MEFV, IRF8, ADA, PEPD, and NBAS gene variants and elevated serum cytokines in a patient with unilateral sporadic Meniere’s disease and vascular congestion over the endolymphatic sac

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ABSTRACT

The etiology and underlying mechanism of Meniere’s disease (MD) development are still unknown, although inflammation and autoimmunity have been implicated as underlying mechanisms. The human endolymphatic sac (ES) has been reported to have innate and adaptive immune capacity in local immune reactions. In vivo demonstration of inflammation of the ES in patients with MD is missing in the literature. We report the case of a 47-year-old female patient diagnosed with unilateral MD with genetic variants and cytokine markers indicating inflammation and vascular congestion of the ES. Endolymphatic hydrops in the right cochlea (grade 2) and vestibulum (grade 3) were detected using MRI. She carried heterozygous variants in MEFV (c.442G > C), IRF8 (c.1157G > T), ADA (c.445C > T), PEPD (c.151G > A), NBAS (c.4049T > C), CSF2RB (c.2222C > T), HPS6 (c.277G > T), IL2RB (c.1109C > T), IL12RB1 (c.1384G > T), IL17RC (c.260_271del GCAAGAGCTGGG), LIG1 (c.746G > A), RAG1 (c.650C > A), and SLX4 (c.1258G > C). In the serum, the levels of granulocyte colony-stimulating factor (G-CSF), macrophage inflammatory protein 1α, and IL7 were significantly elevated, and the level of IL2Rα was reduced.

1. Introduction

The etiology and underlying mechanism of Meniere’s disease (MD) development are still unknown, although clinical subgroups have been proposed (Frejo et al., 2016, 2017). However, inflammation resulting from various causes might be a common underlying mechanism of MD because intratympanic dexamethasone induces acceptable vertigo control in 91% of cases (Boleas-Aguirre et al., 2008). Macrophage-like cells and lymphocytes have been detected in the endolymphatic sac (ES) of patients with MD (Yazawa et al., 1989). Both innate and adaptive immune capacity of the human ES in local immune reactions have been indicated by gene expression analyzed using a combination of microarray analysis and immunohistochemistry (Moller et al., 2015). The existence of fully functional macrophages involved in innate and adaptive immune reactions in the human ES was further suggested by a high-resolution immunohistochemistry study (Kampfe Nordstrom et al., 2018). However, in vivo demonstration of inflammation of the ES in patients with MD is still missing in the literature. We report the case of a patient with unilateral sporadic MD with vascular congestion over the ES carrying multiple gene variations and elevated serum cytokines.

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2. Case presentation

2.1. Clinical data

A 47-year-old woman complained of episodic vertigo and fluctuating hearing loss, tinnitus, and aural fullness in the right ear with a 6-year duration. Vertigo lasted for a maximum of 5 h and occurred once a year, but was exacerbated over the past 3 years. Five years ago, slight tinnitus appeared in the left ear but was not associated with the vertigo attack. Fluctuating hearing loss in the right ear without vertigo recurred 30 times over the past year. She received intravenous infusions of alprostadil, vinpocetine, and dexamethasone, as well as hyperbaric oxygen therapy. Later, 5 intratympanic injections of dexamethasone were administered. Since the symptoms were not controlled, she visited author J.Z. and was diagnosed as having definite MD with MRI evidence of endolymphatic hydrops in both the cochlea and vestibule of the right ear (Fig. 1). She did not complain of symptoms of any other diseases and denied a familial history of MD. She received an intratympanic sustained delivery of dexamethasone on June 24, 2020, which was repeated 4 times. Her hearing recovered to normal (her own feeling) after the 3rd delivery on August 13, 2020, but fluctuated.

