Spatial channel interactions in cochlear implants

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Abstract
The modern multi-channel cochlear implant is widely considered to be the most successful neural prosthesis owing to its ability to restore partial hearing to post-lingually deafened adults and to allow essentially normal language development in pre-lingually deafened children. However, the implant performance varies greatly in individuals and is still limited in background noise, tonal language understanding, and music perception. One main cause for the individual variability and the limited performance in cochlear implants is spatial channel interaction from the stimulating electrodes to the auditory nerve and brain. Here we systematically examined spatial channel interactions at the physical, physiological, and perceptual levels in the same five modern cochlear implant subjects. The physical interaction was examined using an electric field imaging technique, which measured the voltage distribution as a function of the electrode position in the cochlea in response to the stimulation of a single electrode. The physiological interaction was examined by recording electrically evoked compound action potentials as a function of the electrode position in response to the stimulation of the same single electrode position. The perceptual interactions were characterized by changes in detection threshold as well as loudness summation in response to in-phase or out-of-phase dual-electrode stimulation. To minimize potentially confounding effects of temporal factors on spatial channel interactions, stimulus rates were limited to 100 Hz or less in all measurements. Several quantitative channel interaction indexes were developed to define and compare the width, slope and symmetry of the spatial excitation patterns derived from these physical, physiological and perceptual measures. The electric field imaging data revealed a broad but uniformly asymmetrical intracochlear electric field pattern, with the apical side producing a wider half-width and shallower slope than the basal side. In contrast, the evoked compound action potential and perceptual channel interaction data showed much greater individual variability. It is likely that actual reduction in neural and higher level interactions, instead of simple sharpening of the electric current field, would be the key to predicting and hopefully improving the variable cochlear implant performance. The present results are obtained with auditory prostheses but can be applied to other neural prostheses, in which independent spatial channels, rather than a high stimulation rate, are critical to their performance.

(Some figures in this article are in colour only in the electronic version)
1. Introduction

Electric stimulation serves as an effective means of activating nerves and controlling activity patterns in a wide range of man–machine interfaces, including sensory neuroprostheses (Rutten 2002), brain–computer interfaces (Pesaran et al 2006) and functional electrical stimulation (Peckham and Knutson 2005). Generally speaking, the electric stimulation of the neural tissue has a main advantage and a main drawback. The main advantage of electric stimulation is that the fast distribution of charges in the neural tissue allows the accurate reproduction of complicated temporal firing patterns. The main drawback is that the spatial selectivity of the excitation pattern is severely limited by the widespread nature of electric fields in the extracellular tissue.

As the most successful neural prosthesis to date, the cochlear implant (CI) has partially restored functional hearing to more than 200,000 profoundly deafened persons. Modern multi-channel CIs can deliver rapid electric stimulation at a rate as high as several thousands of hertz per electrode but have only 12 to 22 intracochlear electrodes, much less than the 3000 inner hair cells in a normal human cochlea (e.g., Zeng et al 2008). In spite of the success, modern CI users still face two major challenges at present. Firstly, the individual performance varies greatly among the present implant users. The reasons for this great variability remain unclear (e.g., Skinner et al 2002). Secondly, compared with normal-hearing listeners, CI listeners still have extreme difficulty in music perception, speech recognition in noise, and tonal language understanding (e.g., Zeng 2004). In addition to a lack of encoding of the temporal fine structure in the modern CIs (e.g., Wilson et al 1991), insufficient independent channels due to spatial interaction are likely responsible for the great individual variability and difficulty in music perception and speech recognition in noise (e.g., Zeng 2004, Smith et al 2002, Shannon et al 2004).

To increase the number of independent spatial channels, recent CI research has focused on either reducing channel interaction or taking advantage of channel interaction. First, coiled electrode arrays or positioners have been designed to place the electrodes closer to the neurons than earlier electrode arrays, presumably reducing stimulation threshold and channel interaction (e.g., Tykocinski et al 2001, Stickney et al 2006). Second, bipolar or tripolar stimulation mode can produce more narrowly focused electric fields than monopolar stimulation (e.g., Bierer 2007, Srinivasan et al 2010). Third, stimulation waveform shapes, such as anodic-first pulses, pseudo-monophasic pulses or tri-phasic pulses, may also increase spatial selectivity (e.g., van Wieringen et al 2005). Finally, stimulating two or more electrodes may produce a ‘virtual channel’ that is perceptually distinctive from each of the two real channels represented by the physical electrode (e.g., Bonham and Litvak 2008, Wilson et al 2003). Despite significant effort, these methods of increasing the number of independent channels have not yet produced improved CI performance (e.g., Berenstein et al 2008, Donaldson et al 2011, Chua et al 2011). The main reason for this unsatisfactory result is likely to be a lack of fundamental understanding of channel interactions in particular and its relationship to performance in general.

