be identified and corrected. Chlorhexidine mouthwashes may help. Symptoms can often be controlled with hydrocortisone hemisuccinate pellets or trimacinolone acetonide in carboxymethyl cellulose paste four times daily, but more potent topical corticosteroids may be required. Systemic corticosteroids are best given by a specialist. Thalidomide is also effective but is rarely indicated.

Malignant ulcers
Oral carcinoma may present as a solitary chronic ulceration (see previous article).

Mouth ulcers in systemic disease
Ulcers may be manifestations of disorders of skin, connective tissue, blood, or gastrointestinal tract.

The main skin disorders are lichen planus, pemphigus, pemphigoid, erythema multiforme, epidermolysis bullosa, and angina bullosa haemorrhagica (blood filled blisters that leave ulcerated areas after rupture). In view of the clinical consequences of pemphigus, accurate diagnosis of oral bullae is important, and referral for direct and indirect immunofluorescence of biopsy tissue is often indicated.

Drug induced mouth ulcers
Among the drugs that may be responsible for mouth ulcers are cytotoxic agents, antithyroid drugs, and nicorandil.

Non-ulcerative causes of oral soreness
Erythema migrans (benign migratory glossitis, geographic tongue)
This common condition of unknown aetiology, which affects about 10% of children and adults, is characterised by map-like red areas of atrophy of filiform tongue papillae in patterns that change even within hours. The tongue is often fissured. Lesions can cause soreness or may be asymptomatic.

Management—There is no reliably effective treatment, although some have reported efficacy for zinc supplements. Similar lesions may be seen in Reiter's syndrome and psoriasis.

Burning mouth syndrome (oral dysaesthesia, glossopyrosis, glossodynia)
This condition is common in people past middle age and is characterised by a persistent burning sensation in the tongue, usually bilaterally. The cause is unclear, but response to topical anaesthesia suggests it is a form of neuropathy. Discomfort is sometimes relieved by eating and drinking, in contrast to the pain from ulcerative lesions, which is typically aggravated by eating.

Organic causes of discomfort—such as erythema migrans, lichen planus, a deficiency glossitis (related to deficiency of iron, folate, or vitamin B-12), xerostomia, diabetes, and candidiasis—must be excluded, but these are only occasional causes. More often there is an underlying depression, monosymptomatic hypochondriasis, or anxiety about cancer or a sexually transmitted disease. Burning mouth syndrome is more common in Parkinson's disease.

Management—Reassurance and occasionally psychiatric consultation, vitamins, or antidepressants may be indicated, but they are not reliably effective.

Desquamative gingivitis
Widespread erythema, particularly if associated with soreness, is usually caused by desquamative gingivitis. This is fairly common and is seen almost exclusively in women over middle age (see earlier article).
Orofacial pain

Most orofacial pain is caused by

- Local disease, especially dental, mainly a consequence of caries (see earlier article)
- Psychogenic states
- Neurological disorders (such as trigeminal neuralgia). Similar features are seen in the rare SUNCT syndrome (short lasting, unilateral, neuralgiform headache attacks with conjunctival injection and tearing)
- Vascular disorders (such as migraine). Recent evidence suggests that chronic pain may occasionally be related to thrombosis or hypofibrinolysis causing small areas of jaw ischaemia and necrosis; this has been termed neuralgia-inducing cavitational necrosis
- Referred pain (such as angina).

Psychogenic orofacial pain

This is an ill defined entity that includes burning mouth syndrome, atypical facial pain, atypical odontalgia, and the syndrome of oral complaints.

The pain is often of a dull, boring, or burning type of ill defined location. Most patients are women who are middle aged or older. They typically have constant chronic discomfort or pain, rarely use analgesics, sleep undisturbed by pain, have consulted several clinicians, have no objective signs and have negative investigations, and have recent adverse life events such as bereavement or family illness and also multiple psychogenic related complaints.

Management—Attempts at relieving pain by restorative treatment, endodontia, or exodontia are usually unsuccessful. Many patients lack insight and will persist in blaming organic diseases for their pain. Some patients are depressed or hypochondriacal and may respond to fluoxetine or dosulepin hydrochloride. However, many refuse drugs or psychiatric help. Those who will respond invariably do so early in treatment.

Atypical odontalgia presents with pain and hypersensitive teeth typically indistinguishable from pulpitis or periodontitis but without detectable pathology. It is probably a variant of atypical facial pain and should be treated similarly.

