From Uncommon Infection to Multi-Cranial Palsy: Malignant External Otitis Insights

Salvatore Ferlito1, Antonino Maniaci1, Milena Di Luca1, Calogero Grillo1, Lorenzo Mannelli2, Marco Salvatore2, Ignazio La Mantia1, Giacomo Spinato3,4, and Salvatore Cocuzza1

Abstract

Purpose: The progression of the otitic infectious process toward diseases of particular severity is often unpredictable, just as it is challenging to manage the patient over time, even after the apparent resolution of the disease. We aim to define a radiological reading key that allows us to correctly and promptly treat the disease, avoiding the possible severe complications.

Methods: We conducted a retrospective study of 13 cases of basal cranial osteomyelitis (SBO) due to malignant external otitis, by the ENT Department of the University of Catania. Through a standardized approach and following the latest guidelines, we have evaluated all patients performing a standardized and personalized radiological protocol according to the stage of the patient’s pathology and modulating the treatment consequently.

Results: Clinical signs have been observed such as otorrhea (100%), otalgia in 13/13 patients (100%), granulations in external auditory canal (100%), preauricular cellulitis in 9/13 patients (69%) headache 6/13 cases (46%), dysphonia 4/13 cases (31%). HRCT of the temporal bone proved useful in identifying even minimal bone lesions in 13/13 (100%) while improving MRI in vascular and nervous involvement, although in 1/13 patient with nerve palsy clinical symptomatology preceded radiological evidence. The 99mTc 3-phase planar bone scintigraphy was positive for SBO in 9/13 cases (69%) during the initial phase and, in 100% of the cases in images delayed to 2-3 hours. Subsequent checks up to 1 year, using the Ga 67 scintigraphy, excluded the presence of recurrences in 100% of patients.

Conclusion: The osteomyelitis of the base of the skull is a severe complication of malignant external otitis, often not always easily diagnosed. Recurrence can occur up to 1 year after stopping therapy. Imaging techniques such as Tc and MRI are relevant for the initial diagnostic approach and the staging of the pathology and its complications. Nuclear medicine imaging plays a fundamental role in the evaluation of related osteoblastic activity, especially in the remission phase of the disease.

Keywords
necrotizing otitis externa, skull base, radiological findings, osteomyelitis

Introduction

Necrotizing “malignant” external otitis (MEO) is a rare necrotizing infection caused by Pseudomonas Aeruginosa, which spreads from the squamous epithelium of the external auditory canal to the surrounding tissues, first described by Meltzer in 1959.1 Patients with otalgia and otorrhea persistent despite ambulatory therapy for several weeks. Typically, the infection spreads from the EAC through the base of the skull, causing bone erosion with the development of soft tissue granulomas.23 The treatment of the disease provides an initial phase of aggressive pharmacological treatment.

Radiodiagnostic imaging methods perform a fundamental role in the global evaluation of prognosis and the treatment of malignant external otitis and in particular in its complications of the skull base. The imaging techniques related to the

1 Department of Medical and Surgical Sciences and Advanced Technologies
2 “GF Ingrassia,” ENT Section, University of Catania, Catania, Italy
3 IRCCS SDN, Naples, Italy
4 Department of Neurosciences, Section of Otolaryngology and Regional Centre for Head and Neck Cancer, University of Padova, Treviso, Italy
5 Department of Surgery, Oncology and Gastroenterology, Section of Oncology and Immunology, University of Padova, Padova, Italy

Corresponding Author:
Antonino Maniaci, ENT Department of University of Catania, Via Santa Sofia, Catania 95100, Italy.
Email: tnmaniaci29@gmail.com
initial diagnostic approach are CT and MRI, with the first one, to evaluate erosion and demineralization. In contrast, the details of the soft tissues and the base of the skull are involved using the second one. Other diagnostic methods support the diagnosis of SBO are Tc-99 m-MDP, related to the increase in osteoblastic activity of the disease.4

The natural course of the otitic infectious process towards diseases of particular gravity is often unpredictable, putting a strain on the choice of the clinician who faces the pathology, among other things of rare frequency. Moreover, even the post-therapeutic management of the patient in apparent healing is difficult, considering the insidious reurrences. Our goal is to define a radiological reading key allowing us to treat the disease correctly and promptly, avoiding possible and severe complications. Through the analysis of radiological criteria correlated with clinical data, it is indeed possible to define the correct approach of the otitic patient.

