Otitis externa: review and clinical update

Abstract

Otitis externa can take an acute or a chronic form, with the acute form affecting four in 1 000 persons annually and the chronic form affecting 3-5% of the population. Acute disease commonly results from bacterial (90% of cases) or fungal (10% of cases) overgrowth in an ear canal subjected to excess moisture, or to local trauma. Chronic disease often is part of a more generalised dermatologic or allergic problem. Symptoms of early acute and most chronic disease include pruritus and local discomfort. If left untreated, acute disease can be followed by canal oedema, discharge, and pain, and eventually by extra-canal manifestations. Topical application of an acidifying solution is usually adequate in treating early disease. An antimicrobial containing ototopical is the preferred treatment for later-stage acute disease, and oral antibiotic therapy is reserved for advanced disease or those who are immunocompromised. Preventative measures reduce recurrences, and typically involve minimising ear canal moisture, trauma, or exposure to materials that incite local irritation or contact dermatitis.


Introduction

Otitis externa (OE) can take acute or chronic forms. The acute form is primarily of bacterial origin and annually affects four in 1 000 persons in the United States. The chronic form is commonly of a fungal or allergic origin or is the manifestation of dermatitides. It affects 3-5% of the population. Acute OE is unilateral in 90% of patients; it peaks in persons seven to 12 years of age, declines after 50 years of age, and often, is associated with high humidity, warmer temperatures, swimming, local trauma, and hearing aid or hearing protector use.

Pharmacology

The ear canal is a skin-lined, 2.5-cm cul-de-sac. The lateral 33% of the canal has a cartilaginous infrastructure covered by a layer of sebaceous and apocrine glands and hair. The medial canal has an osseous support devoid of adnexal structures. The glands produce a thin layer of cerumen that provides protection via a modestly antimicrobial lysozyme. The cerumen also has a pH level of 6.9, which discourages microbial growth. Insufficient cerumen predisposes to infection, whereas thickened cerumen, which can be caused by genetics, metabolism, or age, fosters retention of water and debris. Ear canals self-clean via a lateral epithelial migration towards the external os, a process that slows with age.

Before World War II, fungi were thought to be the primary cause of acute OE, but research by the US military in the South Pacific confirmed a predominantly bacterial process as the cause. Approximately 50% of bacterial cases involve Pseudomonas aeruginosa, followed in incidence by Staphylococcus aureus, and then various aerobic and anaerobic bacteria. The incidence of fungal infection is only 10%. Less than five per cent of acute disease cases can be attributed to furunculosis (usually staphylococcal),
Signs or symptoms lasting three months or longer indicate chronic OE. Although it can result from inadequately treated acute OE, it is usually of non-bacterial origin. A common cause of chronic OE is allergic contact dermatitis from such things as metal earrings, chemicals in cosmetics or shampoos, or the plastic in hearing aids or protection devices. Generalised skin conditions such as atopic dermatitis (i.e. eczema) or psoriasis can be difficult to treat in the narrow ear canal. Food sensitivity is a potential origin of chronic OE, associated with atopic dermatitis in up to 48% of patients.

A variant of chronic OE is a type IV cell-mediated hypersensitivity reaction to components of an ototopical used to treat acute OE. Dermatophytid eruptions also can manifest in the canal, and are reactions to haematogenously spread fungal by-products from a primary focus, e.g. nails, scalp, or vagina. A subtype of chronic OE is reported in 6-40% of patients, following insertion of a pressure equalisation tube that leads to middle-ear drainage, purulent or not, causing canal maceration.

**Evaluation**

The onset of acute OE generally is over a few days to a week. Initially, it is characterised by a scant, odourless secretion, as well as mild discomfort, and pruritus associated with modest erythema. If disease progresses to a moderate stage, the erythema increases and is joined by oedema (particularly in the thicker lateral canal), seropurulent secretions, and pain exacerbated by tragal pressure or movement of the auricle (Figure 1). In the severe stage pain is intense, the lumen of the canal obstructed, and extra-canal signs such as auricular cellulitis, parotitis, or adenopathy, are likely.

