A Case Series of Malignant Otitis Externa Mimicking Malignancy

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ABSTRACT
Malignant otitis externa is an inflammation of the external auditory canal with preceding osteomyelitis of the temporal bone and the adjacent structures that could be potentially lethal. Malignant otitis externa may present with cranial nerve involvements and massive spread of disease mimicking nasopharyngeal carcinoma or any other malignancies on imaging. Two elderly patients who presented with severe otalgia and significant facial nerve palsy and lower cranial nerve palsies showing extensive spread of disease are reported in this case series. They both had resolution of disease after a prolonged course of antibiotics and cortical mastoidectomy for disease clearance in one of them.

KEYWORDS
malignant otitis externa; Pseudomonas aeruginosa; osteomyelitis; cranial nerves palsies

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INTRODUCTION

Malignant otitis externa (MOE) is an aggressive infection of the external ear canal and temporal bone. MOE was first reported by Toulmouche in 1838 (1). Another name for MOE is necrotizing otitis externa or osteomyelitis of the temporal bone. Patients with MOE usually presents with painful ear discharge with hearing loss, tinnitus and throbbing headaches that is not responding to medical treatment. The ear canal is edematous with granulation tissue at the osteocartilaginous junction on otoscopic examination. The usual causative organism is Pseudomonas aeruginosa and the infection can be lethal with extension of the infection to the skull base and central nervous system causing cranial nerve palsies, meningitis and brain abscesses. The commonly involved cranial nerve is the facial nerve as it exits the stylomastoid foramen, subsequently the glossopharyngeal, vagus and accessory nerves as they exit the jugular foramen and the hypoglossal nerve as it exits the hypoglossal canal. In petrous apex involvement, the trigeminal and abducens nerve can be affected and in cavernous sinus and clivus involvement, the olfactory, trochlear and abducens nerve may be affected as well. We report two cases of malignant otitis externa in two elderly female patients presented with lower cranial nerve palsies and extensive spread of disease described on computed tomography (CT).

CASE REPORT 1

This patient is a 50-year-old lady who presented with dysphagia, lethargy and hoarseness for 2 weeks. She was having intermittent episodes of left ear discharge and pain for 3 months with throbbing headache and vomiting episodes. She was previously treated for acute otitis externa.

She appeared comfortable under room air with stable vital signs. On otoscopic examination, the left ear canal was filled with pus, oedematous and there was a mass occupying the canal arising from the posterosuperior wall. The mass was smooth surfaced and didn’t bleed on probing while the left tympanic membrane couldn’t be visualised. The right ear was normal. Her intraoral examination revealed the curtain sign with left palatopharyngeal weakness. However, there was no trismus. There was a firm fullness felt on the left preauricular region measuring 7 × 6 cm but there was no signs of inflammation. She also had left facial nerve palsy House-Brackmann Grade 2. Flexible nasopharyngolaryngoscopy showed pooling of saliva and the left vocal cord was immobile in paramedian position with a phonation gap consistent with left recurrent laryngeal nerve palsy. Her pure tone audiometry showed left moderate to profound mixed hearing loss and right mild to severe sensorineural hearing loss.

The total white blood cell count was raised up to 16.5 × 10^9/L. Both ESR and C-reactive protein were also raised with a reading of 123 mm/hr and 40 mg/L respectively. She had electrolytes imbalance with hyponatraemia and hyperkalemia secondary to her poor oral intake.

She was admitted for initiation of intravenous antibiotics, corrections of electrolytes and dietary optimization. She was started on ryles’ tube feeding and was referred to the Endocrine team to manage her diabetic status.

This lady had underlying diabetes mellitus, on insulin regime with a diabetic foot ulcer and hypertension. Her foot ulcer was fairly clean and was dressed daily. Her sugar control in the ward was below the range of 10 mmol/L and her HbA1c showed poor sugar control with reading of 14.5%.

Swab was obtained from the left ear for culture and sensitivity and it grew Pseudomonas Aeruginosa. Biopsy was taken from the left external ear canal mass and was later reported as inflammatory granulation tissue by the pathologist.