Material and Methods

We conducted a retrospective study of 13 cases of cranial osteomyelitis (SBO) due to malignant external otitis, from January 2000 to December 2019, at the ENT Department of the University of Catania. The average age range was 54 to 82 years, composed of 10/13 men (77%) and 3/13 women (23%). Regarding risk factors, 9/13 patients (69%) had diabetes mellitus, 3/13 patients (23%) had undergone chemotherapy and, 1 patient (8%) underwent bone marrow transplantation. All included patients presented characteristic symptomatology for an average time of about 2 months ± 1, not responding to the conventional antibiotic therapy. The age, sex and medical history of these patients have been recorded. Initially, the patients underwent a thorough clinical examination, demonstrating clinical features such as deep pain (temporal, parietal, retroauricular, retroorbital), headache, intermittent otorrhea and spiking fever. Moreover, were reported preauricular cellulitis and mastoiditis, hearing floor granulation and cranial nerve involvements. The evaluation of the systemic inflammatory status was performed utilizing hematological examinations (routine blood tests, HGT analysis, VES monitoring, and PCR). The auricular swab and biopsies from the granulation tissues were analyzed by histopathological examination (HPE). An immuno-absorbent test linked to the enzyme for human immunodeficiency virus (HIV) was also performed to analyze the status of patients’ immune systems.

In particular, in each subject at the end of the clinical evaluation, a Radiodiagnostic protocol was performed consisting of high-resolution computed tomography (HRCT) of the temporal bone (axial and coronal sections) and Magnetic Resonance Imaging (MRI) (axial and coronal sections).

While the Tc-99 m-MDP test was performed to strengthen the diagnosis of the osteomyelitis process immediately after the 2 procedures mentioned above, the Ga 67 scintigraphy was executed within 1 month of stopping antibiotic therapy in all cases.

Besides, each patient was evaluated by 2 independent radiologists with at least 15 years of experience in the Head and Neck district.

The Positron Emission Tomography (PET) was performed in 2 patients as an alternative follow-up method.

The final resolution of the infectious process was expressed only after 12 months of well-being from the end of the treatment with concomitant compliance with the established Radiodiagnostic criteria with the integration of clinical symptoms and laboratory tests.

Results

The most common clinical features observed were: otorrhea (100%); otalgia in 13/13 patients (100%); granulations in the EAC (100%); preauricular cellulitis in 9/13 patients (69%); headache 6/13 cases (46%); dysphonia 4/13 cases (31%).

All patients had cranial neuropathy at the time of clinical diagnosis. Facial nerve palsy was observed in 12/13 cases (92%): mainly associated with other cranial nerves paralysis 7/13 (54%), the isolated involvement resulted only in 5/13 (38%) (Table 1).

Moreover, 3/13 patients presented IX, X, XI and XII cranial nerve palsy (23%) while 1/13(8%) patient, who was subjected to chemotherapy, showed isolated VI CN paralysis.

Furthermore, 1/13 (8%) patient who performed Bone Marrow transplantation, showed an associated involvement of the VI and VII CN (Table 2).

The involvement of the temporomandibular joint was observed in 3/13 cases (23%). Only 1/13 patient (8%), presenting a case of paralysis of the VII, developed meningitis, antibiotics bilateral coceleopathy, arteritis and thrombosis of the internal carotid.

HRCT of the temporal bone showed in all cases common findings such as soft tissue density in the EAC and the middle ear, the opacification of mastoid cells of air with cortical bone erosion and bone seizure in all patients (100%) (Figure 1). Moreover, HRCT revealed in 2/13 patients (23%) the erosion of the EAC floor extending anteriorly in the temporomandibular joint, while in 1/3 patient with a positive temporomandibular clinic did not demonstrate visible lesions.

In the presence of clinical symptoms characteristic of cranial nerve paralysis, the initial MRI performed showed the corresponding radiological involvement in 11/13 (85%) patients. The subsequent 1-month MRI check-in, the remaining 2/13 (15%) cases for imaging was positive.

The MRI showed in 1/3 patients with temporomandibular symptoms but without evident lesions to the CT, the anterior diffusion along the peri mandibular adipose planes of the infectious process (Figure 2).