Evaluation includes taking a history of the onset and nature of symptoms; and of prior issues with skin disorders or local trauma, particularly via cotton swab or bobby pin. Patients with diabetes, those who are immunocompromised, and those with local circulatory insufficiency, e.g. from irradiation, are prone to rapid escalation from mild-to-severe manifestations.
Examination includes the ear canal, tympanic membrane, the auricle, and cervical nodes, as well as a cutaneous survey for other dermatologic manifestations. Cerumen or debris that blocks the canal is cleared to verify tympanic integrity. Concomitant and often erroneous diagnoses of acute OE and otitis media are common because the tympanic membrane in acute OE often is erythemic.8,9,19

Pneumatic otoscopy helps rule out middle ear disease. Debris usually can be cleared with a small Frazier suction tip (5 or 7 Fr) or an ear curette or spoon. Lavage is avoided until tympanic integrity is documented. Cerumen in acute OE tends to be hydrated by the otorrhoea, making it easier to remove. If the cerumen is thick or otherwise adherent, or if the patient is intolerant, a liquid ototopical (used overnight) can aid in loosening canal debris.

Treatment

Topical therapy for canal disease was described more than 3 000 years ago; astringents and alcohol were common.5,11,20 Two per cent acetic acid, sometimes diluted in half by 90-95% alcohol, is effective for prophylaxis against acute OE and, with or without the addition of a steroid, for the treatment of mild disease. The caveat is that these solutions can cause stinging and irritation if applied to highly inflamed skin.2-4,8,10,19,22

More advanced disease warrants the use of an ototopical that includes an antimicrobial agent (Table I). Regardless of the ototopical selected, penetration to the epithelium is mandatory; any obstruction should be cleared. Warming the ototopical to body temperature (e.g. by placing it in a shirt pocket or a warm room) before application helps the patient avoid the dizziness from caloric stimulation that cold liquid can incite. Instructing the patient to lie on his or her side with the affected ear up for a few minutes after the administration of ear drops aids migration to the medial canal. Having the patient or someone else pump the tragus a few times improves this process. There is wide variability in the self-administered dosing of ear drops, and relying on another person to place the drops or to perform the pump manoeuvre, better standardises the process.8,9,21,23

A cotton ball temporarily placed in the external os before the patient reassumes an erect position will absorb excess liquid. If the canal is narrowed at least 50% by oedema, placement of a wick (which can be anything from gauze to a preformed cellulose sponge) ensures ototopical access to the medial canal. When indicated, a return visit in two to three days for removal of the wick is necessary. Ototopical therapy usually should continue for five to 10 days depending on disease severity, or for three days after the last symptoms.

For other than mild acute OE, analgesics are appropriate treatment and can range from nonsteroidal anti-inflammatory drugs to mild narcotics.4,8,9,19,24 The ototopical should include an active antimicrobial agent, not just an inhibitor such as acetic acid.2,4,8,16,19,22 There are no randomised controlled