This study addresses this lack of understanding by focusing on spatial channel interactions at the electrical, neural and perceptual levels. Figure 1 illustrates three major sources of channel interactions and different measures that can be used to delineate these sources. At the physical level, the degree of electric field interactions between two stimulating electrodes depends on both electric stimulation parameters, such as amplitude and duration (e.g., Grill and Mortimer 1996), and electrode–tissue interface parameters, such as electrode configuration and impedance (e.g., Frijns et al 1995, Jolly et al 1996, Rattay et al 2001, Ranck 1975). Modern CIs have employed an electric field imaging (EFI) technique to measure potential distributions across the entire electrode array in the cochlea (e.g., Vanpoucke et al 2004a, 2004b).

Channel interactions also arise at the neural level, which can be totally independent of electric field interactions. Imagine an extreme case where only one neuron is present: no matter how many independent electrodes are provided, there is only one functionally independent channel at the neural level. Another example of neural interaction is in the time domain: electric fields can be simultaneously turned on and off, but neural interactions still occur if two stimuli are presented within the refractory period of the nerve fibers, e.g., up to several milliseconds (van den Honert and Stypulkowski 1984). Modern CIs use a non-stimulating electrode in the cochlea to record the electrically evoked compound action potential (ECAP) from a population of nerve fibers (Brown et al 1990, 1996, Rubinstein 2004). The ECAP can be used to quantify channel interactions at the neural level (Abbas et al 2004, Cohen et al 2003, 2004, Hughes and Abbas 2006b, 2006a, Eisen and Franck 2005, 2004).

Channel interactions can be reflected by behavioral measures, which may have a peripheral origin, or a
central origin, or both. For example, detection thresholds may be lowered by in-phase, sub-threshold simultaneous stimulations of two adjacent electrodes, but elevated by out-of-phase stimulation (de Balthasar et al. 2003, Eddington et al. 1978, Favre and Pelizzone 1993). This change in threshold is likely a result of electric field summations at the physical level. Another example is the elevated perceptual threshold in the presence of prior stimulation, which probably reflects refractory properties at the neural level (Nelson and Donaldson 2001). Finally, loudness summation, which is measured as increased loudness in response to two or more physically independent stimuli, is presumably a result of neural processing at the central level (Chatterjee and Oba 2004, Chatterjee and Shannon 1998, Abbas and Brown 1988, McKay et al. 1995, Shannon 1983).

This study used low-rate (⩽100 Hz) stimulation to focus on spatial channel interaction because potentially confounding temporal factors, such as residual charge and refractoriness, rarely last longer than several milliseconds. At these low rates with at least a 10 ms inter-stimulus interval, electric stimulation is unlikely to affect either the electrical field distribution or the neural and behavioral responses. For example, the electric dynamic range changes significantly for stimulation rates higher than 300 Hz but is essentially constant for stimulation rates lower than 100 Hz (Shannon 1985, Zeng et al. 2003, Eddington 2003, Eddington 2003). In contrast to previous studies, this work systematically measured channel interactions at physical, physiological and behavioral levels in the same CI subjects, allowing the assessment of relative contributions of electrical, neural and perceptual factors to channel interactions. The first experiment recorded the distribution of intracochlear potentials as a function of the electrode position in response to the simultaneous stimulations of one or two electrode sites. The second experiment measured ECAP growth functions at each of the 16 electrodes as well as neural interactions in three locations near the apical, middle and basal part of the electrode array. The third experiment evaluated changes in detection thresholds and loudness summation as a function of the phase difference between simultaneous stimulations of two electrodes. Several new measures were introduced to assess quantitatively the width of the excitation pattern and the degree of asymmetry in these measured channel interaction patterns. Correlation analysis was conducted to delineate the relationship of these different channel interaction measures and their relative contribution to speech performance. The implications of the present results are discussed in relation to the development of future CIs and other neural prostheses.

2. Methods

2.1. Subjects

Five CI users participated in this study. Table 1 displays the demographic information regarding these subjects. Three subjects (S1, S2 and S3) used a Clarion CII device, and the other two subjects (S4 and S5) used a HiRes 90 K device (Advanced Bionics Corporation, Valencia, CA). All subjects were implanted with a HiFocus electrode array without the positioner (1.1 mm separation between electrodes). The age of the subjects at the time of the experiments ranged from 46 to 71 years. The duration of implant use ranged from 1 to 5 years. The Institutional Review Board of the University of California, Irvine, approved the subject recruitment and testing protocols.