Planar triple-phase bone scintigraphy 99mTc was positive for SBO in 9/13 cases (69%) during the initial phase, as there was a blood flow increased in the perfusion phase (Figure 3). Moreover, increased soft tissue tracer absorption in the blood pool resulted. Delayed images at 2-3 hours, showed 100% of
The cases achieved the diagnosis of SBOM for an increase in tracer absorption in the temporal region of the skull.

Ga 67 scintigraphy was performed in all cases within 1 month of stopping antibiotic therapy (Figure 4). Gallium citrate was located in areas of osteomyelitis due to bacterial or granulocyte incorporation and to binding with transferrin and lactoferrin at infection sites. Due to the high permanence of the gallium in the soft tissues, although the first images were acquired already 6 hours after the injection, the investigation had to be continued for 24-48 and sometimes 72 hours to obtain sufficiently informative images. Images with Ga 67 were qualitatively worse than those with 99mTc due to the more considerable emission energy but thanks to the localization mechanisms, it allowed to recognize chronic osteomyelitis more easily in 13/13 (100%) patients.

The PET showed the presence of tracer uptake in the temporal site at 2/2 (100%) studied.

### Table 1. Clinical Features of SBO Patients.

| Pt. No | Sex, Age (Y) | Risk factors | Clinical features | Nerve palsy | Ear Swab & HPE | Treatment |
|--------|--------------|--------------|------------------|-------------|----------------|-----------|
| 1      | M, 77        | Diabetes mellitus type II | Otorrhea, granulation in EAC, otalgia, headache, preauricular cellulitis | VII | P aeruginosa | Ceftazidime 2 g tds IV for 6 wk; ciprofloxacin 500 mg bd for >6 mth |
| 2      | M, 65        | Chemotherapy | Otorrhea, granulation in EAC, otalgia, dysphonia | VII and X | S. Aureus | Ceftazidime 2 g tds IV for 6 wk; ciprofloxacin 500 mg bd for >6 mth |
| 3      | M, 69        | Diabetes mellitus type II | Otorrhea, granulation in EAC, otalgia, headache, preauricular cellulitis | VII | P aeruginosa | Ceftazidime 2 g tds IV for 6 wk; ciprofloxacin 500 mg bd for >6 mth |
| 4      | M, 76        | Diabetes mellitus type II | Otorrhea, granulation in EAC, otalgia, headache | VII | P aeruginosa | Ceftazidime 2 g tds IV for 6 wk; ciprofloxacin 500 mg bd for >6 mth |
| 5      | F, 72        | Chemotherapy | Otorrhea, granulation in EAC, otalgia, preauricular cellulitis | VI | No organism identified | Ceftazidime 2 g tds IV for 6 wk; ciprofloxacin 500 mg bd for >6 mth |
| 6      | M, 82        | Diabetes mellitus type II | Otorrhea, granulation in EAC, otalgia, dysphonia, preauricular cellulitis | VII, XI, XII | P aeruginosa | Ceftazidime 2 g tds IV for 6 wk; ciprofloxacin 500 mg bd for >6 mth |
| 7      | M, 59        | Diabetes mellitus type II and HIV | Otorrhea, granulation in EAC, otalgia | VI, VII | S. Aureus | Ceftazidime 2 g tds IV for 6 wk; ciprofloxacin 500 mg bd for >6 mth |
| 8      | M, 54        | Bone Marrow transplantation (LMA) | Otorrhea, granulation in EAC, otalgia, headache, preauricular cellulitis | VI and VII | P aeruginosa | Ceftazidime 2 g tds IV for 6 wk; ciprofloxacin 500 mg bd for >6 mth |
| 9      | F, 80        | Diabetes mellitus type II | Otorrhea, granulation in EAC, otalgia, preauricular cellulitis | VII, XI, XII | P aeruginosa | Ceftazidime 2 g tds IV for 6 wk; ciprofloxacin 500 mg bd for >6 mth |
| 10     | M, 79        | Diabetes mellitus type II | Otorrhea, granulation in EAC, otalgia, dysphonia, preauricular cellulitis | VII | S. Aureus | Ceftazidime 2 g tds IV for 6 wk; ciprofloxacin 500 mg bd for >6 mth |
| 11     | F, 68        | Diabetes mellitus type II | Otorrhea, granulation in EAC, otalgia, preauricular cellulitis | VII | P aeruginosa | Ceftazidime 2 g tds IV for 6 wk; ciprofloxacin 500 mg bd for >6 mth |
| 12     | M, 74        | Chemotherapy | Otorrhea, granulation in EAC, otalgia, dysphonia, headache, preauricular cellulitis | VII, XI, XII | P aeruginosa | Ceftazidime 2 g tds IV for 6 wk; ciprofloxacin 500 mg bd for >6 mth |
| 13     | M, 65        | Diabetes mellitus type II | Otorrhea, granulation in EAC, otalgia, dysphonia, | VII, X | P aeruginosa | Ceftazidime 2 g tds IV for 6 wk; ciprofloxacin 500 mg bd for >6 mth |