Table I: Common ototopicals for the treatment of otitis externa

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<tr>
<th>Agent</th>
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<tr>
<td>2.0% acetic acid, with or without steroid</td>
<td>Inexpensive. Effective for mild acute OE of bacterial or fungal origin in immunocompetent patients. Typically combined with a steroid if used for acute OE, but not for prophylaxis. It is best applied every four to six hours, and can sting and locally irritate.</td>
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<tr>
<td>2.75% boric acid or 90-95% isopropyl alcohol</td>
<td>Inexpensive. Effective for mild acute OE of bacterial or fungal origin in immunocompetent patients, but mainly applied as prophylaxis after swimming. Alcohol commonly mixed 50/50 with 2% acetic acid because the alcohol evaporates quickly, and affords a drier ear canal. It can sting and locally irritate.</td>
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<tr>
<td>Aminoglycoside</td>
<td>Moderately expensive. Usually an ophthalmic preparation, e.g. gentamicin, tobramycin, but effective for bacterial acute OE. It is best applied every six hours. It is minimally irritating, and has ototoxic potential.</td>
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<tr>
<td>Neomycin, polymyxin B, hydrocortisone</td>
<td>Relatively inexpensive. Effective for acute OE of bacterial origin, and best applied every six hours. It is neomycin sensitising in 5-18% of patients, and has ototoxic potential.</td>
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<tr>
<td>Fluoroquinolone, with or without steroid</td>
<td>Moderately expensive to expensive. Effective for bacterial acute OE. Best applied twice daily. Is minimally irritating, and infrequently sensitising. It is the only approved agent if tympanic membrane is perforated.</td>
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<tr>
<td>Steroid</td>
<td>Inexpensive to moderately expensive. Addresses underlying dermatitis, e.g. atopic dermatitis and psoriasis, if it is the cause of chronic OE. Solution easier to apply than cream. Application frequency depends on the agent selected. Cutaneous atrophy with prolonged use.</td>
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<tr>
<td>Tolnaftate or clotrimazole</td>
<td>Relatively inexpensive. Meant for general cutaneous use, but effective for acute or chronic OE of fungal origin (whether primary, or from suprainfection after antibacterial ototopical). The solution is easier to apply than cream, and is best applied every six hours. It is minimally irritating.</td>
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<tr>
<td>Various agents: chloroxylenol, an antiseptic; pramoxine, an anaesthetic; benzocaine, an anaesthetic</td>
<td>Few data published about effectiveness. All can incite contact dermatitis. Topical anaesthetics can mask the symptoms of advancing infection.</td>
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trials that directly compare oral with topical antimicrobial therapy, and few that compare ototopicals. However, the clinical effectiveness of ototopicals, which can achieve local tissue concentrations approximately 1 000 times that of systemic administration, is persuasive, and they have fewer incidents of systemic resistance or side-effects.3,8,17

Other topicals range from aminoglycosides (e.g. neomycin and gentamicin) to fluoroquinolones with or without a concomitant steroid. Ototoxicity from aminoglycosides has been associated with open middle-ear spaces or prolonged use. They are to be avoided if the eardrum is not intact.3,5,19,22 Neomycin tends to be sensitising in 5-18% of patients, and can incite contact dermatitis (Figure 2).1,4,22 Neomycin tends to be sensitising in 5-18% of patients, and can incite contact dermatitis (Figure 2).1,4,22,25,26 If acute OE fails to completely resolve, the physician should be aware that the preservatives, benzalkonium chloride, thimerosal, and propylene glycol, can incite local sensitisation.1,4,9,15,20,24-26

The fluoroquinolone preparations require only twice-daily dosing, and some are approved for use with a non-intact tympanic membrane. The addition of a steroid to a fluoroquinolone diminishes the symptomatic period by 0.8 days, and must be balanced against the small risk of the steroid acting as a sensitising agent.20,22Regardless of the ototopical selected, for other than mild cases, physicians should consider seeing the patient for one further visit to verify resolution of disease and to consider cleaning any remaining canal debris.

Ten per cent of acute OE is of fungal origin, but this percentage is higher in acute OE not fully responding to antibacterial drops. In this situation, the initial ototopical choice and the possibilities of a contact sensitivity or a fungal suprainfection, must be considered.3,4,5,16 Uncomplicated fungal infections commonly manifest with whitish cotton-like strands (e.g. Candida) with or without interspersed small black or white fungal balls (e.g. Aspergillus; Figure 3). Mixed bacterial and fungal infections are common after inadequate ototopical treatment of bacterial acute OE. Most fungal infections are mild and can be treated with a 2% acetic acid and/or a 90-95% alcohol solution. More established disease requires topical agents such as 1% clotrimazole or tolnaftate.3,16,27

Figure 2: An auricle and ear canal affected by acute otitis externa, and exhibiting a type IV hypersensitivity reaction to neomycin. Note the drainage pattern of the ear drops that have incited a cutaneous reaction on the ear lobe.