2.2. Electric field imaging

All electric stimuli were generated and controlled by the Bionic Ear Data Collection System software (BEDCS v1.17.208, 2006, Advanced Bionics Corporation, Valencia, CA). Monopolar electrode configuration (Ex) was used, where x represents the intra electrode number ranging from the most apical (1) to the most basal position (16), with the reference electrode being located near the implant receiver case outside of the cochlea.

EFI spatial profiles were obtained by applying an electric pulse at one electrode while recording the voltage potential at the remaining electrodes, using the customized BEDCS backward telemetry system. EFI spatial profiles were recorded for stimulating the apical electrode E1, middle electrode E9 and basal electrode E15. To test linearity, EFI spatial profiles were recorded by the simultaneous stimulation of two locations at E1 and E9, E9 and E15 or E1 and E15, respectively. The stimuli were single, charge-balanced, biphasic pulses with 53.9 μs/phase in the EFI experiment. The stimulation level was set to 50 μA with one-electrode stimulation or 25 μA with two-electrode simultaneous stimulation. The intracochlear potentials were acquired with a 9-bit analog-to-digital converter at a sampling rate of 55.6 kHz. The gain setting of the internal recording was 30.4 dB. The recording started before the onset of the stimulus and had a total duration of 8 ms.

### Table 1. Subject demographical information. The duration of deafness was defined as the time when the subject first noted hearing loss to the time when they received a CI. Percentage correct scores of HINT sentence recognition using clinically mapped speech processors were obtained in quiet and in speech-spectrum-shaped noise (10 dB signal-to-noise ratio).

<table>
<thead>
<tr>
<th>Subject</th>
<th>Age (years)</th>
<th>Device type</th>
<th>Duration of implant use (years)</th>
<th>Duration of deafness (years)</th>
<th>Sentence score in quiet (%)</th>
<th>Sentence score in noise (%)</th>
<th>Etiology</th>
</tr>
</thead>
<tbody>
<tr>
<td>S1</td>
<td>71</td>
<td>CII</td>
<td>5</td>
<td>20</td>
<td>94</td>
<td>70</td>
<td>Hereditary</td>
</tr>
<tr>
<td>S2</td>
<td>63</td>
<td>CII</td>
<td>3</td>
<td>61</td>
<td>72</td>
<td>33</td>
<td>Spinal meningitis</td>
</tr>
<tr>
<td>S3</td>
<td>68</td>
<td>CII</td>
<td>5</td>
<td>62</td>
<td>91</td>
<td>40</td>
<td>Unknown</td>
</tr>
<tr>
<td>S4</td>
<td>47</td>
<td>HiRes 90 K</td>
<td>1</td>
<td>2</td>
<td>74</td>
<td>6</td>
<td>Sudden hearing loss*</td>
</tr>
<tr>
<td>S5</td>
<td>46</td>
<td>HiRes 90 K</td>
<td>2</td>
<td>11</td>
<td>100</td>
<td>80</td>
<td>Unknown</td>
</tr>
</tbody>
</table>

*S4 had normal hearing in the left ear. The subject was implanted as a result of severe tinnitus after sudden hearing loss.*
Figure 2. Forward-masking artifact subtraction technique (top panel) and the resulting ECAP response (bottom panel). The ECAP amplitude is defined as the difference between the negative (N1) and positive (P1) peaks of the compound action potential. See the text for details.

2.3. Electric compound action potential

ECAP recordings were obtained using the same equipment and parameters as the EFI experiment, except for the following parametric settings. First, a 60 dB gain was used in the preamplifier to capture the relatively small neural potentials. Furthermore, 32 sweeps were averaged to increase the signal-to-noise ratio. The inter-sweep interval was 48 ms, resulting in a nominal rate of about 20 Hz for the ECAP recording. Finally, the stimulation level was selected to cover the entire dynamic range of the single electric stimulus. The dynamic range was defined as the electric current difference between the threshold and the most comfortable loudness level (MCL). The threshold of the single pulse was obtained using an adaptive, three-interval, forced choice (3IFC) with trial-by-trial feedback (Levitt 1971), while the MCL of the same single pulse was measured using a method of limits with the ascending sequence only (Zeng et al. 1998).

A forward-masking subtraction technique was used to remove the stimulus artifact (Brown et al. 1990). This subtraction technique took advantage of the neural refractory properties and used three biphasic (32.32 μs/phase) pulses, including (A) masker only, (B) probe only and (C) masker and probe with a masker–probe interval of 452.6 μs, which was assumed to be shorter than the absolute refractory period of the auditory fibers.

