

# Cochlear Implant Impedance Fluctuation in Ménière's Disease: A Case Study

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**Objective:** To contribute to the understanding of hearing fluctuation in Ménière's disease (MD) by disseminating a case study of a cochlear implanted ear with ongoing fluctuation of electrode impedances with episodic tinnitus and no associated vestibular symptoms.

**Study Design:** Retrospective case review.

**Setting:** Tertiary referral audiology clinic.

**Patient:** Man, born in 1936, with a total hearing loss in the right ear because of Mumps at age 8 years and a fluctuating progressive hearing loss in the left ear because of Ménière's disease since age 63 years.

**Intervention:** Sequential bilateral cochlear implantation right ear in August 2002 and left ear in March 2006.

**Main Outcome Measure:** Impedance measurements of implanted intracochlear electrodes via common ground stimulation using proprietary programming software.

**Results:** Electrode impedances in the MD showed significant ongoing variation since implantation, whereas the contralateral non-MD ear remained stable over a period of 9 years.

**Conclusion:** Electrode impedances in the ear with MD showed a variation pattern similar to that found in the hearing fluctuation characteristic of the disease. These findings raise the possibility that the same physiological mechanisms of hearing fluctuation may be responsible for intracochlear electrode impedance changes. We hypothesize that impedance fluctuation is because of changes in the permeability of the blood-labyrinth barrier because of cyclic immune activity in the inner ear which alters the electrical resistance between scala tympani and blood. **Key Words:** Cochlear implant—Electrode impedance—Hearing fluctuation—Impedance fluctuation—Ménière's disease—Tinnitus.

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Hearing fluctuation is a characteristic of Ménière's disease (MD) well documented in the literature (1). Despite hearing fluctuation, studies have shown that hearing aids are beneficial to alleviate hearing loss (2,3) but as the disease progresses they become less effective and many patients become eligible for cochlear implant (CI). Studies have shown that CI can successfully rehabilitate hearing in patients with MD who meet the audiological criteria for candidacy (4–7).

Interestingly, there have been reports that some patients continue to experience hearing fluctuation after receiving a CI in the MD ear (4–6). Graham and Dickins (8) were the first authors to report fluctuation of electrical thresholds in ears with MD and CI. Lustig et al. (4) also reported hearing fluctuation in patients with MD post-CI, noting that some patients experienced alterations in implant performance in association with fluctuations in vestibular symptoms over a follow-up period of 1 to 5 years. Two other studies recently reported hearing fluctuation in patients with MD post-CI, with incidence ranging

from 33 to 55% depending on sample size ( $N = 13, 8$ ) and follow-up period (3.5–2 yr) (5,6). CI remapping was sufficient to restore speech perception and hearing (5,6). Neuburger et al. (9) reported 16 patients with 18 affected ears in whom impedance increases were clearly demonstrated without any sign of previous inflammation.

Endolymphatic hydrops is widely accepted as the underlying cause and physiology of hearing fluctuation remains poorly understood. A clear explanation of the mechanisms of these fluctuations would be invaluable to our understanding of MD.

## This Study

This article reviews a unique case study of a patient with bilateral cochlear implants who presented significant electric hearing fluctuation in the ear affected with MD comparing to the contralateral non-MD ear.

## METHODS

Patient consent was sought and granted before writing this case study.

The patient was implanted with a Cochlear Nucleus CI24CS in August 2002 in the right ear after 50 years of total hearing loss attributed to mumps. The left ear was diagnosed with MD in 2001 and was implanted with a Cochlear Nucleus CI24RECA