Fig. 1. Inner ear MRI 24 h after intratympanic administration of Gd-DTPA. T2-sampling perfection with application-optimized contrasts using a flip angle evolution (SPACE) sequence did not indicate either inner ear fibrosis or vestibular schwannoma (A, B). Cochlear grade 2 (C) and vestibular grade 3 (E) endolymphatic hydrops (EH) in the right ear (R) was demonstrated by heavily T2-weighted 3-dimensional fluid-attenuated inversion recovery reconstructed using a magnitude plus zero-filled interpolation (hT2FLAIR-MZFI) sequence acquired 24 h after intratympanic administration of Gd-DTPA mixed with dexamethasone (Zou J. et al., 2022). There was no EH in the left ear (L in D and F). CN: cochlear nerve; FN: facial nerve; LSCC: lateral semicircular canal; SM: scala media; ST: scala tympani; SV: scala vestibuli; VA: vestibular aqueduct; VN: vestibular nerve; Sa: saccule; Ut: utricle. Scale bars = 3.0 mm.
and progressed even after two further sustained deliveries of dexamethasone were repeated. Laboratory examinations showed that the levels of the complement proteins C3 (0.73 g/L) and C4 (0.13 g/L) were below the normal range. Routine blood examination showed elevated monocytes (11.2%) and basophils (1.1%). A routine urine test showed positive white blood cells (25/μL) and epithelial cells (15.4/HP). Blood biochemical examination showed reduced total proteins (44 g/L) and aspartate aminotransferase (13 U/L) and elevated high-density lipoprotein (1.59 mmol/L) and lipoprotein-α (393 mg/L). The other results, including those for immunoglobulins, transferrin, β2-microglobulin, α1-microglobulin, ceruloplasmin, anti-streptolysin “O” antibody, erythrocyte sedimentation rate, rheumatoid factor, C-reactive protein, total triiodothyronine (T3), 3,3',5-triiodo-L-thyronine (3,3',5-T3), tetraiodothyronine (T4), calcitonin, thyroid stimulating hormone, thyroglobulin, anti-thyroglobulin antibodies, anti-thyroid peroxidase antibody, anti-
thryotropin receptor antibody, and coagulation function, were in the normal range. Immunological detections of hepatitis B and C virus were negative. She received endolymphatic sac delivery of steroids on May 12, 2021, as previously reported (Zou, 2019). Significant vascular congestion was visualized surrounding the ES during surgery (Fig. 2). A follow-up infusion of dexamethasone (5 mg/mL, 1 ml) to the area of the endolymphatic sac was performed through a preplaced catheter on July 13, 2021. From November 1 to December 16, 2021, the patient had daily mild vertigo (nonrotational) and one episode of rotational vertigo. Her hearing loss remained fluctuating and progressive (Fig. 3). The aural fullness went away on December 28, 2021.

### 2.2. Gene sequencing

Genomic DNA was extracted from the peripheral blood sample, and the whole exomes of 423 candidate genes potentially involved in diseases associated with immune reactions were selected according to Human Phenotype Ontology (http://human-phenotype-ontology.github.io/) and sequenced with next-generation sequencing (Zou et al., 2016). The data were analyzed, and the clinical significance of allelic variants was judged as previously reported (Zou, 2019). The following heterozygous variants were detected (Table 1): an autosomal-dominant inherited variant in Mediterranean fever (MEFV) (c.442G > C), an autosomal-dominant/recessive inherited variant in interferon regulatory factor 8 (IRF8) (c.1157G > T), autosomal-recessive inherited but pathologic variants in adenosine deaminase (ADA) (c.445G > T) and peptidase D (PEPD) (c.151G > A), and an autosomal-recessive inherited but limited pathologic variant in neuroblastoma-amplified sequence (NBAS) (c.4049T > C). The abovementioned variants in the patient were verified using Sanger sequencing (Fig. 4). Autosomal-recessive inherited variants were also detected in the following genes: colony stimulating factor-2 receptor β (CSF2RB) (c.2222C > T), Hermansky-Pudlak syndrome 6 (HPS6) (c.277G > T), interleukin-2 receptor β (IL2RB) (c.1109C > T), IL12 receptor β1 (IL12RB1) (c.650G > C), IL7 receptor (IL7R) (c.1384G > T), and IL17 receptor A (IL17RA) (c.746G > A), recombination activating gene 1 (RAG1) (c.650C > A), and SLX4 (c.1258G > C, c.5072A > G).

### 2.3. Quantification of cytokines/chemokines

Two milliliters of blood was taken from each volunteer at 7:00 a.m. before breakfast, and the serum was stored at −80 °C until measurements. Forty-eight human cytokines/chemokines were quantified using a Bio-Plex Pro Human Cytokine Screening Panel (Bio-Plex Suspension Array System; Bio-Rad Laboratories Inc., Hercules, CA, USA) according to the manufacturer’s instructions (data will be reported separately). Compared to the values of the 39 healthy control volunteers, the levels of granulocyte colony-stimulating factor (G-CSF), macrophage inflammatory protein 1α, and IL7 in the serum of the patient were significantly elevated (exceeding the mean ± 2 SD). The level of IL2Rα was below the range of the standard curve (SI-Table 1).