Pt no = patient number; y = years; M = male; CN = cranial nerve; tds = 3 times a day; IV = intravenously; wk = weeks; bd = twice a day; mth = months; EAC = external auditory canal; F = female.

### Table 2. Different Ways of Spreading the Infective Process and the Organs Involved.

| Direction | Site | Diffusion way |
|-----------|------|---------------|
| ANTERIOR  | • TMJ | SANTORINI FISSURE |
|           | • PAROTID GLAND | |
|           | • MASTATORY SPACE | |
| POSTERIOR | • MASTOID | PETRO-TIMPANIC SUTURE |
| MEDIAL    | • VERTEBRAL SPACE | HEARING TUBA |
|           | • PARAPHARYNGEAL SPACE | FARINGO-BASILARY BAND |
| SUPERIOR  | • SKULL BASE | VASCULAR-NERVOUS FORAMS |
| INFERIOR  | • CI | PREVERTEBRAL ABSCESS |
|           | • OCCIPITAL BONE | HEMATOGENOUS WAY |

The PET showed the presence of tracer uptake in the temporal site at 2/2 (100%) studied.
At subsequent checks at 1 month and 3 months, all patients presented complete regression of the infectious process and radio diagnostic evidence but with the presence of typical radiological sequelae. Despite the effective treatment of their infection, all patients did not resolve cranial nerve dysfunction disorders, presenting significant neurological sequelae, in particular regarding VII function.

Refer to the below figures:

**Figure 1.** CT scan: the evaluation of facial nerve involvement is possible through high-resolution CT. On left image, negative imaging but positive clinical data in a patient with facial nerve paralysis. On right image (after a month), the erosion of the cortical bone and a considerable enlargement in the mastoid portion of the facial nerve are revealed.

**Figure 2.** MRI Tw2 FFE: wide anterior diffusion in the subtemporal adipose planes and in the soft tissues that reaches the site of the bony structures of the temporomandibular joint and posteriorly into the mastoid portion of the petrous bone.

MOE occurs predominantly in elderly diabetics or immunocompromised, and the most common pathogen is Pseudomonas Aeruginosa, but cases of Staphylococcus, Candida and Aspergillus have also been reported.

Several nosological criteria have emerged in the literature to confirm the diagnosis of malignant external otitis. Levenson’s criteria are used for diagnosis of necrotizing otitis externa, including refractory otitis externa, severe nocturnal otalgia, and purulent otorrhea associated with Pseudomonas infection and granulation of the EAC in an immuno-compromised or diabetic patient and positive temporal bone scanning with Tc99.

Although in most cases the patient has a disproportionate otalgia compared to the clinical picture, nonspecific symptomatology characterized, for example, by persistent headache and development of cranial neuropathy is possible.

MOE is the end stage of a severe infection that originates from the EAC and could progress through cellulitis to chondritis, osteitis and at the end of osteomyelitis. Once periostitis develops, the infection progresses rapidly across the skull base, replacing the progressively compact bone with granulation tissue. As a result, facial and other cranial nerve palsies can occur. Parotitis and trismus secondary to masseter myositis and involvement of the temporomandibular joint can also develop in SBO.

Diffusion along the temporal bone through the fissure of Santorini commonly involves the styloid mastoid foramen (containing the facial nerve) and the jugular foramen (containing the glossopharyngeal, vagal and accessory nerves). The infectious process, from the fissure of Santorini, spreads to retro condylid fat, parapharyngeal fat, temporomandibular joint, and masticatory muscles.