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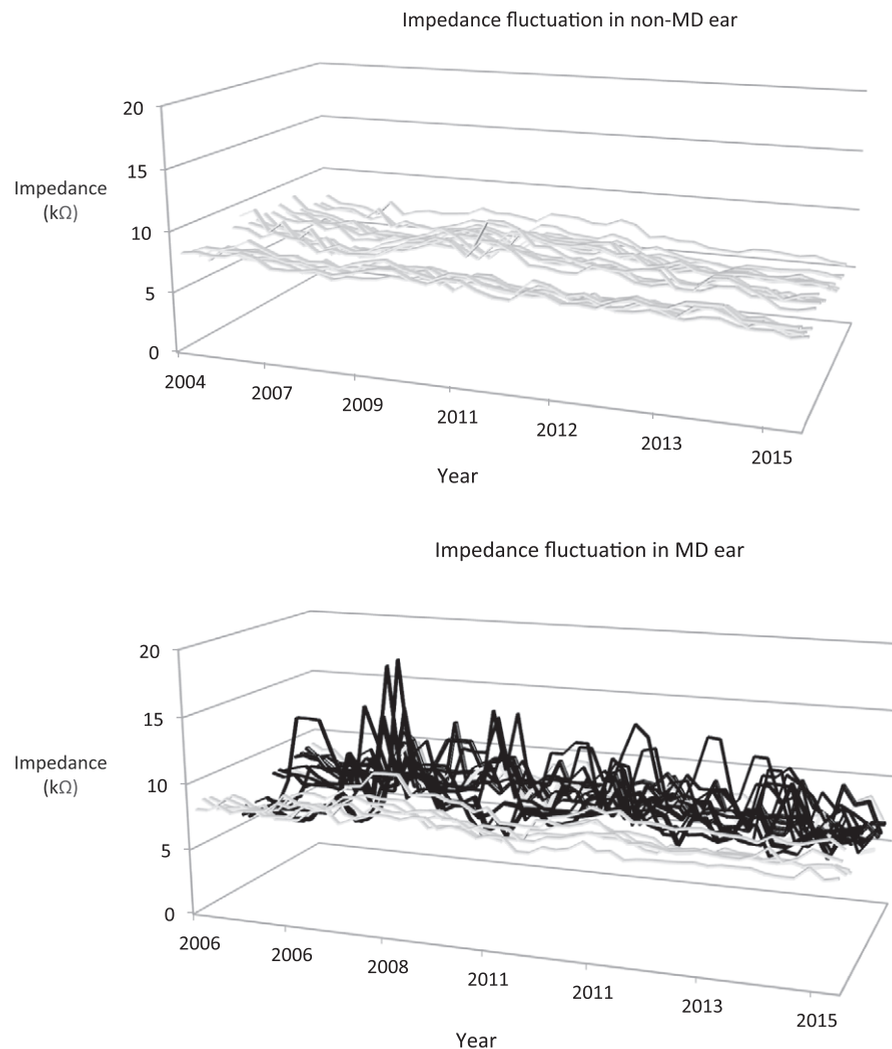
in March 2006 after fluctuating hearing loss progressed to severe levels. Both cochlear implants were the “contour” design of different generations. The CI24CS was the first generation and the soft tip of the CI24RECA is the only difference between the two electrodes. Both implants have 22 tonal topic intracochlear electrodes, with electrode 1 being the most basal (high frequencies) and 22 the most apical (low frequencies).

At the time of this review the patient had 37 visits to the clinic after receiving the second implant in 2006. During each visit the patient’s experience was noted including subjective report of hearing, speech intelligibility, hyperacusis, tinnitus, and vestibular symptoms; electrode impedances were measured in each ear; and implant was remapped if required. The authors found that simple remapping of the implant was able to address impedance fluctuation and perceived distortion in speech per-

ception and hearing through the implant. Mapping was conducted by subjectively measuring threshold (T) and comfortable (C) levels and balancing each electrode for equal loudness perception. Since the initial maps, electrodes 1 and 2 were disabled in right ear and electrode 1 in left ear because of nonauditory percept.

Thirty-eight measurements were recorded from 20 active intracochlear electrodes from the right CI (non-MD) using Custom Sound Cochlear Ltd proprietor software as shown in Figure 1A and 50 measurements from 21 active electrodes from the left CI (MD) as shown in Figure 1B. It should be noted that the left CI had more than one measurement within one session. These results were exported into an Excel spreadsheet for analysis.

For the purposes of our analysis, common ground mode impedance measurements were used and results obtained before



**FIG. 1.** A, Impedance measurements in kΩ of the 20 active intracochlear electrodes in the non-MD ear over time. Each line represents one electrode. At the front of the graph is electrode 3, which is the most basal and at the back is electrode 22, which is the most apical. B, Impedance measurements in kΩ of the 21 active intracochlear electrodes in the MD ear over time. Each line represents one electrode. At the front of the graph is electrode 2, which is the most basal and at the back is electrode 22 which is the most apical. Black lines represent electrode impedance fluctuation >3 kΩ between measures; gray lines represent fluctuation <3 kΩ between measures. MD indicates Ménière’s disease.

**TABLE 1.** Mann–Whitney test results

Variability	Right CI Non-MD	Left CI MD	Z Value	Significance (2-Tailed)
Standard Deviation	0.65	1.21	−2.95	0.003*
Variance	0.46	1.75	−2.91	0.004*

MD indicates Ménière’s disease.  
\*Statistically significant ( $P < 0.01$ ).