### 3. Discussion

According to a previous report (Frejo et al., 2017), the patient had unilateral MD type 1. It was reported that patients with Cogan’s syndrome present audi vestibular symptoms similar to those of patients with Meniere’s syndrome (Greco et al., 2013). However, the present patient did not have the ocular symptoms or systemic disease with a pathological mechanism of vasculitis that is associated with Cogan’s syndrome. Although a histological study was unavailable, we demonstrated severe local vascular congestion, indicating vasculitis, over the ES in this patient with unilateral sporadic MD. Genetic analysis of the patient demonstrated variants in MEFV, IRF8, ADA, PEPD, and NBAS, among others.

Mutations in MEFV, the gene encoding pyrin (NLRP3), have been defined as variants mostly associated with monogenic systemic autoinflammatory diseases (Szekanecz et al., 2021). The c.442G > C and c.329T > C mutations in MEFV appeared in a Han Chinese patient with adult-onset Still’s disease, which has been advocated as being in the spectrum of autoinflammatory diseases (Ou-Yang et al., 2018). IRF8 is expressed in myeloid cells (macrophages, monocytes, dendritic cells) and forms part of a group of core genes in which genetic variants constitute risk factors for a large spectrum of chronic inflammatory diseases in humans (Salem et al., 2020). However, the influence of the c.1157G > T mutation on the protein function of IRF8 is unknown.

Adenosine deaminase (ADA) is a ubiquitous enzyme important for the degradation and salvage of purine metabolites, and the
inherited absence of ADA leads to the accumulation of these metabolites that are toxic to rapidly proliferating cells such as lymphocytes, resulting in severe combined immunodeficiency (SCID). In addition to increased susceptibility to infections, ADA deficiency has also been reported to be associated with sensorineural loss (Albuquerque et al., 2004; Tanaka et al., 1996; Xu et al., 2019). The impact of the c.445C>T mutation on the protein function of ADA is harmful. PEPD encodes prolidase, which catalyzes the hydrolysis of a dipeptide containing a C-terminal proline or hydroxyproline into constituent amino acids, and the most studied substrates of the enzyme are imidodipeptides, which are generated during the breakdown of collagen. It was reported that oxidative stress stimulates collagen breakdown via inductions of matrix metalloproteinases and induces inflammatory homeostasis (Eni-Aganga
et al., 2021). The c.151G > A mutation is harmful for the function of prolidase. It is possible that a disturbance of the function of prolidase promotes the inflammation induced by oxidative damage due to decreased clearance of collagen catabolites.

NBAS functions as a component of an endoplasmic reticulum (ER) tethering complex involved in retrograde Golgi-ER transport and the formation of large transport vesicles for bulky cargo (e.g., collagen) at the ER exit site in the secretory pathway. There are three clinical subgroups associated with the affected region of the NBAS protein: the combined phenotype (associated with the β-propeller); infantile liver failure syndrome type 2 (ILFS2; associated with Sec39); and short stature, optic atrophy, and Pelger-Huët anomaly (SOH; associated with the C-terminal) (Staufner et al., 2020). Although there is limited impact on the function of NBAS caused by the c.4049T > C mutation, it is possible that a disruption in Golgi-ER transport and the formation of large transport vesicles at the ER exit site induced ER stress and inflammation in the current patient.

