Development and evaluation of the Nurotron 26-electrode cochlear implant system

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ABSTRACT

Although the cochlear implant has been widely acknowledged as the most successful neural prosthesis, only a fraction of hearing-impaired people who can potentially benefit from a cochlear implant have actually received one due to its limited awareness, accessibility, and affordability. To help overcome these limitations, a 26-electrode cochlear implant has been developed to receive China’s Food and Drug Administration (CFDA) approval in 2011 and Conformité Européene (CE) Marking in 2012. The present article describes design philosophy, system specification, and technical verification of the Nurotron device, which includes advanced digital signal processing and 4 current sources with multiple amplitude resolutions that not only are compatible with perceptual capability but also allow interleaved or simultaneous stimulation. The article also presents 3-year longitudinal evaluation data from 60 human subjects who have received the Nurotron device. The objective measures show that electrode impedance decreased within the first month of device use, but was stable at a slight increase at the end of two years. The subjective loudness measures show that electric stimulation threshold was stable while the maximal comfort level increased over the 3 years. Mandarin sentence recognition increased from the pre-surgical 0%-correct score to a plateau of about 80% correct with 6-month-use of the device. Both indirect and direct comparisons indicate indistinguishable performance differences between the Nurotron system and other commercially available devices. The present 26-electrode cochlear implant has already helped to lower the price of cochlear implantation in China and will likely contribute to increased cochlear implant access and success in the rest of the world.

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1. Introduction

Since the United States Food and Drug Administration (FDA) approved the first cochlear implant in 1984, more than 300,000 hearing-impaired people worldwide have used electric stimulation of the auditory nerve to derive benefits from restoring speech perception in post-lingually deafened adults to developing language in pre-lingually deafened children [Clark, Hochmair, Wilson,
Eisenberg, Chouard and Merzenich references in the same issue]. Cochlear implant technology has evolved from a single-electrode analog device to multi-electrode devices with interleaved stimulation and in vivo neural recording. Three major companies, Cochlear Corporation in Australia, Med El in Austria and Advanced Bionics LLC in the United States, control essentially the entire market, with Cochlear Corporation being the dominating player. Despite clearly documented benefits, technological advances and commercial success, the cochlear implant is still limited to about 10% of potential candidates in developed countries and much less in developing countries (Zeng, 2007). The high cost of the cochlear implants has not changed in the last 30 years despite the increased volume and is still a major factor limiting market access. Developing a low-cost, high-performance cochlear implant system has long been recognized as an effective, perhaps the only means to increase competition and hopefully market access, particularly in developing countries (An et al., 2007; Wilson et al., 1998; Zeng, 1995).

The present article describes the conceptualization and process in developing and evaluating the Nurotron 26-electrode cochlear implant system, which started with a technology transfer from the University of California in 2006, received CFDA market approval in 2011 and CE Mark in 2012. As of July 2014, the Nurotron device has been implanted in 1500 deaf subjects in China, 2 in Columbia and 2 in India. First, the philosophy is presented on the design and development of the overall system specification. Second, specific design and verification of the system components are laid out from the external sound processor and radio frequency transmission to the internal receiver, stimulator and intracochlear electrode array. Third, systematic evaluation results are presented from objective measures to subjective functional assessments in 60 human subjects who participated in the initial CFDA clinical trial. Finally, relevant research and socio-economical impact are discussed.

2. System specifications

Cochlear implant technologies have converged in the last 30 years – from single-electrode analog stimulation to multi-electrode interleaved stimulation (e.g., Wilson et al., 2008; Zeng et al., 2008). Fig. 1 presents the functional block diagram of a contemporary cochlear implant system, consisting of an external unit, a transcutaneous radio-frequency (RF) transmission unit, an internal unit, and a fitting unit that is not worn by the user but used only by clinicians to adjust connection and stimulation parameters for optimal performance. The external unit is often called a speech or sound processor and contains a digital signal processor (DSP) to control signal flow from environmental sounds to RF transmission. The internal unit includes a hermetically sealed application-specific-integrated-circuit (ASIC) that derives power and decodes information from the RF signal, while delivering electrical stimulation to the electrodes, measuring feedback signals from the electrodes and transmitting these measurements back to the sound processor or the clinical fitting system.