6 weeks post-switch-on were excluded, as unstable impedances are expected during this initial period. Impedances of electrodes 1 and 2 of the right ear and electrode 1 of the left ear were also excluded from analysis as they were not enabled in the maps.

**Statistical Analysis**

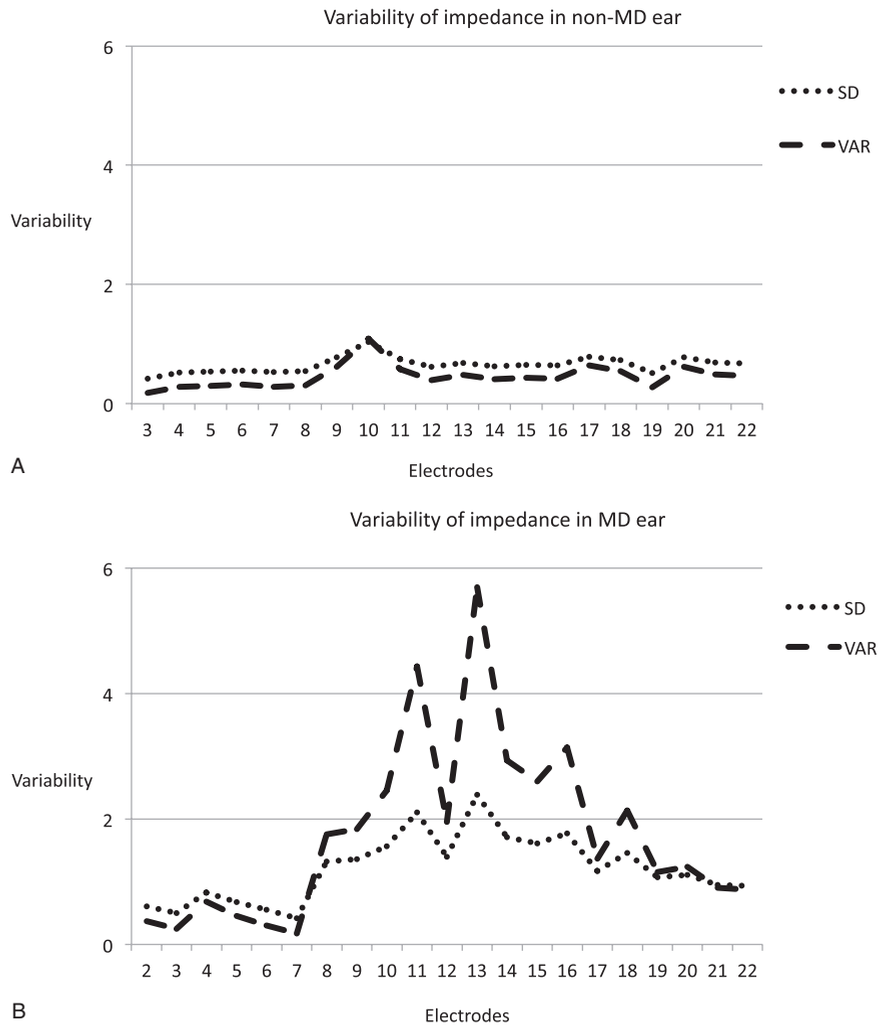
Descriptive statistics were obtained using Microsoft Excel. Figures and graphs were created to explore the variability between ears as well as within and across electrodes over time.

Mann–Whitney test was performed to quantify the variability (standard deviation, variance) between ears using SPSS.

**RESULTS**

The pattern of impedance fluctuation over time was clearly different between the two ears. Figure 1A depicts the results of the non-MD ear, which shows that all electrodes fluctuated less than 3 kΩ between measurements. Figure 1B depicts the results for the MD ear showing electrodes 9, 10, 11, 12, 13, 14, 15, 16, 18, 19, 20, and 22 which fluctuated more than 3 kΩ between measurements (represented by black lines) and the remaining electrodes which fluctuated less than 3 kΩ (represented by gray lines).

Mann–Whitney test confirmed a statistically significant difference in the variability of impedance measurements between ears (Table 1).



**FIG. 2.** A, Variability of impedance measurements at each electrode in the non-MD ear. Dotted line represents standard deviation (SD) of measurements from mean within each electrode, and dashed line represents variance (VAR) or spread of measurements within each electrode. B, Variability of impedance measurements at each electrode in the MD ear. Dotted line represents standard deviation (SD) of impedance measurements from mean within each electrode, and dashed line represents variance (VAR) or spread of measurements within each electrode. MD indicates Ménière’s disease.