Figure 2 explains the rationale of using these three stimuli and shows the exemplary recording waveform. The masker only condition (A) produced a neural response to the masker $R_m$ plus its electric artifact $A_m$. The probe only condition (B) produced a neural response to the probe pulse $R_p$ plus its electric artifact $A_p$. The combined masker and probe condition (C) produced the same neural response to the masker but no neural response to the probe due to the refractory effect. Because the combined condition still produced the same electric artifacts $A_m$ and $A_p$, adding (A) and (B) and subtracting (C) effectively removed the neural response to the masker and canceled the electric artifacts, yielding a neural response to the probe. Finally, the neural response was enhanced by off-line averaging the responses to the anodic- and cathodic-leading pulses, which produced slight differences in latency between the two responses that needed to be corrected by the off-line procedure (Klop et al. 2004). The bottom trace of figure 2 shows a representative ECAP waveform. The ECAP amplitude is defined as the difference between the first negative peak (N1) and the following depolarization (P2) peak.

By applying the forward masker to one electrode and the probe to another electrode, we can estimate the spatial overlap in the neural excitation pattern between the two electrodes. In contrast to the electric field interaction, which only counts the overlap between two electric fields, the extent of the neural interaction actually measures the relative overlap between the population of neurons excited by the masker and the probe. To systematically measure the neural interaction across the cochlea, the masker was presented to all electrodes other than the recording electrode. The probe electrode was E1, E9 or
E15, with the adjacent electrode E2, E10 or E16 serving as the recording electrode, respectively.

2.4. Threshold interaction

The threshold was measured using a 3IFC with trial-by-trial feedback (Levitt 1971). The stimulus level was decreased with two correct responses and increased with one incorrect response. A reversal occurred with a change in the response direction from either correct to incorrect responses or vice versa. The step criteria were either 40 trials or 8 reversals, whichever occurred first. The step size for varying the signal level was 10 \( \mu \)A for the first two reversals and 2.5 \( \mu \)A for the next three to eight reversals. The threshold was obtained by averaging the values from the reversals with the smaller step size and approximated the 71% correct point on the psychometric function.

To measure behavioral channel interaction, the simultaneous stimulations of two electrodes were used with electrode E1, E9 or E15 being the first electrode site, while the remaining odd-numbered electrodes were being used as the second site. The stimuli were 100 Hz, 500 ms biphasic (107.8 \( \mu \)s/phase) pulse trains. The electric current of the second stimulation site was presented at two sub-threshold levels (20% and 60% of the threshold). Thresholds were obtained first at all electrodes with anodic-leading pulse trains. A positive probe level indicated that the probe and the masker were in phase, while a negative probe level indicated that the probe and the masker had opposite phases.

Similar to previous studies (Eddington et al. 1978, Shannon 1983, Stickney et al. 2006), the following simple linear equation was used to calculate the behavioral channel interaction index at the threshold level (CIIth),

\[
I_{\text{probe}} = I_{\text{th probe}} - \text{CIIth} \times I_{\text{masker}},
\]

where \( I_{\text{probe}} \) is the probe level needed to reach threshold in the presence of a masker, \( I_{\text{masker}} \) is the masker level and \( I_{\text{th probe}} \) is the probe threshold without the masker.

2.5. Loudness interaction

Different from threshold interaction, in which the masker was always presented at a sub-threshold level, the probe and the masker were both presented at a supra-threshold level (i.e. audible) in loudness interaction. Three supra-threshold levels, namely, very soft, medium soft and soft, were used in the loudness interaction experiment.

The MCL for all single electrodes was first measured using the method of limits with an ascending sequence only and used as a reference point. A loudness balance procedure was then used, in which two stimuli (A and B) were presented to the subject: A was single-electrode stimulation at MCL and B was dual-electrode stimulation with a fixed masker level. The subject was instructed to bracket the point of equal loudness by first adjusting the probe level on stimuli B to a level where B was just noticeably louder than A, then to a level where B was just noticeably softer, and then to the level of equal loudness. The subject clicked the ‘up’ or ‘down’ button on the computer screen to adjust the level. One electrode was used as the probe and the other electrode was used as a masker. The probe and masker electrodes were reversed to counter balance any pitch related influences on loudness judgment. For each pair of electrodes, six data points were obtained to construct the equal loudness contour.

To derive a new loudness interaction model, an exponential loudness growth function was used for single-electrode stimulation (Zeng and Shannon 1994),

\[
L = k e^{\alpha E},
\]

where \( L \) is the perceived loudness as an exponential function of electric stimulation \( E \), and \( k \) and \( \alpha \) are constants that depend on stimulation modes and parameters. For two-electrode simultaneous stimulation, the total loudness is presumably determined by three components: the no-overlapping component from electrode 1, the overlapping component between electrodes 1 and 2 and the no-overlapping component from electrode 2,

\[
L = (1 - \text{CII}_1)k e^{\alpha E_1} + \text{CII}_1 k e^{\alpha (E_1+E_2)} + (1 - \text{CII}_2)k e^{\alpha E_2},
\]