Figure 3: External auditory canal with Aspergillus overgrowth manifesting as a cottony matrix topped by small black balls.
Complications

When initial therapy fails, physicians should consider other possible reasons, including the accuracy of the initial diagnosis (Table II). In the unusual event of disease progression to extra-canal manifestations (e.g., auricular cellulitis, cervical adenopathy, or parotitis), addition of an oral antimicrobial is appropriate and culture is prudent. Oral antibiotics also should be considered for moderate acute OE in older patients; in patients who are immunocompromised; and in patients with diabetes, coexisting otitis media, or malignant external otitis. Malignant external otitis is an osteomyelitis of the ear canal (Figure 4). It often involves the adjacent mastoid and should be suspected when canal skin necrosis or granulation appear; the pain is disproportionate; the patient's temperature exceeds 102.2°F (39°C); or facial paralysis, vertigo, or meningeal signs occur.

A furuncle can occur in the ear canal de novo or as the result of acute or chronic inflammation. Cultures are taken at the time of incision and drainage, and ototopical and oral antibiotics are prescribed.

Prevention

Common precipitants of acute OE are moisture and trauma. One study showed that children with acute OE were more likely to have had their ears cleaned with cotton swabs, had wax removal, been swimming, or had pressure equalisation tubes inserted within the previous 10 days, compared with children without acute OE. The likelihood of acute OE can be reduced through daily prophylaxis with acidifying or alcohol drops during the at-risk period (e.g., swim season, scuba diving trip), use of a hair dryer on the lowest setting with or without a head tilt to aid fluid clearance after swimming or bathing, and the avoidance of cotton swabs. The use of hypoallergenic ear canal moulds (hearing aid or water exclusion varieties) with or without tight swim caps to diminish recurrent infections is controversial. Prevention is particularly important in patients who are immunocompromised; those with a systemic dermatologic condition, contact sensitivities to an ototopical, or who perspire excessively; or those for whom water sports are an avocation or common recreation.

The authors

J. David Osguthorpe, MD, is a professor in the Otolaryngology and Communicative Sciences Department and the Dermatology Department at the Medical University of South Carolina in Charleston. Dr Osguthorpe received his medical degree from the University of Utah in Salt Lake City. He trained in general surgery and in head and neck surgery at the University of California, Los Angeles Medical Center and in skull base surgery at the University of Zurich in Switzerland.

David R Nielsen, MD, is executive vice president and chief executive officer of the American Academy of Otolaryngology–Head and Neck Surgery. He received his medical degree from the University of Washington and completed his residency training in otorhinolaryngology at the University of Iowa.

Table II: Considerations when otitis externa fails to respond to initial therapy

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<td>Contact dermatitis (original issue or reaction to ototopical)</td>
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<td>Failure to adhere to preventive measures, e.g. temporary cessation of swimming</td>
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<tr>
<td>Faulty, or infrequent administration of ototopical</td>
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<tr>
<td>Immunosuppression (requires prolonged therapy and possibly oral antibiotics)</td>
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<tr>
<td>Inadequate penetration of ototopical because of canal debris or narrowing</td>
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<tr>
<td>Malignant external otitis (requires intravenous antibiotics and debridement)</td>
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<tr>
<td>Misdiagnosis, including dermatological conditions (e.g. control systemic psoriasis and seborrhoea), dermatophytid reaction, cancer, self-instrumentation trauma</td>
</tr>
<tr>
<td>Ototopical not effective against bacteria or fungi involved (or it is a mixed bacterial and fungal problem)</td>
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Figure 4: Malignant external otitis, with pus draining from the necrotic ear canal and underlying osteomyelitic bone. The adjacent auricle demonstrates the swelling and loss of cartilaginous architecture characteristic of chondritis.
University of Utah School of Medicine, and completed a residency in otolaryngology at the University of Utah Affiliated Hospitals, both in Salt Lake City. Dr Nielson also completed a fellowship in otology at Providence Hospital, Southfield, Mich.

Author disclosure

The authors have nothing to disclose.

References