The Nurotron cochlear implant system was developed based on the following design philosophy. First, the device should meet technical specifications that are critical to safety and performance of a contemporary cochlear implant. The European standard EN 45502-2-3 (2010) and additional modern practices were followed to conform to safety considerations; multi-channel, multi-stimulation strategies were employed to meet performance needs. Second, the device should have a flexible architecture to meet future needs such as virtual channels or fine structure encoding that could improve speech recognition in noise, tonal language understanding and music appreciation. With this philosophy in mind, the ASIC contained 4 current sources to provide simultaneous stimulation of 2 or 3 electrodes to potentially produce 47 or more spectral channels. Third, the device should have the capacity to support other neural prosthetics applications such as auditory nerve implants, auditory brainstem implants, retinal implants, or deep brain stimulation. To provide this level of expandability, the design included 2 reference electrodes and 24 active stimulating outputs that could be configured as 2 × 12, 3 × 8, or 4 × 6 surface electrodes or penetrating bundles.

Fig. 2 illustrates the Nurotron cochlear implant system. Panel A shows the behind-the-ear sound processor that contains dual microphones, control buttons, display lights, battery case, and an RF transmission coil. A body-worn sound processor (not shown) is also available for extended battery life or as preferred by some user groups. Panel B shows the internal components of the system that contains a gold RF receiving coil, a removable magnet (with “X” sign in the middle of the coil to designate polarity), a titanium case, a plate platinum reference electrode (beside the logo in the case), a ring platinum reference electrode (at the exit of the electrode array), and a straight 24-contact electrode array. The following section describes specific design and verification for the Nurotron cochlear implant system.
The CIS strategy employs a fixed number of analysis and stimulation channels, with the actual number being determined by the number of a subject’s usable electrodes up to 24. The APS strategy is similar to an “n-of-m” strategy, with the maximum number of analysis channels being 24 while the number of stimulation channels being typically 6 to 8, dependent on the energy distribution of the input sound and the number of available electrodes. The APS strategy increases stimulation rate for improved representation of temporal fine structure. The Symphony strategy combines virtual channel and peak selection strategies to additionally improve the spectral fine structure, in which simultaneous stimulation of two electrodes is used to generate intermediate pitch percepts between two these two electrodes.

Fig. 3 shows the spectrum (upper panel) and electrogram (lower panel) of a Mandarin sentence “今天你去那里?” (Where are you going today?). In this example, the electrogram shown is the output generated by the APS strategy, which divides a continuous frequency band between 100 and 8000 Hz into 128 linearly-spaced frequency bands. The 128 frequency bands are then combined into 24 analysis bands based on the Greenwood map spacing, of which 8 channels with peak energy are selected to deliver electrical stimulation to the appropriate electrode sites. The APS was the default strategy used to produce performance data reported in the user performance result section of this study.

3. Design and verification
3.1. Sound processing

Advanced sound processing is critical to cochlear implant performance. First, sound pre-processing is needed to ensure audibility and clarity of speech sounds. The Nurotron standard pre-processing includes sensitivity control, automatic gain control, single microphone noise reduction, and mixed input selection. Second, sound processing needs to convert analog sound into electrical pulses. The Nurotron DSP employs 3 processing strategies, including continuous-interleaved-sampling (CIS), advanced peak selection (APS) and virtual channel (Symphony) strategies. The CIS strategy employs a fixed number of analysis and stimulation channels, with the actual number being determined by the number of a subject’s usable electrodes up to 24. The APS strategy is similar to an “n-of-m” strategy, with the maximum number of analysis channels being 24 while the number of stimulation channels being typically 6 to 8, dependent on the energy distribution of the input sound and the number of available electrodes. The APS strategy increases stimulation rate for improved representation of temporal fine structure. The Symphony strategy combines virtual channel and peak selection strategies to additionally improve the spectral fine structure, in which simultaneous stimulation of two electrodes is used to generate intermediate pitch percepts between two these two electrodes.

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3.2. Radio frequency transmission

The RF unit serves 4 functions, including (1) providing power to the implant circuit with reasonable transmission efficiency, (2) transmitting data to the implant circuit with high reliability, (3) synchronizing between the external processor and the implant circuit while providing timing for the latter, and (4) transmitting feedback signals to monitor the implant circuit power and status and to measure electrode impedance and neural responses to electric stimulation. To accomplish these functions, the Nurotron external processor employs Pulse Width Modulation (PWM) to encode digital signals and Amplitude Shift Keying (ASK), a method of amplitude modulation, to balance transmission efficiency and reliability. A 16-MHz sinousoid is chosen as the carrier frequency. A Class-E amplifier is employed to achieve high efficiency by maintaining the loaded RF circuit in a quasi-resonant status.