She was started on intravenous ceftiraxone but once the culture grew Pseudomonas Aeruginosa, her antibiotics were upgraded to meropenem for better blood brain barrier penetration and was given for 2-weeks. She was also given gentamicin with betamethasone ear drops.

She underwent high resolution computed tomography (HRCT) to see the extent of the disease. As shown on Figure 1, there was a soft tissue density in the left middle ear, part of the external ear canal and the left mastoid air cells. The ossicles and incudomalleolar joint were intact. Even though the first genu and the tympanic segment of the facial nerve were not well visualized, the rest of the facial nerve course were normal.

The HRCT also showed features of skull base osteomyelitis with abscesses in the left parapharyngeal extending to left fossa of rosenmuller, left torus tubarius, retropharyngeal, masticator space and left temporomandibular joint as shown in Figure 2.

There was an intracranial extension with meningeal enhancement in the left temporal region with bony erosions at the petrous apex, left anterior occipital condyle, squamous part of temporal bone, sphenoid bone, tegmen

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Fig. 1 HRCT temporal, axial cuts showing soft tissue density in the left mastoid air cells (A). The ossicles were intact (B). The left facial nerve course was normal (C).
After 2 weeks of intravenous meropenem, she was given oral ciprofloxacin for 4 weeks. Her symptoms improved as her otalgia and otorrhea resolved. Although her hoarseness persisted, she was well and was discharged home with oral ciprofloxacin.

She was seen again in clinic 2 weeks after and she was still pain free. Her ear symptoms such as otalgia and otorrhea improved. She was feeding via ryles tube with evident aspiration. However, there was a new finding of left hypoglossal nerve palsy showing significant muscle wasting of the left lateral tongue deviating to the left. The previously seen left facial nerve palsy reverted back to normal during examination. The other pre-existing left glossopharyngeal and left vagus nerve palsies showed no improvement. She was referred to the speech therapist for swallowing exercise. She was planned for a bone scan to monitor her disease progression and treatment response but patient refused for any further intervention.

CASE REPORT 2

The second case reported is a 53-year-old lady who presented with severe right ear otalgia for almost 5 months, throbbing in nature. She described it as a constant pain associated with right sided headache affecting her daily routine. The pain was relieved by oral analgesia but it only lasted a while. She also complaint of right ear fullness with tinnitus and reduced hearing. Otherwise she denied of any fever and there was no giddiness nor vertigo. The symptoms were not preceded by any respiratory infection or trauma.

She visited the local clinics multiple times and was given tropical ear drops and oral antibiotic but her symptoms were not relieved. She had underlying diabetic and hypertension, well controlled on oral medications and insulin injections.

She appeared comfortable under room air with no acute respiratory distress. There was right facial nerve palsy House-Brackmann Grade 2. On otoscopic examination, the right external ear canal was filled with a mass and the tympanic membrane could not be visualized; however the left ear canal was clear with intact tympanic membrane. She also had right glossopharyngeal and right vagus nerve palsies with curtain sign elicited intraorally and uvula deviated to the left. Her gag reflex was absent. The other cranial nerves examinations were unremarkable. There was a right level II lymph node measuring 2 × 1 cm, firm in consistency with well demarcated borders. Her pure tone audiometry showed right mild to severe mixed hearing loss and left mild to profound sloping sensorineural hearing loss.

She was admitted for intravenous ciprofloxacin given for 2 weeks, ofloxacin topical ear drops and pain control. During admission, her ESR was reported as 127 mm/hour and C-reactive protein was positive with a reading of 48 mg/L. Her total white blood cell count was 12.04 × 10⁹/L. A biopsy was obtained from the right ear canal mass reported as compatible with malignant otitis externa. Nasal endoscopy showed an enlarged right torus tubarius obliterating the right fossa of Rosenmuller. Biopsy of the right torus tubarius revealed no malignancy. Other findings were normal. Swab for culture and sensitivity from the right ear canal was reported as negative for any growth of microorganism.

