Biomechanics and Immunomediated Mechanisms in the Cochlear Implant Rejection

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Abstract

Patients with bilateral and severe neurosensorial hypoacusia are candidates for cochlear implantation. Even though it’s a relatively modern procedure, it has a very low rate of complications (5-10%), making it a safe surgery. The Cohen classification is currently used to measure and group the complications of this surgery, classifying them according to the moment in which they appear whether it’s intraoperative or postoperative time.

We present the case of a female patient who suffered the extrusion of the implant several months after the procedure. She was re-operated successfully, but she returned months later with a second extrusion of the same implant. We did not find any other report of this phenomenon in the revised literature. We present this case with pictures of the repair surgery and the patient’s current condition. We also analyze possible causes of this cochlear implant rejection, citing both magnetomechanical and immunomediated factors.

Keywords: Cochlear implant; Complications; Extrusion; Biomechanic; Immunity

Introduction

One of the first published papers about the complications of cochlear implants (CI) was written by Thielemeir [1]. The complications can be classified as major and minor, depending on their severity. Using this classification Cohen [2] and Lehnhardt [3], placed major complications around 2.5% to 15%, most of them related to the graft and erosion of the posterior wall of the external auditory conduct. Minor complications (facial palsy, instability, tympanic membrane perforations, seromas, etc) varied from 6.2% to 25%, all of them temporary.

It’s interesting to note how various authors state that the rate of complications tends to go down, because of the experience acquired and the correction of flaws in the surgical technique [4,5].

While complications related to rejection of the CI are mentioned in literature, there has been no real analysis of the possible etiopathology of the phenomenon.

Local adverse effects of any CI can be immediate: edema; intermediate: abscess formation and late: foreign body- induced granulomas [6].

Inert materials like silicone and ceramics are biologically active. Evidence of their importance in the etiology of implant rejection is provided by the presence of granulomas and autoantibodies in the implant’s periphery and the absence of infection in the extruded materials analyzed.

By itself, the cochlear implant has fibrogenic capabilities and can migrate. There is a physiologic response from the host to the foreign body: the biomaterial’s placement in the body produces a local acute inflammatory response, similar to a foreign body reaction. Afterwards a fibroblastic reaction, with the presence of collagen an elastin, takes place and finally, the late granuloma formation as an immunomediated reaction occurs [7].

Faced with this rejection phenomenon we can ask ourselves: Why have adverse effects been described only with the CI and not with other uses? Also, what to do when faced with patients with a personal or familial history of autoimmune disease or serum autoantibody presence? Is it possible to look for high risk markers?

Materials and Methods

We present a 56 year old female patient with a medical history of deep bilateral hypoacusia, depression, type II diabetes and a former pack and a half- a day smoking habit. She had bilateral neurosensorial endocochlear hypoacusia (NSH) proven by brainstem auditory evoked potentials (BERA) with a more severe left-ear deficit. She received a left cochlear implant (Medel model PULSARci 100) on July 2006, with good functional adaptation.

During a routine follow-up visit on March 2008, signs of extrusion of the implant were observed. There were no external signs of infection Figure1 and 2. She was taken to the OR for rotation-relocation of the internal receptor of the same CI in a new bone bed, avoiding the scar lines of the myocutaneous flap and reinforcing it in proximity to the skin.

The macroscopic findings of the bone bed of the internal receptor showed osteogenesis that extruded it towards the surface. There was also atrophy on the internal face of the myocutaneous flap that was in contact with the receptor Figures 3-5.

The intraoperative control telemetry was satisfactory.

Results

On a follow-up rehabilitation visit, when the CI was reconnected the patient reported that the voices "sounded far away” and that she relied a lot on lipreading. In the following controls, word discrimination was poor in both open and closed spaces and telephone speech intelligibility also declined.

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Received December 22, 2011; Accepted January 10, 2012; Published January 12, 2012


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The implant’s subjective functionality was also poorer than in the original procedure. At the moment, the patient possesses 6 active electrodes and has started rehabilitation, recognizing up to 3 distinct vocal groups.

During the patient’s clinical follow-up, in November 2008, the start of a new CI extrusion was observed Figure 6. There was thinning in some areas of the cutaneous flap and macroscopic granulomas present on its surface. Given the patient’s circumstances, we planned a new repair procedure, or even the extraction of the extruded implant and a CI placement on the contralateral ear. The patient, nonetheless, rejected any new surgical procedures.

Discussion

Cochlear implant surgery is considered as a safe surgical option that can achieve acceptable results. The complications of the surgery usually follow Cohen’s classification: intraoperative or postoperative [8]. Among the intraoperative, the most noteworthy are CSF leaks, facial nerve injuries, electrode placement alterations and excessive thinning of the posterior wall of the external auditory canal. Postoperative complications can have an early or delayed onset [9]. Wound infection, device migration, facial nerve palsy, meningitis and instability are the most important immediate complications [10,11].

Regarding complications immediate to the surgery we would highlight:

1. Incision: any incision can be used, as long as the flap completely covers the device and it has an adequate blood supply.