Fig. 4A shows the use of pulse width modulation to encode “0s” and “1s”, in which bit “0” is coded by 5 RF cycles whereas bit “1” by 10 RF cycles. Fig. 4B shows the actually received 16-MHz RF waveform (yellow traces) in the implant circuit. Due to the Class-E amplifier’s resonance status, rising ramps and post-modulation ringing are apparent in the received waveform. In addition, environmental interferences, variations in scalp thickness, and relative movement between the external and internal coils all distort the received RF signal. Therefore, a robust decoder is needed to reliably retrieve the digital information. Hysteresis trigger levels are employed so that signal patterns with 3–7 RF cycles are all decoded as bit “0”, whereas that with 8–12 cycles as bit “1”. Fig. 4B shows the recovered envelope of the PWM-coded “0” and “1” (green traces). This PWM-coded bit signal also provides a 1-MHz clock for the implant circuit. For back telemetry, a load modulation method, as commonly used in RFID, uses the same pair of coils to transmit the implant related data to the external processor. The implant related data include internal status of the ASIC, compliance voltage, stimulation waveform, impedance and neural responses.

3.3. Receiver and stimulator

The implant circuit receives RF signals to derive power, then to decode data and command frames to produce electric stimulation. Common techniques such as parity check, cyclic redundancy check, data boundary check, and handshaking protocols are used to validate data transmission integrity. Fig. 5A shows the layout of the Nurotron ASIC receiver and stimulator. The die size is 3594 × 2499 µm². The core is a hybrid CMOS chip with mixed digital and analog circuit. The chip contains 4 current sources, ensuring reliability and enabling both interleaved and simultaneous stimulation, including two-intracochlear-electrode
Several measures have been implemented in the ASIC to ensure safe electric stimulation. First, biphasic pulse stimulation should be charge balanced, resulting in less than 100 nA DC bias current (EN 45502-2-3: 2010). Fig. 5E shows accumulated DC current level as a function of biphasic pulse current level. Without an active discharge circuit in the ASIC, the DC level exceeds the 100-nA safe limit at 800-μA current level and above. With the discharge circuit, the DC level was at 33 nA or less over the entire pulse current level.

Second, to further balance residual charge that may be applied to tissues, a capacitor is serially connected to each of the electrodes. Third, to avoid overstimulation causing tissue damage, maximal charge of any phase is hardware limited to 250 nC (Shannon, 1992).

Fig. 5F shows overstimulation protection model (dashed line) and actually measured maximal charges under different pulse duration and current level combinations (circles). Under no circumstance, does the implemented maximal charge exceed 250 nC.

Implemented on the ASIC chip is a programmable gain amplifier with a maximal gain of 60 dB and a 12-bit analog-to-digital convertor for back telemetry. As a result, a 10 μV change can be detected to support recording of neural responses. Electrode impedance can be measured with 5% tolerance.

### 3.4. Electrodes (24 + 2)

A final step towards cochlear implant hearing is electric stimulation of the auditory nerve via a 24-intracochlear-electrode array. Optimally, this electrode-nerve interface should support high levels of subject performance, minimize the occurrence and severity of insertion related trauma, and demonstrate excellent long-term reliability. The physical characteristics of individual cochlear implant electrode designs have been clearly associated with the incidence of intracochlear trauma in studies using human cadaver temporal bones (Rebscher et al., 2008, 1999; Wardrop et al., 2005a; Wardrop et al., 2005b) and intracochlear trauma has been correlated with significant reductions in subject performance (Aschendorff et al., 2007; Carlson et al., 2011; Finley et al., 2008). Based on these studies, it is important for the electrode array to be sufficiently small to fit within the wide variety of scala tympani dimensions. Additionally, the Nurotron electrode array incorporates greater stiffness in the vertical plane of the cochlear spiral than in the horizontal plane which has been shown to be one successful strategy to minimize the occurrence of vertical deviation into the scala media or scala vestibuli.

