Autoimmune inner ear disease – a case report and brief review of literature

Deepa Panikkath*, Avinash Adiga, and Michael Calmes
Department of Internal Medicine, Texas Tech University Health Sciences Center, Lubbock, Texas, USA

Abstract
Autoimmune inner ear disease (AIED) is a rare syndrome characterized by progressive bilateral sensorineural hearing loss (SNHL). It is a poorly understood form of hearing loss often accompanied by vestibular symptoms and can mimic Meniere’s disease. Diagnosis is mainly based on clinical characteristics and response to immunosuppressive agents. It is one of the rare causes of hearing loss that is potentially reversible with treatment if rapidly initiated. We hereby present a case of a 54-year-old female diagnosed with Autoimmune inner ear disease and a brief review of literature.

Introduction
Autoimmune inner ear disease (AIED) is a relatively rare cause of sensorineural hearing loss probably accounting for less than 1% of hearing loss [1]. Initially described in 1979 by McCabe, it is characterized by rapid progressive hearing loss often occurring bilaterally with or without vestibular symptoms [2]. The pathophysiology of this disease still remains to be fully elucidated. Autoimmunity is thought to play a role based on the presence of autoantibodies directed against the inner ear, its association with other systemic autoimmune disorders, and treatment response to immunosuppressants. Due to lack of a definite diagnostic test, the clinical presentation and treatment response to immunosuppressants establishes its diagnosis.

Case report
A 54-year-old Caucasian female presented with complaints of sudden onset hearing difficulty and dizziness following a brief flu like illness. She had no accompanying tinnitus, headaches, blurry vision, numbness or weakness of extremities. Physical examination showed no neurological deficits but was significant for bilateral hearing loss. Her routine laboratory investigations were normal. MRI of the brain was within normal limits. Audiogram done for evaluation of hearing difficulty revealed asymmetric bilateral sensorineural deficit, right more than the left. Autoimmune workup was done which was positive for an elevated ANA with titer of 1:160 and a speckled pattern seen on immunofluorescence. She also had positive antibody against 68-kD antigen. She was diagnosed with Autoimmune inner ear disease based on her clinical presentation and positive serology for anti-68kD antibodies. She was started on a trial of oral steroids which improved her vestibular symptoms. No significant improvement in hearing was seen. Her clinical course worsened with development of tinnitus. Repeat laboratory test showed persistence of anti-68kD antibody and rising ANA titers to 1:320. She was continued on daily oral steroids and methotrexate 10 mg weekly dose was initiated. With this regime, she had dramatic improvement in her hearing deficit. Her steroids were then tapered to a low dose with addition of diuretics to her treatment regime. She continued to be in remission on this regime.

Discussion
The initial description of AIED was in 1979 by McCabe who described a series of patients with rapidly progressive bilateral sensorineural hearing loss (SNHL) [2]. The exact pathogenesis of AIED remains unclear. Autoimmunity was initially proposed by McCabe as the cause as patients showed marked improvement with steroids [2]. Inner ear is now considered to be an immunocompetent site with the endolymphatic sac and presence of resident T cells capable of mounting adequate immune responses [3,4]. A number of antibodies against various inner ear antigens have been identified in experimental and human AIED. Their exact pathogenic role is still unclear. These include antibodies against subcomponents of crude inner ear antigens like 31 kD protein and the 60 kD protein, antibodies against cochl, connexin 26, myelin protein P0 etc [5,6]. One of the most studied antigen in AIED is the inner ear 68-kd antigen. Autoantibodies against this antigen was initially detected in 1990 by Harris and Sharp in patients with idiopathic bilateral sensorineural hearing loss [7]. In 1994, Moscicki et al. confirmed these findings and demonstrated a correlation between steroid responsiveness and patient’s 68-kD antibody status [8]. This 68-kD antigen was later identified to be heat shock protein 70(hsp 70) [9]. Approximately 30% of patients with AIED also have coexisting systemic autoimmune disease like systemic lupus erythematosus, rheumatoid arthritis, disseminated vasculitis, Sjögren’s syndrome, myasthenia gravis, Hashimoto’s thyroditis etc. [10,11]. The patient reported above demonstrated antibodies against 68-kD antigen and ANA positivity in titers ranging from 1:160 to 1:320 in a speckled pattern. ANA positivity in a speckled pattern have been reported previously by Dayal et al. in patients with AIED [12].

The hallmark of AIED is the presence of rapidly progressive, often bilateral hearing loss over a period of weeks to months. Vestibular symptoms have been reported in almost 50% of patients. Due to...
the constellation of auditory and vestibular symptoms, it can mimic Meniere’s disease. The course often fluctuates with episodic flares and periods of remission [2,13]

Diagnosis is often based on clinical suspicion and improvement of symptoms with immunosuppressants due to lack of specific diagnostic tests. Western blot tests have been widely used to detect anti-68-kd antibodies in the sera. A recent meta-analysis evaluating the diagnostic accuracy of Western blot showed a pooled sensitivity of 0.70 (95% confidence interval [CI], 0.59–0.80) and specificity of 0.98 (95% CI, 0.87–1.00) [14]. Patients also frequently undergo series of tests to rule out common autoimmune diseases. Common lab tests include ESR, CRP, ANA, Rheumatoid factor, and complement levels. Workup should also include tests to rule out syphilis. Imaging studies like MRI of the brain and internal auditory canal and PET scan have failed to demonstrate utility in diagnosis though useful in ruling out other organic causes for hearing loss like acoustic neuroma, intracranial metastases, demyelinating diseases etc. In some cases, MRI may show enhancement of vestibular and cochlear elements.

Prompt and early treatment has a significant impact as AIED is one of the few potentially reversible causes of hearing loss. The most widely used and effective treatment modality in AIED is corticosteroid therapy. Patients are started on steroids without delay when AIED is suspected as disease is very steroid responsive. The initial dosage regimen is 60 mg or 1 mg/kg per day of prednisone or 6-methylprednisolone for a period of 4 weeks [15]. Low sodium diet and diuretics are also beneficial in the management [16]. In most steroid responsive patients, the dose can be lowered or tapered off without relapse. Some patients can have symptoms refractory to steroids or have side effects that limit the use of steroids, in which case immune modulating therapies are indicated. Methotrexate is a commonly used immunosuppressant in AIED and Sismannis et al. first reported the usefulness of methotrexate in the improvement of symptoms of aural fullness, tinnitus vertigo and hearing [17]. Though shown to be useful in controlling the vestibular symptoms, methotrexate was found to be less effective in maintaining hearing improvement [18]. Other immunosuppressants used includes biologic agents like Infliximab, Etanercept, and Rituximab [19,20]. Intratympanic instillation of medications like steroids has shown promising results [21].There are ongoing clinical trials for drug eluting gels that could be injected into the middle ear space to give sustained drug release. Cochlear implants have also been tried in patients with progressive AIED and they have demonstrated good functional outcomes making it a viable therapeutic option in patients wanting to avoid long term immunosuppression [22].

Conclusion

Autoimmune inner ear disease (AIED) is a rare distinct cause of idiopathic sensorineural hearing loss. Patients often develop auditory and vestibular symptoms with a rapid progressive fluctuating course. Characteristics of this disease includes responsiveness to steroids and other immunosuppressive agents. There is no gold standard test to aid in the diagnosis of the disease. Prompt and early treatment with immunosuppressants is essential in preventing irreversible hearing loss. The natural history and pathogenesis of the disease still remains to be fully elucidated and further research is needed for a better understanding of this rare disease entity.

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