Rearranging equation (4), we obtain

\[
E_1 = \frac{1}{\alpha} \log \left( \frac{e^{\alpha E} - (1 - \text{CII}_1) e^{\alpha E_2}}{1 - \text{CII}_1 + \text{CII}_1 e^{\alpha E_2}} \right),
\]

Consider the following two boundary conditions. If \( \text{CII}_1 = 1 \), then total interaction occurs between the two electrodes. Equation (5) is simplified to

\[
E_1 = E - E_2.
\]

If \( \text{CII}_1 = 0 \), then no direct electric interaction occurs between the two electrodes. The total loudness is the sum of loudness of two independent electric stimuli, so equation (5) becomes

\[
E_1 = \frac{1}{\alpha} \log (e^{\alpha E} - e^{\alpha E_2}).
\]

Figure 3 shows three loudness scenarios representative of channel interaction. If there was no interaction between the masker and the probe, then the loudness of dual electrode stimulation would be the sum of the loudness evoked by the masker and the probe (no-interaction scenario where \( \text{CII}_1 = 0 \)). On the other hand, when significant interaction occurred, electric fields would be summed up before exponential transformation for the perceived loudness. The solid diagonal line represents this total interaction scenario (\( \text{CII}_1 = 1 \)). The dashed line represents an intermediate interaction case. The present loudness summation model is used to derive the loudness channel interaction index and to compare with other channel interaction indexes.
Figure 3. Representative channel interactions modeled by loudness summation.

3. Results

3.1. Channel interactions at the physical level

Figure 4 shows the group mean EFI, representing intracochlear potential data as a function of the recording electrode in response to electric stimulation on E1, E9 or E15 (indicated by the vertically arrowed line in the top, middle and bottom panel, respectively). First, note the small standard error of the mean (average = 2.6 mV, range = 1.8–8.0 mV), indicating a highly uniform electric field profile across these five subjects. Second, note an asymmetrical distribution of the EFI profile, with the apical side decaying more steeply (from E8 to E1 in the middle panel and from E14 to E1 in the bottom panel) than the basal side (from E2 to E16 in the top panel and from E10 to E16 in the middle panel). Third, note an apical plateau around 30 mV from E1 to E3 in the middle panel and 20 mV from E1 to E6 in the bottom panel that decays on the basal side.

Figure 5 shows measured (solid circle) and predicted (open circle) intracochlear potentials in response to three dual-electrode stimulation sites (E1 and E9 in the top panel; E9 and E15 in the middle panel; E1 and E15 in the bottom panel) in subject S2. Predicted intracochlear potentials assumed a simple linear summation of the measured intracochlear potentials from single-electrode stimulation. Data from the other four subjects (not shown) were also highly consistent, with the overall root mean squared error being only 2.3% between the measured and predicted values across all subjects. This accurate prediction of the dual-electrode stimulation data from the single-electrode stimulation data, coupled with the highly consistent across-subjects data, indicates that channel interactions at the electrical level can simply be modeled as a linear function.

3.2. Channel interactions at the neural level

Figure 6 shows the ECAP responses as a function of stimulation currents for all electrodes in the five individual subjects (all panels except for the bottom-right one). The dashed lines are linear fit to the individual electrode data (mean $R^2 = 0.81$, SE = 0.02, $p < 0.05$). S1 and S3 have similar ECAP thresholds and dynamic ranges across all electrodes, but S2, S4 and S5 have different thresholds and dynamic ranges between electrodes, with the apical electrodes more likely having lower thresholds than the basal electrodes. These individual differences in the dynamic range, particularly the different individual criterion to determine the MCL, introduce variability in both measures and their related analysis.

The bottom-right panel shows the average ECAP slope as a function of the electrode position with the error bar representing the standard error of the mean. In contrast to the tight distribution in the individual EFI data (figure 4), much greater individual variability is apparent in the ECAP slope data. A pairwise post hoc multiple comparison procedure reveals that the ECAP slope is significantly steeper at the middle electrode E9 than at the most basal electrode E16 (Tukey’s HSD criterion, $F(1.8) = 7.88$, $p < 0.05$). The dependence of the ECAP slope on the stimulation site has also been observed in other CI devices and suggested to be...
proportional to local nerve density (e.g., Brill et al 2009, Eisen and Franck 2004). The steepest ECAP slope at the middle electrode is likely a combined result of the ability of the middle electrode to recruit neurons from both sides of the cochlea and generally greater local nerve survival on the apical side than the basal side (e.g., Nadol 1997). Figure 7 shows individual ECAP spatial profiles (rows) as a function of the masking electrode (x-axis) with electrode E1, E9 or E15 being the stimulation site (the vertical dotted line in each panel). In contrast to the EFI profile, the ECAP data show great between-subjects as well as within-subjects variability. For instance, subject S1 shows higher masking on stimulation electrodes E1 and E9 but no masking on E15; subject S3 shows masking on E9 and E15 but no masking on E1; on the other hand, subject S5 shows only minimal masking on E15 but no masking on E1 and E9. This great inter- and intra-subject variability suggests that channel interactions at the neural level cannot be modeled as a simple linear function. Both the source and the implication of this great variability will be discussed later.