The Nurotron electrode consists of an injection molded silicone elastomer carrier supporting 24 intracochlear stimulating contacts and 2 extracochlear reference electrodes. Fig. 6A shows the flexible tip and 7 most distal electrode contacts. The active contacts have an exposed surface area of 0.2 mm² and are located at intervals of 0.8 mm on the inner surface of the electrode to be oriented toward target neurons in the modiolus. The maximum charge density is 1.25 μC/mm² (=250 nC/0.2 mm²). Taking into account that a platinum contact with smooth surface has a roughness factor of 1.4, the actual maximum charge density is more likely 0.9 μC/mm², well below the safety limit of electric stimulation (Brummer et al., 1983).

Fig. 6B shows the contact sites and leads from the opposite side of the array. The tapered ends of each contact are securely embedded in the silicone carrier. Fig. 6C illustrates the coiled wire lead cable. The total recommended insertion length of the array is 22.0 mm, corresponding to an insertion depth of roughly 400° by a straight electrode array in an average-sized cochlea (e.g., Franke-Trieger et al., 2014). These 24 active intracochlear contacts and the relatively deep insertion support high levels of subject performance as reported below. To accommodate the size of the human scala tympani, the Nurotron electrode array is 0.70 (H) × 0.68 mm(W) at the tip and 0.93 mm(H) × 0.68 mm(W) at the base, which fits

![](image.png)

**Fig. 4.** The Nurotron RF encoding scheme: Panel A shows the use of pulse width modulation to encode “0s” and “1s” (Panel A) and Panel B shows the actually received 16-MHz RF waveform (yellow traces) and the recovered “0” and “1” temporal envelopes (green traces).
within all documented human scala tympani cross sections (Rebscher et al., 2008). Fig. 6D shows that the Nurotron electrode array fits within the outlines of a set of randomly selected human scala tympani cross sections in the base (90°) and near the electrode tip at 360°.

As described above, the Nurotron electrode is mechanically stiffened in the vertical plane as a strategy to reduce the incidence of damage resulting from vertical deviation of the electrode tip. Physical measurements of the array indicate a ratio of 2.5–3.0 between vertical stiffness and horizontal stiffness, similar to that reported for other electrode designs which have demonstrated significant reduction in insertion associated trauma (Rebscher et al., 2008; Wardrop et al., 2005a). Preliminary temporal bone studies, and post-surgical clinical radiography, have confirmed that the Nurotron array can be reliably inserted to the 22 mm recommended insertion depth without significant trauma. To validate reliability, connection of the electrode array to the implanted stimulator, repetitive flex testing of the spiral coiled interconnect cable and the intracochlear electrode itself have been conducted to meet the European standard (EN 45502-2-3:2010). A final design consideration in reducing traumatic insertions, which have been associated with poor cochleostomy placement, incomplete visualization of the basal scala tympani, angular misdirection of the electrode tip or a combination of these three
The tapered ends of each contact are securely embedded in the silicone carrier. The organized vertical arrangement of the wire leads is also visible in this image. Fig. 6C illustrates the coiled wire lead cable. A critical factor in minimizing insertion related trauma is the 10-nC safe limit (EN 45502-2-3:2010). Finally, the 15-min MRI scan produced no significant difference in temperature between the two containers, with a temperature rise in MRI compatibility at 1.5T has been verified.

3.5. Packaging and MRI compatibility

Twenty-five platinum feedthroughs are placed in the bottom of the titanium case and connected to an internal PCB on one end, the 24-intracochlear electrodes and the ring reference electrode on the other. The titanium case has a volume of 12.8 ml (36.4 x 33.0 x 6.9 mm³), but with only 3.9-mm spacing being expected between the skin and the mastoid surface because a 3-mm deep recess is routinely drilled in the bone to securely house the implant and minimize its movement. Three additional platinum feedthroughs on the sidewall of the case are connected to the two ends of the RF coil and the plate reference electrode, respectively. Silicone coating and molding form and shape the feedthroughs on the sidewall of the case are connected to the two ends of the implant. Biocompatibility tests were performed and approved by the CFDA testing facility before human clinical trial. Hermetic seal, measured by the helium-gas leakage rate, is less than 1 x 10⁻⁹ atm cm²/second (EN 45502-2-3: 2010).