Prasad et al. divided temporal bone osteomyelitis into these 3 categories to define the spread of disease: osteomyelitis of the lateral temporal bone (bone involvement of the mastoid and middle ear cleft), medial temporal bone osteomyelitis (bone involvement of the otic capsule, petrous apex, and clivus) and
pan temporal osteomyelitis (showing features of both mastoid/ middle ear cleft and petrous involvement).²,¹³

Lesser et al. related the onset of disease with malignant external otitis, thus describing 3 different types of skull base osteomyelitis. In the first case, the necrotizing external otitis, detected at the clinical and diagnostic examination, extends to the skull base. The development of osteomyelitis represents another possibility after the resolution of the necrotizing external otitis. Finally, osteomyelitis can occur primary without any apparent procedural lateral infection.¹⁴

Moreover, Kwon et al. described 4 spreading patterns of the soft tissue extension: medial, anterior, crossed and intracranial spreading. In addition to the patterns mentioned above, there can be intravascular involvement.¹²

The diagnostic approach of osteomyelitis at the base of the skull can be a challenge, considering that the diffusion model can be non-specific for an infectious process.

Despite the characteristic pain symptoms and the involvement of the cranial nerves, the symptoms and signs of a severe infection can be almost absent.¹⁵

For example, the absence of infectious symptoms such as fever, pain, swelling, and increased ESR and white blood cell counts may be absent in immunocompromised patients, with possible diagnostic error.

Moreover, the involvement of the cranial nerves of the advanced SBO can be associated with difficulties with swallowing and paralysis of the facial nerve, pathological signs commonly associated with a neoplastic process of the base of the skull.¹⁶

Although the osteomyelitis of the cranial base presents local edema of the tissue and extensive diffuse bone destruction at radio diagnostic examinations, imaging can sometimes be highly suggestive but not specific to the pathology as mentioned earlier.¹⁷

Even after the conclusion of follow-up, patients who have developed severe complications, such as cranial nerve involvement, do not have complete recovery of the primary function.

Differential diagnosis of the disease should consider Squamous cell carcinoma of the head and neck, which may also involve soft tissues of the calvarium and preclival and destruction of the base of the skull. Other differential diagnoses are nasopharyngeal carcinoma, multiple myeloma, lymphoma and metastatic processes.¹⁸⁻²¹

Among the imaging methods available to date, computed tomography (CT) is the most commonly used for the initial approach to the disease in question. It allows to quickly and relatively easily evaluate both the involvement of soft tissues and the degree of demineralization of the mastoid bone.¹⁴

Through the use of ultra-thin high-resolution Tc, the evaluation of bone erosion is optimal, but this is not necessarily present, especially in the initial stages of the pathology (Figure 1).

The strength of this modality is the evaluation of bone erosion and demineralization, in particular with the use of high-resolution (ultra) thin CT on several levels.

Since the infectious outbreak originates from an initial process of the external ear, the CT scan can show either the classic finding of edema of the external auditory canal with the inflammation of the surrounding adipose tissue or very slight differences of soft tissues only appreciable comparing them with the contralateral ear.²²

Indeed, a frequent consequence is the anterior development of the pathology up to the temporal mandibular articulation
with the presence of associated destructive phenomena. However, bone involvement is often a late phenomenon, common also to other inflammatory or neoplastic diseases.\textsuperscript{23}

The diagnostic method must also study the extension to the skull of the base of the foramina, in particular, the jugular and stylomastoid foramen, frequently involved.

In this regard, through the analysis of the sub-temporal fat, and parapharyngeal fat plans, it allows us to evaluate the median diffusion of the nasopharynx through the Eustachian tube. However, these plans are more difficult to study than the masticatory or retromandibular one.

Magnetic resonance imaging is the superior imaging technique for anatomical evaluation of soft tissue components compared to CT, considering, for example, signs such as thrombosis and intracranial spread (Figure 2).\textsuperscript{17}

The T1-w sequences allow identifying fat planes adjacent to the base of the skull, the sub temporal region, the space of the mastoid. Even the muscles affected by the inflammatory process demonstrate an altered signal in T1-w, such as arterial and venous vascular involvement. Although it is not possible to detect alterations of the cortical bone, indirect signs are represented by an abnormal signal of the bone marrow (Figure 5). In

\textbf{Figure 4.} Ga 67 scan: Long-term follow-up with Ga 67 (Gallium Citrate) scan highlights the marked accumulation in the temporal region and extends to the skull base (indicated by the arrows).
the T2-w sequences, the infectious focus previously detected at T1-w will appear hypointense due to the particular condition of the diabetic patient’s microcirculation. Surely the T1-w sequences performed after administration of Gadolinium contrast medium associated with fat suppression allow a better interpretation of the soft tissues and the extension to the skull base as well as to identify the severe intracranial complications. Unfortunately, MRI is not indicated in the long-term evaluation of the disease because the method does not discriminate against inflammatory sequelae from an active infectious process.