3.3. Channel interactions at the perceptual level

Figure 8 shows the individual (rows) channel interaction data at the behavioral threshold level on three probe electrode positions (columns). The masker level is plotted as a function of the probe level, with a positive value representing the in-phase condition and a negative value representing the out-of-phase condition. The horizontal dotted line represents no interaction, whereas the dashed line at $-45^\circ$ represents the total interaction. The threshold channel interaction data can be adequately modeled by a linear function (dashed lines, mean $R^2 = 0.82$, SE = 0.03, $p < 0.05$). In general, the channel interaction index, namely the slope of the fitted linear function, decreases as the distance between the probe and the masker increases. For example, the most basal electrode E15 (diamonds) did not interact with the detection of the most apical probe electrode E1 (left columns) but totally interacted with the detection of itself as expected (right columns). The reverse pattern can be observed between the most apical probe and the most basal masker electrodes.

Figure 9 shows the individual loudness summation data (rows) in response to simultaneous two-electrode stimulation at the three different probe electrode locations (columns). The masker level is plotted as a function of the probe level that reaches comfortable loudness, with different symbols representing different masker electrodes. Despite individual differences in absolute stimulation levels reaching comfortable loudness, all subjects produced consistent patterns of results: direct electric field summation accounts for nearby probe and masker electrodes (the $-45^\circ$ diagonal line), whereas loudness summation for distant probe and masker electrodes (the significantly curved line). The present loudness channel interaction model, equation (5), yielded excellent fit to the loudness summation data (mean $R^2 = 0.94$, SE = 0.06, $p < 0.05$).

3.4. Comparison of channel interactions at different levels

To compare these different channel interaction measures, each measure was normalized by dividing it by its corresponding largest value. Figure 10 shows the normalized spatial profiles of the EFI (squares), ECAP (triangles), behavioral detection threshold (circles) and loudness data (diamonds). Rows represent different subjects and columns represent the probe electrode. There is an expected general trend in this highly variable result: the closer the probe is to the masker, the greater are the interactions observed at all levels. In addition, the data can be roughly classified into the following three patterns. First, 6 of the 15 cases show a consistent correlated pattern between all four measures (E9 in S1, E9 in S2, E15 in S3, E9 and E15 in S4 and E9 in S5). Second, five cases show diverse patterns for four measures (E15 in S1, E1 in S2, S3, S4 and S5). Third, four cases show the same pattern between some of the four measures (CII, CIII and ECAP on E1 in S1 and E9 in S3, EFI and ECAP on E15 in S2 and E15 in S5). In the following section, we define two global indices to quantify a spatial profile and to compare two spatial profiles.
Figure 6. ECAP growth functions for all electrodes (different symbols) in five subjects (individual panels labeled by S1–S5). The dashed lines are the best fit to the ECAP data. The bottom-right panel shows the mean ECAP slope (circles) and its standard error (error bars) as a function of the masker electrode in the cochlea.

3.4.1. Half-width of the spatial profiles. A two-parameter rounded exponential (roex) function was used to fit the spatial profile for each measure,

\[ f(x) = (1 - w) (1 - \alpha |x - x_s|) e^{-\alpha |x - x_s|} + w, \]