MRI compatibility at 1.5T has been verified for the Nurotron cochlear implant. In the heat generation test, the implant without the magnet, was placed into a 15-ml container filled with 9 g/l saline. An identical container without the implant served as control. 15-min MRI scan produced no significant difference in temperature between the two containers, with a temperature rise in the implant case or at the electrode tip being less than 2 °C. In addition, the MRI scan produced a charge of 0.06 nC, less than the 10-nC safe limit (EN 45502-2-3: 2010). Finally, the 15-min MRI operation produced no noticeable change in function and performance of the implant device.

4. System evaluation

4.1. Methods

4.1.1. Subjects

Sixty subjects participated in the Nurotron cochlear implant clinical trial from December 2009 to October 2010. They were 34 males and 26 females, with a mean age of 26 ± 12 years old (range = 6–59 years old). The cochlear implantation candidacy criteria followed the standard issued by Chinese Ministry of Health (China, 2007). The mean duration of deafness, defined as bilateral pure-tone-average thresholds (0.5, 1, 2 and 4 kHz) greater than 85 dB HL, was 7 ± 5 years (range = 0.3–15 years). The actual pre-surgical pure-tone-average thresholds were 107 ± 11 dB HL. The etiologies included ototoxicity (n = 31), enlarged vestibular aqueduct syndrome (7), sudden onset hearing loss (5), meningitis (2), noise exposure (2), and unknown causes (13). These subjects were from 18 provinces in China. Each subject and his or her family consented to participate in the clinical trial, with a protocol that was approved by the local Human Research Ethics Committee of each of the five participating hospitals, including Shanghai Fudan University (n = 23), Beijing PLA General Hospital (15), Beijing Tongren Hospital (10), Zhejiang University (7), and Chongqing Medical University (5). At the end of the first year, 3 subjects dropped out of the clinical trial, with 2 being unable to be tested at the 4- and 6-month periods and 1 being disqualified for having a pre-surgical pure-tone-average threshold that was 5 dB lower than the 85 dB HL inclusion criterion. The one-year clinical trial data were submitted to Chinese FDA and received its approval on August 19, 2011. However, follow-up continued with data being collected in 48 subjects until the end of the third year; the additional 9 subjects dropped out due to their inability or unwillingness to be tested at the specified times.

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4.1.2. Surgery

Twelve neurotologists, who all had prior experience in cochlear
implant surgery, used a soft surgical approach to perform mas-
toidectomy and cochleostomy in the 60 participants. An incision
was made in the skin behind the ear. A 0.44-mL, or 440-mm³, bed
was created to securely house the receiver and stimulator case
before entry into the middle ear. The facial recess was opened, and
the bone anterior to the facial nerve was removed to provide wide
exposure of the round window niche. A small cochleostomy hole
was made in a position inferior and anterior to the round window
niche with a 1.0- or 0.5-mm diamond burr. The cochleostomy hole
was drilled until the blue lining of the endosteum became visible.
A small amount of sodium hyaluronate was gently injected into the
scala tympani to prevent fluid leak and entry of foreign bodies
such as bone dust. Suction was prohibited at this stage to avoid loss
of perilymphatic fluid. The electrode array was then inserted into
the scala tympani. The cochleostomy was then sealed with a small
amount of connective tissue. After implantation, electrode
impedance and electrically evoked stapedius reflex were typically
measured to verify device integrity and functionality. The total
time of implantation ranged from 30 to 120 min, dependent on
surgeon’s experience and subject’s condition. Although all 12
surgeons encountered no significant difficulty in inserting the
electrode array, 4 recommended an insertion tool be used in the
future.

To estimate the electrode position and insertion depth (Verbist
et al., 2010; Xu et al., 2000), 48 subjects had X-ray of the cochlear
implant after their surgeries. The average insertion depth was 449 mm,
with a standard deviation of 79 mm, which was slightly deeper than
the designed insertion depth of 400 mm. Of these 48 subjects, 37 had
all of the 24 electrodes properly placed in the cochlea (e.g., Fig. 7A),
whereas 11 had a buckled or mislocated electrode array, especially
near the round window (e.g., Fig. 7B), indicating the need for
further evaluation of the electrode design and surgical procedure.
The effect of a buckled or mislocated electrode array will be
discussed in the discussion section.