2. Flaps: The possible complications include infections and hematomas in the posteroinferior triangle of the flap [12]. It is more likely to occur in adults with an excessive occipitomastoid curve and children because of their smaller size.

3. The mastoidectomy: Because of the CI antenna’s memory, extending the surgery posteriorly could cause a protrusion.
Figure 6: New extrusion of the implant with thinning and granuloma formation at the flap.

of the antenna and its posterior extrusion. This would be a mechanical traction-expulsion process [13].

4. The receptor’s anchoring: The receptor needs to always be securely anchored to bone. Failure to do so could cause an attraction from the magnet to the receptor and its extrusion by a mechanomagnetic mechanism.

Regarding the surgery’s possible early postoperative complications:

1. Displacement of the coil: if the antenna isn’t properly secured to the bone, the material’s memory can cause a prolapse at the superficial layers. The incision has to be separate from the internal device to avoid extrusion by traction.

2. Coil extrusion: The internal device’s position is very posterior to the occipitomastoid suture. Flaps that are thinned in an attempt to reduce impedance, could leave the coil in contact with the incision.

Regarding the late postoperative complications: The extrusion of the implant is an infrequent form of the set of complications that includes mastoiditis, meningitis, facial nerve stimulation and a technical failure in the implanted system [14,15].

To study the possible complications of a CI, while centering on the one that’s the object of this study, that is the extrusion of the implant, we will first analyze the factors that can lead to the rejection of the implant. That is, the pre-surgical factors. After those, the hypothetical factors that could lead to the extrusion of the implant over biological rejection by the host.

Most of the reviewed publications describe that extrusion of the CI is produced by decubitus damage at the myocutaneous flap. On occasions, surgical reintervention is necessary, with a new flap or even re-implantation. In our case, while it’s true that the receptor’s extrusion occurred as it fistulized through the flap, the receptor wouldn’t have been exposed had there been no osteal neogrowth in the receptor’s bone bed. This osteogenesis was probably motivated by an immunemediated rejection by the host and/ or the potent mechanistic extrusive effect by the CI’s magnet [16-18].

According to the available literature, re-implantation of the receptor by itself is enough to adequately resolve the complication. However, in this particular case, this procedure hasn’t solved the problem. That’s why we believe that it might be more appropriate to implant the other ear that to re-implant the ear where the extrusion occurred.

Regarding the biological complications we will say that the local adverse effects of any implant can be immediate: pain, edema; intermediate: swelling, abscess formation, necrosis and altered healing; and late: nodule and granuloma formation.

The different types of granulomas secondary to implants can have different anatomopathological compositions, whether they are sarcoid, tuberculoid, necrobiotic, suppurative or foreign body [6], as was the case in this patient.

If the implant cannot be phagocytized, due to the size of its particles or its vehicle, a foreign body reaction, with granuloma formation and local atrophy, is induced. The anatomopathological base of granulomas is a diffuse infiltrate that affects the dermis, with or without inclusion bodies, histiocytes, and multinucleated cells with sclerotic stroma. They can appear within months or even years. Local and oral treatments with corticosteroids haven’t achieved good results [7]. The definitive treatment is the extraction of the implant [19].

Some components of the CI, like the inner receptor, can use silicone or ceramics among other vehicles. In this case it’s a model PULSAR ci 100-Mede implant, which has an internal receptor made out of ceramics. The ceramic is a type of clay, a plastic mineral substance made from the hydrated silicates of different metals, routinely used in medicine among other industries.

When used for medical purposes, the material’s great hardness can be an advantage or an inconvenience. These compounds, like silicones among others, are inert and biologically active materials. That’s the reason why, before the usage of any implant, the following steps should be part of the protocol: mutagenesis testing, citotoxicity tests, sensibilization testing and trophoblastic activity tests.

This particular case is an extrusion of the CI with ossification of the receptor bone bed and with atrophy and granuloma formation at the flap and surrounding tissues, that had required reintervention, and that has now, once again, presented the same type of late complication. It’s the first reported such case of a second extrusion of a CI, according to our bibliographical review. It’s also the first article that tries to explain the reasons for these chain rejections.

Conclusions

1. Cochlear implantation surgery is considered a safe option with an acceptable result. Its complications can be classified as major or minor, depending on their severity and if a second surgery was needed.

2. This case can be helpful in raising certain fundamental questions like: What type of ethiopatogeny is involved in osteogenesis? And, is this ossification primary or secondary to the rejection? Also, what role does the magnet play in the establishment and progression of the implant’s extrusion? The convexity of the skull could help the antenna’s memory to cause an extrusion of it. But, why does this happen in some patients and not in others?

3. The implant has phylogenetic capabilities and can migrate. There is a physiological response from the host to the foreign material (implant). The placement of the biomaterial produces a local foreign-body reaction in the host. But, why does this happen in some patients and not in others?
4. We open a door to a research subject: finding genetic locators that can help identify before the surgery which patients are more likely to reject the implant.

References