Finally, through the analysis of the diffusion coefficient to the DWI, it is possible to differentiate the infiltrative neoplastic lesions from the indicated pathology.

Nuclear medicine methods allow a functional analysis of the images obtained, gaining importance both in the initial diagnosis and in the follow-up of the disease. The Tc-99 m-MDP test is positive at the beginning of the osteomyelitis process, being related to the increased osteoblastic activity (Figure 3). However, the same method will be used in other conditions characterized by high bone turnover as in malignant diseases. Furthermore, they follow the corresponding clinical improvement, following the corresponding clinical improvement. The principle that allows the detection of infectious outbreaks is the increase in vascular flow due to the infectious process; however, the same specific accumulation of labeled leukocytes can also occur in healthy marrow tissue.

On the other hand, gallium-67-citrate, by binding to actively dividing cells, such as leukocytes involved in the infectious process, allows its use in monitoring treatment response (Figure 4). This method has the disadvantage of being particularly expensive and subjecting the patient to a high dose of radiation.

Finally, the problem of the low spatial resolution of both the described exams is overcome by the SPECT-CT, a technique that thanks to 3D reconstructions improve the anatomical correlation.

Positron Emission Tomography (PET) is an alternative method, whose use in the literature is described in the follow-up of the pathology. The most commonly used tracer is the FDG, a marker that detects the conditions of increased cellular metabolism and precisely for this nonspecific for infectious diseases. This method also has the spatial limitation of the first 2 nuclear ones. Precisely for this reason, new technologies have been developed and are widely used in the diagnosis and control of malignant external otitis, such as PET-Tc and PET-MRI.

**Conclusion**

The osteomyelitis of the base of the skull is a severe complication of malignant external otitis, characterized by a symptomatic pattern that is often but not always clear and easily diagnosed. Therefore, considering the often shaded and subtle interpretation clinic, the correct interpretation of radiological examinations is essential, identifying critical points in the evaluation of the pathology.

The major complication of MOE is the involvement of the skull base, with subsequent paralysis of the cranial nerves and hearing loss. Recurrences of MOE can occur up to 1 year after cessation of therapy, so a patient must be considered cured until that period has elapsed. Furthermore, the pathology is insidious due to the lack of correlation between the severity of this and the associated clinical-symptomatologic picture, making imaging the only way to avoid severe complications. Numerous new high-resolution and highly useful methods are described in the literature, but the relatively low diffusion of these technologies in primary and first-level hospitals must be considered.

**Figure 5.** MRI scan T1w / T2: wide diffusion of the infectious process in the soft tissues starting from CUE. The region of the temporomandibular joint appears to be infiltrated.
Precisely for this reason, the right radiological tips must be better interpreted and recognized to understand the pathology fully. The corresponding clinic must always dictate the use of various radiological methods and the correct timing of the various procedures is necessary.

Authors' Note
All procedures performed in this study involving human participants were in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards.

Declaration of Conflicting Interests
The author(s) declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

Funding
The author(s) received no financial support for the research, authorship, and/or publication of this article.

ORCID iD
Salvatore Ferlito https://orcid.org/0000-0001-7136-9698
Lorenzo Mannelli https://orcid.org/0000-0002-9102-4176
Giacomo Spinato https://orcid.org/0000-0001-5295-5077