4.1.3. Stimuli

Two open-set tests of Mandarin sentence recognition were used
for evaluation of the Nurotron cochlear implant performance. The
House sentence recognition test included 10 lists of 10 phonetically-balanced sentences with each containing 7 key words
(Fu et al., 2011). The 301 sentence recognition test included 12 lists
of 11 sentences with each containing 6–8 key words, for a total of
57 words (Xi et al., 2012). Percentage of correctly identified words
was used as the open-set sentence recognition score. Because no
statistical difference was found between the two tests, the averaged
scores were used in data analysis and reported here. In addition, 3
closed-set tests of Mandarin consonants, vowels, and tones were
used, in which each test had four alternatives and the subject had to
choose one of them (http://www.tigerspeech.com).

The default speech processing was the multi-peak APS strategy.
The default stimulation was monopolar mode using both reference
electrodes (MP1 + 2). Electrode impedance was estimated by a
single biphasic pulse with a 40-μs pulse duration. The threshold
and comfort loudness levels were estimated by a 500-μs, 1000-Hz
biphasic pulse train. The pulse duration was either 25 or 50 μs/
phase; Speech stimuli were presented to the subject at 65 dB SPL in
a sound field condition.

4.1.4. Procedures

As part of the informed consent, all subjects agreed to participate
in a pre-surgical test and 5 post-surgical tests at 1, 2, 4, 6 and
12 months. The additional tests at 24 and 36 months were volun-
tary. During each test, the subject went through a full battery of
audiometric tests including air-conducted pure-tone thresholds,
tifferential electrocardiogram, thoracic roentgenoscopy, visual examination for
physical examination, including vital signs, blood and urine tests,
electrocardiogram, thoracic roentgenoscopy, visual examination for
surgical complications such as swelling, infection, facial paralysis,
and hematoma. All tests were performed independently by

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audiologists and physicians, under supervision of a certified contract research organization.

4.2. Results I: objective measures

During the first year clinical trial period, 6 adverse events were noted, including 2 possibly related to surgery (swelling in the eyelid and swelling in the surgery area) and 4 unrelated to the surgery or the device (2 mild fatty liver cases, 1 acute nephritis, and 1 upper respiratory tract infection). The physicians determined that none of these adverse events was directly related to the Nurotron device.

Fig. 8 shows impedance as a function of electrode position at the device switch-on time (upper panel) and as a function of device usage time (lower panel). The electrode impedance was 13 ± 3 kΩ, which was not significantly different as a function of electrode position [F(23,771) = 0.7, p > 0.05; univariate ANOVA]. However, the impedance averaged over the entire electrode array changed significantly with implant use [F(23,242) = 20.2, p < 0.01]. The average impedance decreased significantly by 6 kΩ from switch on to 1 month, stayed unchanged over the 12-month period (p > 0.05), and increased significantly by 2 kΩ at the 24-month evaluation (p < 0.05; Post Hoc tests with Bonferroni correction for these and other comparisons in the remainder of the results section).

4.3. Results II: psychophysical measures

The upper panel of Fig. 9 shows threshold (T) and comfortable (C) levels as a function of electrode position at the device switch-on time. Both T and C levels showed a significantly increasing trend as a function of electrode position at the time of switch on [F(23,711) = 3.1, p < 0.01, F(23,717) = 3.3, p < 0.01, for T and C levels respectively]. Compared with the most apical electrode (#1), post-hoc analysis revealed significant higher T levels for electrode 23 and 24 and higher C levels for electrodes 22, 23 and 24 (p < 0.05). Note that these different T and C levels cannot be attributed to electrode impedance, which was not significantly different between electrodes (Fig. 8).

The lower panel of Fig. 9 shows the averaged T and C levels over the entire electrode array as a function of the implant usage time. The T level had been stable over the 36 month period [F(7,226) = 0.4, p > 0.05]. Interestingly, the C level increased over time [F(7,226) = 2.3, p < 0.05], with the C level at 36 months being significantly greater than that at the time of switch on [p < 0.05]. This increased C level possibly reflects the increased tolerance to electric stimulation of the auditory nerve (Zeng, 2013).

4.4. Results III: functional measures

Fig. 10 shows that sentence recognition varied greatly among subjects (dotted lines) but improved significantly as a function of implant usage over the 36-month period [F(7,426) = 132, p < 0.01].
5. Discussion

In this section, the Nurotron device is compared against other devices from both technical and performance perspectives. Several relevant studies using the Nurotron device are also summarized and discussed. Finally, the socio-economic impact of the present developmental effort is discussed.