Magnetic resonance imaging (MRI) was done towards the end of the 2 weeks of admission to look for any evidence of cholesteatoma or malignant lesions. As shown in Figure 4, there was a diffuse enhancing soft tissue lesion inferior to the right temporal lobe involving the right masticator space, right carotid space, right parapharyngeal space, pharyngeal mucosal space and retropharyngeal space. The lesion extended till the dural layer superiorly, inferiorly until the upper oropharynx and medially to the nasopharynx.
She was given another 6 weeks of oral ciprofloxacin to complete 8 weeks all together. Her symptoms improved clinically with reducing pain and resolved ear discharge but she underwent right cortical mastoidectomy to obtain a tissue biopsy as we were still uncertain of her diagnosis. Intra-operatively there was sagging of the supero-posterior wall with intact tympanic membrane. When the tympanomeatal flap was raised, there was a soft tissue mass seen within the right mastoid antrum and right middle ear which was later reported as chronic inflammation. The middle ear mucosa appeared thickened with mucoid discharge.

Post operatively, she was doing well. Her pain was well controlled and there was no more ear discharge. There was improvement in the cranial nerve functions with right facial nerve House-Brackmann Grade 1 and uvula was central during intraoral examination. Her total white blood cell count dropped to 9.8 × 10^9/L and her ESR reading was decreasing to 111 mm/hour with a negative reading of C-reactive protein. Her pure tone audiometry also showed improvement with right moderate mixed hearing loss and left mild to profound sensorineural hearing loss.

DISCUSSION

Malignant otitis externa is a fulminant osteomyelitis involving the external auditory canal and the skull base following an episode of external ear infection. Being termed as a misnomer, it behaves aggressively like a malignancy. Patient in old age or patients with diabetis mellitus and debilitating condition are at higher risk for MOE (1).

Patients with MOE presents with severe otalgia and throbbing headaches just like the cases reported in this series. The intensity of pain is usually measured with the visual analogue scale (VAS). Both our patients had significant pain score according to the VAS. They usually show no improvement to local treatment and due to the ongoing infection, they may also have a discharging ear. The most commonly grown organism in MOE is Pseudomonas aeruginosa (95%) (2) as reported in our first case report.

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The otoscopic examination will reveal an edematous ear canal with granulation tissue at the osteocartilaginous junction. Besides the granulation tissue at the osteocartilaginous junction, there can also be a mass in the external ear canal. Biopsy of this mass is important to help to exclude other pathologies such as malignancies or cholesteatoma (3, 4) but most frequently, these biopsies obtained may not be significant (5) delaying the establishment of diagnosis.

MRI and CT scans are used to determine the anatomical extent of the disease and intracranial complications if any. Features of MOE on CT is reduced skull base density and bony erosions. Bony erosions of the skull base and petrous apex as seen in the first case report raises the suspicion of malignancy. In MOE, bony erosions on CT is only evident when at least 30% demineralisation of bone has happened which is usually in the later stage of the disease (6). MRI on the other hand determines the soft tissue changes, exact location and extent of the disease. MRI is highly sensitive but not so specific for MOE. The Technetium Tc 99 methylene diphosphonate bone scanning is the most useful imaging tool to evaluate the positive findings based on binding of the osteoblasts (3) but they may be positive in cases of malignancy as well (7).

As the disease progresses, patients can rarely present with cranial nerve involvement. Intracranial involvement can potentially alter the patient’s mental status and even cause mortality. Cranial nerve involvement indicates a poor prognosis of the disease leading to death due to the complications (8, 9). The spread of infection from the external ear is through the fissures of Santorini and the osteocartilaginous junction. The lower cranial nerves are commonly affected in the skull base osteomyelitis as reported in both our cases. The other cause of lower cranial nerve palsy in these areas is the nasopharyngeal
carcinoma (NPC) that is related to the postriorlateral spread of the disease as well as the jugular foramen infiltration (10).

The facial nerve is the first and most common to be affected in MOE as it exits the stylo mastoid foramen (7) and both our cases had facial nerve palsy on presentation. This is followed in order by the glossopharyngeal, vagus and accessory nerves as they exit the jugular foramen and the hypoglossal nerve as it exits the hypoglossal canal (11). The trigeminal and abducens nerve can be affected as well if there is petrous apex involvement. The spread of infection to the cavernous sinus and clivus may cause the olfactory, trochlear and abducens nerve palsy (12). If the treatment is initiated early, the cranial nerve palsies in MOE can be reverted back to normal. Both our cases had facial nerve and lower cranial nerves involvement and they showed improvement after treatment suggesting MOE rather than malignancy.