References
1. Meltzer PE, Kelemen G. Pyocyanous osteomyelitis of the temporal bone, mandible and zygoma. Laryngoscope. 1959;69(10):1300-1316. doi:10.1288/00005537-195910000-00006
2. Prasad KC, Prasad SC, Mouli N, Agarwal S. Osteomyelitis in the head and neck. Acta Oto-Laryngologica. 2007;127(2):194-205. doi:10.1080/00016480600818054
3. Chandler JR, Grobman L, Quencer R, Serafini A. Osteomyelitis of the base of the skull. Laryngoscope. 1986;96(3):245-251.
4. Adams A, Offiah C. Central skull base osteomyelitis as a complication of necrotizing otitis externa: imaging findings, complications, and challenges of diagnosis. Clin Radiol. 2012;67(10):e7-e16. doi:10.1016/j.crad.2012.02.004
5. Chandler JR. Malignant external otitis. Laryngoscope. 1968;78(8):1257-1294. doi:10.1288/00005537-196808000-00002
6. Hariga I, Mardassi A, Bellaj Younes F, et al. Necrotizing otitis externa: 19 cases’ report. Eur Arch Otorhinolaryngol. 2010;267(8):1193-1198. doi:10.1007/s00405-009-1194-2
7. Corey JP, Levandowski RA, Panwalker AP. Prognostic implications of therapy for necrotizing external otitis. Am J Otol. 1985;6(4):353-358
8. Babiatzki A, Sade J. Malignant external otitis. J Laryngol Otol. 1987;101(3):205-210
9. Cohen D, Friedman P. The diagnostic criteria of malignant external otitis. J Laryngol Otol. 1987;101(3):216-221
10. Levenson MJ, Parisier SC, Dolitsky J, Bindra G. Ciprofloxacin: drug of choice in the treatment of malignant external otitis (MEO). Laryngoscope. 1991;101(8):821-824. doi:10.1288/00005537-199108000-00004
11. Sreepada GS, Kwartler JA. Skull base osteomyelitis secondary to malignant otitis externa. Curr Opin Otolaryngol Head Neck Surg. 2003;11(5):316-323
12. Kwon BJ, Han MH, Oh SH, Song JJ, Chang KH. MRI findings and spreading patterns of necrotizing external otitis: is a poor outcome predictable? Clin Radiol. 2006;61(6):495-504. doi:10.1016/j.crad.2006.01.010
13. Prasad SC, Prasad KC, Kumar A, Thada ND, Rao P, Chalasani S. Osteomyelitis of the temporal bone: terminology, diagnosis, and management. J Neurol Surg P B, Skull Base. 2014;75(5):324-331. doi:10.1055/s-0034-1372468
14. Lesser FD, Derbyshire SG, Lewis-Jones H. Can computed tomography and magnetic resonance imaging differentiate between malignant pathology and osteomyelitis in the central skull base? J Laryngol Otol. 2015;129 (9):852-859. doi:10.1017/s0022215115001991
15. Sharma P, Agarwal KK, Kumar S, et al. Utility of (99m)Tc-MDP hybrid SPECT-CT for diagnosis of skull base osteomyelitis: comparison with planar bone scintigraphy, SPECT, and CT. Jpn J Radiol. 2013;31(2):81-88. doi:10.1007/s11604-012-0148-6
16. Ducic Y. Management of osteomyelitis of the anterior skull base and craniovertebral junction. Otolaryngol Head Neck Surg. 2003;128(1):39-42. doi:10.1067/mhn.2003.9
17. van Kroonenburgh AMJL, van der Meer WL, Bothof RJP, van Tilburg M, van Tongeren J, Postma AA. Advanced imaging techniques in skull base osteomyelitis due to malignant otitis externa. Curr Radiol Rep. 2018;6(1):3-3. doi:10.1007/s40134-018-0263-y
18. Slattery WH III, Brackmann DE. Skull base osteomyelitis. Malignant external otitis. Otolaryng Clin N Am. 1996;29(5):795-806
19. Iannella G, Stasolla A, Pasquariello B, Re M, Magliulo G. Tymanomastoid cholesterol granuloma: radiological and intraoperative findings of blood source connection. Eur Arch Otorhinolaryngol. 2016;273(9):2395-2401. doi:10.1007/s00405-015-3820-5
20. Magliulo G, Iannella G, Chinigio Appiani M, Re M. Unilateral vestibular schwannoma associated with a Jacobson’s schwannoma. Eur Rev Med Pharmacol Sci. 2014;18(9):1422-1425.
21. Galletti B, Gazia F. Rhinocerebral mucormycosis with dissemination to pontine area in a diabetic patient: treatment and management. Clin Case Rep. 2019;7(7):1382-1387. doi:10.1002/ccr3.2255
22. Mortazavi MM, Khan MA, Quadri SA, et al. Cranial osteomyelitis: a comprehensive review of modern therapies. World Neurosurg. 2018;111:142-153. doi:10.1016/j.wneu.2017.12.066
23. Chan LL, Singh S, Jones D, Diaz EM Jr, Ginsberg LE. Imaging of mucormycosis skull base osteomyelitis. AJNR Am J Neuroradiol. 2000;21(5):828-831.