5.1. Technical comparison

Table 1 compares key technical parameters of the Nurotron cochlear implant against presently available devices by the three major cochlear implant manufacturers. Although the Nurotron device was conceived in 2005 and developed into a product in 2008, it is still technically comparable by today's standards. A reason for this comparability is that cochlear implant advances in the last decade have been mostly in pre-processing and cosmetics, with little changes in speech processing strategies, RF transmission and receiver and electrode arrays.

In terms of the external unit, all devices, including the Nurotron device, have similar technical parameters from a relatively wide input dynamic range (75–80 dB) and a frequency range (~8000 Hz) to the default processing strategy (CIS-like) and the number of maps in the processor (4–5). Note that the Nurotron processor has the shortest battery life, reflecting a lack of design consideration for optimal power consumption. Also note that the Nurotron processor lacks wireless connectivity, reflecting design priority considerations in 2005. Both power consumption and connectivity problems are being addressed by Nurotron, but are beyond the scope of discussion for the present study.

Although there is no manufacturer-specific systematic report on in vivo electrode array status, buckled or mislocated electrode arrays are believed to be present in all devices, e.g., as high as 40% electrodes mislocated in scala vestibuli in some cases (Holden et al., 2013; Kong et al., 2012; Skinner et al., 2007; Wardrop et al., 2005a, 2005b). It is unclear whether the incidence of the presently
reported 11 cases of buckled electrodes is comparable to other
devices, and whether these cases are due to the ineptitude of
some of the surgeons themselves, or their unfamiliarity with a new
device. Nevertheless, post-surgical analysis showed that the
average number of unusable electrodes is 4 (range = 0–13) in
subjects with buckled or mislocated arrays, compared with 1
(range = 0–10) in other subjects. The corresponding average sen-
tence recognition score was 70 ± 18%, compared with 79 ± 19% correct.
Neither the difference in the number of unusable elec-
trodes nor that in the sentence recognition score reaches a statis-
tically significant level (p > 0.05), possibly due to the relatively
small sample size. To highlight the complicated subject and device
interactions, the subject with the most unusable electrodes (=13)
actually had one of the highest sentence recognition score (=92%).
Except for having the most intracochlear electrodes, the key
technical parameters for the Nurotron RF and internal units are
roughly in the middle of other currently available devices. For
example, the 16-MHz RF carrier is lower than the 49-MHz AB de-
vice but higher than the 5-MHz Nucleus and 12-MHz Med El de-
vice; the 40-kHz total stimulation rate is higher than the 32-kHz
Nucleus device but lower than the 83-kHz AB and 51-kHz Med El
devices; the 4 current sources are more than the 1-source Nucleus
device but less than the 16-source AB and 24-source Med El de-
vice. As can be seen from the remainder of the discussion, these
differences in technical parameters reflect philosophical differ-
ences in design but have not produced any measurable differences
in performance.

5.2. Performance comparison

The Nurotron device has produced similar performance to other
devices in terms of basic objective and subjective measures. The
present electrode impedance (Fig. 8) and stimulation level (Fig. 9)
effects are generally consistent with a large body of existing studies
from other cochlear implant devices (e.g., Henkin et al., 2003, 2005,
2006; Hughes et al., 2001; Mosca et al., 2014; van Wermesken-
er et al., 2006). In addition to a larger sample size and longer
observation time (3 years vs. 1–2 years), the present study extends
previous studies in the following ways. First, the absence of elec-
trode position effect on impedance (Fig. 8A) suggests that electrode
impedance reflects mostly the electrode physical properties such as
area and the dynamic interplay between electrode and nearby
tissues in electric stimulation (e.g., Clark et al., 1995), whereas the
presence of electrode position effect on loudness levels (Fig. 9A)
suggests that behavioral measures reflect more the overall electric
field property, the survival nerve extent and distribution, and central factors (e.g., Tang et al., 2011). The non-monotonic function of
impedance versus time (Fig. 8B) and the monotonic function of
loudness versus time (Fig. 9B) lend further support for the idea that
increased loudness tolerance with prolonged electric stimulation
reflects more a decreased central gain in the brain rather than
changes in electric properties in the cochlea (Zeng, 2013). The
greater amount of this loudness change in children than in adults
(e.g., Henkin et al., 2003, 2005, 2006; Hughes et al., 2001) is
consistent with greater plasticity in the developing brain.
The Nurotron device has produced functionally indistinguishable
performance from other devices in speech recognition mea-
sures. Fig. 12A shows comparison in sentence recognition between
4 cochlear implants. The Advanced Bionics data were from 51
subjects who had used the HiRes processing strategy for 3 months
after prior 3-month experience with conventional strategies, CIS or
SAS (Koch et al., 2004). The Med El data were 14 subjects who had
used the fine structure processing strategy after an average 1-year
experience with the CIS strategy (Arnoldner et al., 2007). The
Cochlear data were from 55 subjects who participated in the Nu-
cleus Freedom North America clinical trial and had used the
Freedom device for 6 months (Balkany et al., 2007). As a compari-
son, the Nurotron data were from the 6-month performance, which
was significantly higher than the 3-month performance but lower
than, although not significantly different from, the 1-year perfom-
ance (Fig. 10). Despite the fact that Advanced Bionics and
Cochlear used English sentences, Med El used German sentences
and Nurotron used Mandarin sentences, and additionally these
clinical trials were conducted by different researchers and under