Navin reported 43% of cranial neuropathies in a retrospective analysis of 23 MOE cases where he mentioned that 60% cases were with facial nerve palsy, 30% with lower cranial nerves palsy in combinations of glossopharyngeal, vagus, accessory and hypoglossal nerve and 10% had an extended cranial nerve palsies involving the facial nerve, lower cranial nerves and other cranial nerves. Out of the 10 cases reported, 4 died due to unrelated causes, 5 had resolution of disease but no cranial nerve palsy improvement and 1 required prolonged treatment of oral ciprofloxacin. And it was speculated that facial nerve palsy does not resolve after treatment because of its longer course of involvement compared to the others. This case series reported a recovery rate of 87% with 0% of mortality (13). In another journal, Ethan et al. reported that involvement of facial nerve palsy showed a significant extension of disease to the parapharyngeal and nasopharyngeal region in addition to the mastoid on computer tomography (4) just like the two cases reported in this series.

The choice of antibiotics for treatment is from the fluoroquinolone group namely ciprofloxacin that can attain high level of bone and soft tissue penetration, and third generation cephalosporins such as ceftazidime (2) and aminoglycosides such as gentamicin may be used in ciprofloxacin-resistant patients.

Surgical options are kept in view for local debridement, abscess drainage and bony sequestrum removal. Even though our patient in the first case was given intravenous meropenem for 2 weeks, she was then given a long 4 weeks course of oral ciprofloxacin and for our patient in the second case, she was treated with 8 weeks course of ciprofloxacin and they both responded well to the treatment.

There may be many instances where MOE was mistaken for NPC because both these diseases invade the bony structures aggressively. Both the cases in this series had imagings with involvement of fossa of rossenmuller mimicking NPC. Goh et al reported 14 cases with skull base osteomyelitis where the imagings showed a nasopharyngeal bulge involving the fossa of rossenmuller (14). He also mentioned that in MOE, the soft tissue enhancement is greater or equal to the mucosa and the spread of disease is along the fascial plane preserving the architecture whereas it was vice versa in NPC. The presence of abscesses in imagings should also lead to infection rather than malignancy.

Another significant distinguishing factor of MOE is the lateral structure involvements such as TMJ and parotid gland involvement. NPC usually does not extend laterally except in late cases due to perineural invasion or contiguous spread along the auriculotemporal nerve (15).

Clival involvement on MRI is shown for both MOE and malignancy (16). But for MOE specifically, the clival involvement is shown enhanced with narrow space hypointensity on T1 weighted images but hyperintense on T2 weighted images with enhancement of dura, parapharyngeal fat plane effacement and soft tissue mass in skull base (17). Ozgen et al reported the important function of diffusion-weighted sequence on MRI in differentiating MOE and malignancy (18). He said that malignant tumours such as NPC and lymphomas have very low apparent diffusion coefficients (ADC), while benign lesions such as MOE have higher ADC.

The raise of inflammatory markers such total white blood cell count, ESR and CRP should also lead the suspicion of infection rather than malignancy. MOE patients may have a reactive node but not in typical fashion like NPC. NPC patients commonly present with nodal metastasis starting from the retropharyngeal region extending to level II, III, IV, V and supraclavicular.

MOE may spread causing further complications such as lateral sinus thrombosis, internal jugular vein thrombosis, ophthalmoplegia and blindness which can mimic malignancy. It is also important to correlate with the clinical history and examination where in MOE, patients commonly with underlying diabetes mellitus presents with ear pain and ear discharge with ear swabs growing Pseudomonas Aeruginosa.

CONCLUSION

Although MOE with cranial nerve involvement indicates extensive progression of the disease, it doesn’t worsen the prognosis of the disease itself. In view of aggressive features of disease in imaging, it is important to differentiate MOE from malignancy as early intervention with long-term high dose antibiotics may lead to resolution of disease although the cranial nerve functions might not be reverted back to normal in all cases.

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