where \( x_s \) corresponds to the probe electrode number (1, 9 or 15), and \( \alpha \) and \( w \) correspond to the decay length and decay height of the profile, respectively (Patterson et al. 1982). The roex fit was applied to four different locations, including E1, E9a (apical side of E9), E9b (basal side of E9) and E15. The spatial profile half-width was defined as the distance in the number of electrodes, at which the roex profile had a value of 0.8. Compared with other models, such as a cubic or linear function, the roex fit yielded more consistent and interpretable results (\( R^2 = 0.97 \) for EFI, \( R^2 = 0.67 \) for ECAP, \( R^2 = 0.85 \) for CIIth and \( R^2 = 0.97 \) for CII). Figure 11 shows the mean and standard error of the half-width derived from the four measures at locations E1, E9a (apical side), E9b (basal side) and E15. Several interesting observations can be made from these data. First, the EFI half-width decreases significantly from the apex to the base (mean = 3.3 at E1 and 1.5 at E15; \( F(3,16) = 20.9, p < 0.05 \)). This decrease in the EFI half-width from the apex to the base is consistent with several previous psychophysical and physiological studies, which found larger electrode interactions at the apex than at the base (Favre and Pelizzzone 1993, Eisen and Franck 2004, Cohen et al. 2003, Abbas et al. 2004, Hughes and Abbas 2006b, Chatterjee and Shannon 1998, Stickney et al. 2006). On the other hand, no such systematic difference is apparent in the half-width for the other three measures. Second, the average ECAP half-width across electrodes (mean = 1.47; SE = 0.49) is significantly narrower than the EFI width (mean = 2.28; SE = 0.37; \( F(1,38) = 9.71, p < 0.01 \)). This difference can be understood because not all electric fields would excite a neural response and, additionally, even if an electric field is strong enough, it may not elicit any neural response due to ganglion cell loss. Third, the EFI half-width data have the least variability (SE/mean = 0.08), the ECAP the most (0.26) and the CIIth (0.19) and CII (0.18) in the middle. The variability analysis suggests that behavioral channel interactions are likely dependent upon both the uniform intracochlear electric field distribution and the heterogeneous nerve survival pattern.

3.4.2. Asymmetry of the spatial profiles. For a given spatial measure \( S \), its asymmetry index (AI) can be defined as the...
percentage of the mean difference between corresponding apical \( (i = 2–8) \) and basal \( (i = 10–16) \) values divided by the mean value of the overall spatial profile

\[
AI (\%) = \frac{100 \sum_{i=2}^{8} (S_i - S_{i+8})}{\sum_{i=2}^{16} S_i / 14},
\]

where a positive AI corresponded to more spread on the apical side than on the basal side; a zero AI indicated perfect symmetry between the apical and basal sides in the spatial profile.

Figure 7. Individual ECAP spatial profiles (five rows labeled by S1–S5) as a function of the masker electrode produced by stimulating E1 (left panels), E9 (middle panels) and E15 (right panels).

Figure 12 shows a box plot of the AI for the five spatial profiles at the E9, including EFI, ECAP slope, ECAP, CIIth and CIIl. One consistent finding is that the EFI profile has a positive AI for all subjects (mean = 11.42%; SE = 2.15%; \( t(4) = 6.92, p < 0.01 \)), indicating greater spread of current toward the apex than the base. This asymmetry in the intracochlear voltage profile was noted in previous studies and was thought to be a consequence of the low-resistance path of the current flow from the apex to the base (e.g., Jolly et al 1996, Vanpoucke et al 2004a, 2004b). The other consistent finding is that the two neural and the two behavioral measures, particularly the ECAP slope, have much greater variability than the electric measure. The lack of apparent dependence of the two behavioral measures on either the EFI or the ECAP measure reinforces the suggestion from the half-width analysis that the behavioral measure is dependent upon both electric and neural properties.

4. Discussion

4.1. Mechanisms of channel interaction

Channel interaction is subject to electric stimulus and its transformation from electrical potentials to neural responses and perceptual interpretation. The transformation from electric potentials to neural responses is primarily determined by an ‘all-or-none’ threshold mechanism, followed by a facilitating or inhibitory process when the two electric stimuli
Figure 8. Individual threshold channel interaction profiles (five rows labeled by S1–S5) for three probe conditions: E1 (left panels), E9 (middle panels) and E15 (right panels). Different symbols represent different masker electrodes. The dashed lines are the best fit to the threshold channel interaction index (see equation (1) in the text for details).

are close to each other in time. Because the highest rate was 100 Hz, equivalent to a 10 ms interval between two adjacent pulses, the present result was unlikely affected by this temporal interaction of the electric-to-neural transformation. The transformation from neural firing to perceptual measures, such as threshold and loudness, is not clear but is certainly a...
Figure 9. Individual loudness channel interaction profiles (five rows labeled by S1–S5) for three probe conditions: E1 (left panels), E9 (middle panels) and E15 (right panels). Different symbols represent different masker electrodes.
nonlinear mechanism. The effect of these transformations on threshold and loudness has been studied in electric hearing, assuming a generally uniform nerve survival pattern (Bruce et al 1999, Xu and Collins 2005). If the neurons are uniformly distributed along the cochlea, then channel interactions are determined by the electric potential distribution and its transformations only. Thus, all channel interaction spatial profiles should be relatively similar to the electric field profile, including more masking on the apical side than the basal side. Subjects 3, 4 and 5 generally showed these predicted patterns (figure 10) and therefore presumably had a relatively uniform distribution of the survived neurons. In contrast, in cases of the poor electrode–neuron interface due to a dead region, or cochlear ossification, or simply the great distance between electrodes and neurons (e.g., Bierer et al 2010, Moore 2004), these electrodes will function interactively because they will likely activate the same group of the closest neurons. For example, the saturated channel interaction index for apical electrodes from 1 to 5 in S1 and basal electrodes from 8 to 16 in S2 (figure 10), indicated either the dead regions underneath these electrodes or the large distance between the electrodes and the neurons. Unfortunately, both possibilities were consistent with the elevated evoked neural thresholds for these electrodes (figure 6). Alternative methods need to be explored to differentiate between dead region and distant stimulation in the future.