The impacts of subsequent autosomal-recessive inherited variants on protein function are unknown. CSF2RB encodes the β-chain of granulocyte-macrophage colony-stimulating factor receptor (GM-CSFR), which forms a high-affinity receptor in the presence of the α-chain upon interacting with GM-CSF, and the activation promotes the survival, proliferation, maturation, and functional activation of a spectrum of hemopoietic lineages, including committed myeloid progenitors, neutrophils, monocytes, and eosinophils (Lock et al., 1994). The HPS6 gene is named Hermansky-Pudlak syndrome 6 and encodes the HPS6 protein, which is a subunit of the BLOC-2 protein complex that directly interacts with the dynactin p150Glued subunit of the dynein–dynactin motor complex and acts as a cargo adaptor for the retrograde motor to mediate the transport of lysosomes from the cell periphery to the perinuclear region (Li et al., 2014). Variants in the HPS6 gene were also reported in Chinese patients with Hermansky-Pudlak syndrome (Wang et al., 2021). IL2RB encodes the β subunit of the IL-2 receptor, which forms a complex with the γ subunits upon IL-2 and IL-15 stimulation and plays a central role in the control of the immune system. Patients with homozygous mutations in the IL2RB gene were shown to present with autoantibodies, hypergammaglobulinemia, and bowel inflammation, among other symptoms and signs (Zhang et al., 2019). Human IL12RB1 encodes IL12Rβ1, which is an essential component of the IL12-and IL23-signaling complexes and promotes delayed type hypersensitivity and autoimmunity, as well as the immunological reaction that limits, for example, tuberculosis that might be accompanied by leukocytoclastic vasculitis (Robinson, 2015). IL17RC encodes the IL-17 receptor C (IL-17R C) subunit that forms the IL-17R complex together with the IL-17RA subunit and mediates inflammatory activities after binding to IL-17. The IL-17 signaling axis has been implicated in the development of rheumatoid arthritis and the impairment of IL-17 signaling in the disease progression of psoriasis (Ho et al., 2010). LIG1 encodes DNA ligase 1, which is the primary replicative ligase and interacts with proliferating cell nuclear antigen (PCNA) via an N-terminal PCNA-interacting protein box, and mutations in this gene have been implicated in several immunological diseases (Jurkiw et al., 2021). RAG1 encodes the RAG1 protein, which modulates V(D)J recombination, which assembles and diversifies immunoglobulin and T-cell receptor genes in developing B and T lymphocytes (Brecht et al., 2020). SLX4 encodes the protein SLX4, which is involved in DNA repair and the maintenance of genome stability (Young et al., 2021).

Regarding the immunological phenotype, the levels of G-CSF, MIP1α, and IL7 were significantly elevated, while the level of IL2Rα was decreased in the serum. Elevated levels of G-CSF and MIP1α are associated with active inflammatory cellular infiltration. G-CSF stimulates neutrophil production and release from the bone marrow and promotes injury at sites of inflammation. It was reported that G-CSF is also involved in endotoxemia-enhanced blood-labyrinth barrier trafficking by cooperating with TLR4 (Urdang et al., 2020). MIP1α is a C–C chemokine that recruits B cells, cytotoxic T cells, and CD4+ T cells to the inflammatory site (Schall et al., 1993). IL-7 is mainly produced by epithelial and stromal cells and regulates T-cell homeostasis, proliferation, and survival. Beneficial therapeutic results were reported in T-cell-mediated chronic inflammatory diseases by targeting the expansion of pathogenic memory immune cells with anti-IL-7Rα mAbs (Belarif et al., 2018). The IL-2/IL-2Rα signaling pathway is essential for the regulation of both the size and the content of the peripheral lymphoid compartment, probably by influencing the balance between clonal expansion and cell death following lymphocyte activation. IL-2Rα-deficient mice develop autoimmune disorders, such as hemolytic anemia and inflammatory bowel disease (Willerford et al., 1995). Elevated soluble IL-2Rα concentrations in adults indicate the activation of lymphocytes during infection and inflammation and with autoimmune disease (Downes et al., 2014). However, we cannot conclude the clinical relevance of the reduced soluble IL-2Rα level in the current patient. Although autoantibody spectrum test results were negative, we cannot rule out the existence of autoantibodies against antigens of the inner ear or endolymphatic sacs.

C3 and C4 are important opsonins that bind to and facilitate the clearance of immune complexes and apoptotic debris, which would otherwise promote inflammation and autoimmune diseases. Angiogenesis was found to be increased in C3−/− mice, as demonstrated by an increased capillary/muscle fiber ratio and increased proliferating endothelial cells (Gotz et al., 2021). Complete C4 deficiency was reported in SLE patients who finally developed fatal central nervous system vasculitis (Rupert et al., 2002).

The current unilateral sporadic MD patient demonstrated genetic variants and varied serum cytokine levels as well as vascular congestion over the ES, indicating vasculitis. Although malfunction of the abovementioned genes is involved in autoinflammation and autoimmunity, the clinical relevance of the detected coexistent genotypes in MD phenotypes needs further investigation.

Declaration of competing interest

The authors declare no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.joto.2022.03.001.

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