Temporomandibular joint pain-dysfunction syndrome

(myofascial pain-dysfunction syndrome, facial arthromyalgia)

This common disorder afflicts young women mainly. Symptoms are highly variable but characterised by

- Recurrent clicking in the temporomandibular joint at any point of jaw movement, and there may be crepitus especially with lateral movements
- Periods of limitation of jaw movement, with variable jaw deviation or locking but rarely severe trismus
- Pain in the joint and surrounding muscles, which may be tender to palpation.

Patients with a night time habit of clenching or grinding the teeth (bruxism) may awake with joint pain which abates during the day. In people who clench or grind during working hours the symptoms tend to worsen towards evening and sometimes have a psychogenic basis.

Different aetiological factors that have been implicated include muscle overactivity (such as bruxism and clenching), disruption of the temporomandibular joint, and psychological stress (such as anxiety and stressful life events). Precipitating factors may include wide mouth opening, local trauma, nail biting, and emotional upset. However, there is rarely one specific aetiology, and a combination of factors is often contributory. Occlusal factors do not in general seem to be important.
**Diagnosis**—This is arthropathy. Radiographic changes are uncommon, and arthrography or magnetic resonance imaging is seldom indicated.

**Management**—Most patients recover spontaneously, and therefore reassurance and conservative measures are the main management. These include rest, jaw exercises (opening and closing), a soft diet, and analgesics. If these are insufficient, it can be helpful to use plastic splints on the occlusal surfaces (occlusal splints) to reduce joint loading, heat, ultrasound treatment, anxiolytic agents, or antidepressants. A very small minority of patients fail to respond to the above measures and require local corticosteroid or sclerosant therapy, local nerve destruction, or, often as a last resort, joint surgery.

Crispian Scully is dean and Rosemary Shotts is honorary lecturer at the Eastman Dental Institute for Oral Health Care Sciences, University College London, University of London (www.eastman.ucl.ac.uk). The ABC of oral health is edited by Crispian Scully and will be published as a book in autumn 2000.

Crispian Scully is grateful for the advice of Rosemary Toy; general practitioner, Rickmansworth, Hertfordshire.

**Further reading**

- Van der Waal I. *The burning mouth syndrome*. Copenhagen: Munksgaard, 1990

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**A patient who changed my practice**

**The internet and a “small miracle”**

I have just returned from a mother’s day concert at my 6 year old’s primary school. The first “welcome” statement was made by a friend of hers, A, in a loud clear voice—a remarkable achievement for this particular child.

I have known A since she was a baby, watching her and two younger siblings pass through the baby clinic and reach normal development milestones. A was always a quiet child in company, but I was surprised to hear my daughter, in A’s class at nursery, remark one day, “You know Mummy, A never speaks at school.” There was no hint of developmental delay, and at home A interacted quite normally with her family. The transition to primary school saw a persistence of A’s determined silence—no verbal interaction at all with her class mates or her teachers, although her basic literacy and numeracy skills developed in line with those of her peers.

A’s parents were worried but remained patient and expectant—they at least knew she had much more normal behaviour at home. By the beginning of her second year at school A had still not uttered a single word at school. She also refused to remove her shoes and socks for physical education in front of others and would eat nothing all day, neither school dinners nor a packed lunch. A’s parents asked for a specialist review, wondering if any form of therapy would lead to more normal childhood interaction. No specific help resulted from this psychiatric assessment, but at least A now had a label “selective mutism,” and in today’s world a label by itself can begin to unlock doors.

I have to say my heart sank a little at the sight of sheets of internet printouts in A’s mother’s hand when she came in to see me a couple of weeks after the psychiatric clinic appointment. This was not because I resent patients accessing health information but because I don’t know how to judge the quality or validity of this information—I don’t know how to use it to make clinical decisions. But I was impressed. A series of case reports and parents’ stories of children seemingly similar to A who had responded dramatically to short courses of fluoxetine.

This drug is not licensed for children in the United Kingdom, but local drug information pharmacist was able to locate a small trial describing its use in children with selective mutism.1 A’s parents and I talked about the concerns relating to the use of unlicensed medication, and I thought that I had to share my reservations explicitly, drawing up a clear contract acknowledging our shared responsibility in using this drug on their child.

Within two weeks of starting the drug, A was recording taped messages for her teacher and beginning to participate in physical education. After six weeks she is chattering happily with her friends at school and has been to her first party alone. She has been transformed into a totally “normal” 6 year old, and her parents are slowly withdrawing the fluoxetine.

I am convinced that the use of fluoxetine has played a central part in this huge change in A’s behaviour, and I am equally sure that without the internet her parents could not have accessed this information. So if my heart sinks again at the sight of a patient’s internet printout, I will simply remind myself of the small miracle of A and suspend my prejudgment.

Dr Jolley general practitioner, North Shields