4.2. Relation to function

Channel interaction is frequently related to CI speech performance (e.g., Bierer 2010). This study produced no significant correlation between speech performance (table 1) and any of the channel interaction measures. The largest, but still insignificant, correlation was found between speech scores in quiet and threshold interaction ($R^2 = 0.29$) and speech scores in quiet and ECAP slope ($R^2 = 0.23$). One reason for a lack of correlation was the discrepancy in the stimulation rate between...
Figure 11. Mean and standard error of the half-width for the EFI (first bar), ECAP (second bar), threshold (CIIth, third bar) and loudness (CII, fourth bar) spatial profiles at different locations of the probe electrode (x-axis). See the text for details.

Figure 12. Box plot of the AI for the EFI, ECAP slope, ECAP, threshold (CIIth) and loudness (CII) spatial profile. The box represents the 25–75% range, the error bars represent the min–max range, and the line within the box represents the median. See the text for details.

the present measures (<100 Hz) and speech processors (>500 Hz). Another reason for this lack of correlation was the small sample size in this study. To get meaningful correlation data, strategically simplified channel interaction measures need to be conducted over a large sample size. For example, the loudness interaction would be correlated with the detection of spectral peaks and vowel recognition (e.g., Henry and Turner 2003).

4.3. Implications

It is worth noting that alternative approaches have been proposed to reduce the spatial channel interaction problem in electric stimulation. Instead of intracochlear electrode placement, the auditory nerve implant directly places a thin-film microelectrode array in the auditory nerve bundle, shortening the stimulation path and reducing channel interaction (Middlebrooks and Snyder 2007). Similarly, a new version of the auditory brainstem implant uses bundled needle-like electrodes to penetrate the tissue and directly place the electrode next to the nerve (McCreery 2008). To avoid the widespread problem of electric stimulation altogether, optical stimulation has been used to greatly reduce channel interaction because only those neural tissues that are in the direct path of the laser can be stimulated (Izzo et al. 2007). It is also worth noting that, except for auditory prostheses that require high rate stimulation, other neural prostheses use the stimulation rate that rarely exceeds several hundred hertz. Instead, spatial channel interaction is a particularly significant problem in these neural prostheses. For example, a retinal implant has to provide sufficient independent pixels to support a meaningful visual function (e.g., Chader et al. 2009). A vestibular implant has to provide independent functional channels while limiting current spread to avoid undesirable auditory or other facial nerve stimulation (Fridman et al. 2010). Similarly, targeted electric stimulation is important for avoiding side effects while increasing efficacy in spinal cord stimulation, deep brain stimulation and other neural modulation devices (e.g., Kuncel and Grill 2004, Sankarasubramanian et al. 2011, Pikov et al. 2010). It is conceivable that the present approach characterizing spatial channel interactions at the physical, neural and behavioral levels can be generalized to all forms of physical energy, from electric to magnetic or optical stimulation, as well as to most neural prostheses and modulation devices.

5. Conclusions

A functional channel carries independent information. The number of functional channels can be different from, and generally less than, the number of physical electrodes, depending on electric stimulation parameters and the interfacing neural tissue. This study used low-rate (<100 Hz) stimulation to focus on spatial channel interaction. Although spatial channel interactions have been measured at electric, neural and behavioral levels previously, this study provided systematical measures from the same subjects, allowing direct comparison and identification of inter- and intra-subject variability sources. By quantitatively comparing global as well as local spatial profiles across these different measures, this study found the following results:

(1) Electric potential distribution is broad but uniform within- and between-subjects. Channel interaction at the physical level can be modeled as a linear system in the present cochlear implant system.

(2) In contrast, neural survival patterns as reflected by evoked potentials vary significantly, producing highly variable neural interactions within and between subjects.

(3) A novel model combining linear electric field summation and exponential loudness growth can capture channel interaction at a behaviorally relevant level. Behavioral channel interaction is determined by at least three factors, including the electrical field distribution, the peripheral nerve survival pattern and the central nonlinear transformation.

In the foreseeable future, electricity will likely remain the dominant means of stimulation in neural prostheses. As
a result, channel interaction in electric stimulation remains a significant challenge. It is critical to understand the distribution of the electric field in a biological environment, and the interaction between the electric field and the nerve tissue for these factors plays a key role in determining the ultimate performance of neural prostheses.

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