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different protocols, all 4 devices produced functionally equivalent performance between 60 and 80% correct for sentence recognition in quiet.

In a more tightly controlled study, researchers at Beijing 301 Hospital (Li et al., 2014) directly compared performance between Nucleus and Nurotron devices by recruiting two groups of 15 subjects who were matched in age (29 ± 13 vs. 25 ± 9 years old), duration of deafness (7 ± 5 vs. 7 ± 4 years), gender (9 males and 6 females vs. 10 males and 5 females), and other audiological and etiological factors. Using the same surgical and evaluation protocols including the same Mandarin speech test materials, they found no statistically difference in performance between these two devices. Fig. 12B re-plots the subjects’ sentence recognition data after 2-year usage of their respective Nucleus and Nurotron devices. The Nucleus users produced 87 ± 14% correct score while the Nurotron users produced 83 ± 21% score. The high level of speech performance by the Nurotron device as well as its ability to improve quality of life has been independently verified by other researchers at Beijing Tongren Hospital (Liu et al., 2014), Central South University Xiangya Hospital (Yu, 2013), and China Rehabilitation Research Center for Deaf Children (Yu, 2013).

5.3. Socio-economical impact

Thanks to pioneers such as William House, Blair Simmons, Robin Michelson (Eisenberg ref and Merzenich ref in this issue), as well as the three 2013 Lasker Award winners: Graeme Clark, Ingeborg Hochmair and Blake Wilson, the cochlear implant has helped restored partial hearing to more than 300,000 adults and children worldwide. Unfortunately, only a fraction of the current cochlear implant users reside in developing countries, despite the fact that these countries boast more than 80% of the world population. While a lack of competition, a lack of awareness and a lack of access all contribute to some extent, high price is the prohibitive factor limiting the widespread use of the cochlear implant (Zeng, 2007).

To our knowledge, the Nurotron cochlear implant represents the first time an implantable, active medical device has ever been developed and manufactured in a developing country. Although the commercial operation is still in infancy, the successful development of the Nurotron device, as shown here, has not only broken technical barriers, but also produced a significant socio-economical impact. Thanks to this emerging product and the competition it has brought about, at present, 3 deaf children in China can benefit from the cochlear implant for the price that would allow only child to receive a device several years ago. It is expected that the cochlear implant will continue to drop in cost while improving performance, benefitting hearing-impaired people living in both developed and developing countries.

6. Summary

The present article describes the development and evaluation of a 26-electrode Nurotron cochlear implant that is comprised of an external sound processor, a radio-frequency transmission link and an internal receiver and stimulator. The default sound processing strategy uses multi-peak interleaved stimulation, while the default mode is monopolar stimulation with an active electrode being one of the 24 intracochlear electrodes and both extra-cochlear electrodes being the reference. Sixty severe-to-profoundly hearing-impaired subjects had participated in a 1-year clinical trial with some of them being continuously followed up for 3 years. Similar to other commercially available cochlear implants, the Nurotron device showed a decrease in electrode impedance within the first month implant use, a stable threshold level, and an increased comfort level over the 3-year period. Mandarin speech recognition significantly improved from the pre-implantation level to 4-month usage, reaching a plateau high-level performance of about 80% correct after 6-month usage. Both indirect and direct comparisons indicated that the Nurotron 26-electrode cochlear implant and other commercially available devices produced indistinguishable performance.

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